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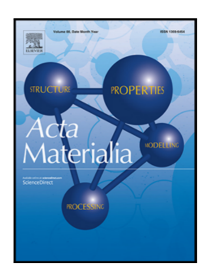
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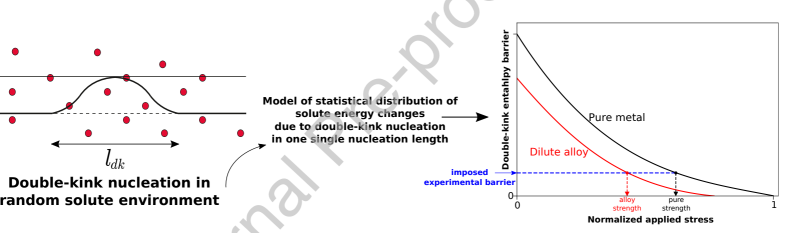
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Theory of double-kink nucleation in dilute BCC alloys

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Abstract

Yielding in pure BCC metals and dilute substitutional alloys occurs by doublekink nucleation and propagation along screw dislocations . At low temperatures, the yield stress is controlled by double-kink nucleation. Here, an analytical statistical model is presented to predict the stress- and length-dependent doublekink nucleation barrier in dilute BCC alloys solely in terms of the double-kink process in the pure metal and the solute/screw-dislocation interaction energies in the dilute alloy. Consistent with early literature, dilute alloying always reduces the double-kink nucleation barrier (softening) independent of solutes or matrix. The model is extensively validated via simulations in model Fe-Si alloys described by interatomic potentials. The model is then compared to experiments on real Fe-Si, W-Ta, and W-Re alloys, showing qualitative agreement consistent with the accuracy of the inputs. A cross-over from the dilute limit to the nondilute limit, where there is hardening, is analyzed using the present theory and the non-dilute theory of Maresca et al. The analysis for Fe-Si is consistent with a cross-over at $\approx 2 - 3$ at. %Si, as observed experimentally, and qualitatively consistent with W-Ta and W-Re. The present theory plus the recent theory of Maresca et al. together provide a coherent predictive framework for strengthening of screw dislocations over the full range of concentrations from extremely dilute ($\ll 1$ at.%), to dilute (up to a few at.%) and non-dilute alloys including High Entropy Alloys.

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Keywords: BCC; Dilute alloys; Double-kink nucleation; Screw dislocations; Solute softening

1. Introduction

In BCC metals and dilute substitutional alloys, plastic deformation is controlled by the thermally-activated motion of screw dislocations by a two-step process. The first step is the nucleation of a pair of kinks along an initial long straight screw dislocation. The second step is the migration of these kinks along the screw that ultimately advances the entire screw dislocation forward by one Peierls valley. In pure BCC metals, the double-kink nucleation is the rate-limiting process because kink migration has a negligible barrier. In dilute solid solution BCC alloys, the solutes modify the barriers for both double-kink nucleation and kink migration. The controlling process is that with the largest barrier; this requires a statistical determination of the relevant barriers as a function of the applied stress and concentration. In general, the double-kink 12 nucleation barrier is reduced by solutes because nucleation occurs in that region 13 where the solute fluctuations in the random alloy will most-favor formation of 14 the double kink. Kink migration must then overcome the largest solute-induced barriers for migration along the remaining line length. With increasing solute 16 concentration, the conventional understanding is that the nucleation barrier is 17 steadily reduced and the kink migration barrier steadily increased such that 18 there can be a cross-over from control by nucleation to control by migration. 19 This can lead to a transition from softening (reduced flow stress, relative to the pure metal, with increasing concentration) to hardening (increasing flow stress 21 with increasing concentration). Such behavior has been predicted via models [1] and observed in recent Monte Carlo simulations in both Fe-Si [2] and W-Re 23 [3], although results depend on details of the models and simulation methods. The softening by solutes in the dilute limit was recognized long ago [4, 5, 6]. Considering the attractive force of impurity atoms on a dislocation line, Weertman [4] showed that the impurity atoms could reduce the Peierls energy in their

immediate neighborhood, leading to a reduction in the required stress. Softening can be achieved, however, with either attractive or repulsive solutes (see Figure 1). For repulsive solutes, an initial straight dislocation is simply pushed away from the solute, and the double-kink nucleation barrier is reduced. There 31 is substantial evidence of softening due to small additions of many solutes in 32 many BCC elements at low temperatures [5, 7]. Suzuki [6] developed a statistical model for solute effects on double-kink nucleation and kink-migration processes. He envisioned a single solute/dislocation interaction energy E_0 and 35 considered that somewhere along the dislocation line there could be m solutes within the double-kink nucleation length. Suzuki then postulated that the de-37 crease in double-kink nucleation energy would be $-m|E_0|/2$ where the absolute value accounts for the fact that both attractive and repulsive solutes could reduce the barrier and where the factor of 1/2 was postulated as the reduction in barrier following simple chemical kinetics models. The value of m was then computed as the largest value along a dislocation line length L. The Suzuki model 42 thus contains the key features of the problem, but with crucial assumptions about the solute/dislocation interaction energies and their effects on the nucleation barrier that limit quantitative predictions. That is, the single parameter E_0 , the statistical factor m, and the factor of 1/2, are all ad-hoc approximations that are not accurate in realistic systems.

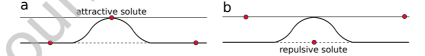


Figure 1: (a) An attractive and (b) a repulsive solute pulling and pushing the initial dislocation line away, respectively, towards the next Peierls valley. In both cases, the double-kink nucleation barrier is reduced.

In spite of the above early studies, the recent literature [1, 8] makes a different set of assumptions. Retaining only a single solute/dislocation interaction energy E_0 , these works postulate that the rate of double-kink nucleation along a long line is changed by the factor $c e^{-E_0/kT}$ where c is the solute concentration.

This model thus effectively considers only individual solutes (the very dilute limit) with no statistical effects and, moreover, predicts that repulsive solutes $(E_0 > 0)$ decrease the double-kink nucleation rate (leading to increased yield stress). Solutes with positive and negative interaction energy are thus predicted 55 to be fundamentally different [8] (hardening vs. softening). This is in conflict with experiments at the yield point. For instance, as shown later below, data of 57 Stephens [7] shows softening at low temperature and low solute concentrations for both W-Ta and W-Re alloys even though first-principles DFT predicts the 59 interaction energy of Ta to be positive and that of Re to be negative [8] (see 60 figures 2(a) and (c)). Other comparisons with experiments were made using 61 hardness data [1], corresponding to plastic strains of \sim 7-10% while the data in Ref. [7] shows that behavior at yield and at a few percent plastic strain can lead to conflicting conclusions regarding softening versus hardening. 64 Prior analyses [1, 9] also included an additional direct change to the Peierls 65

Prior analyses [1, 9] also included an additional direct change to the Peierls stress in an alloy. This contribution can be viewed as the Peierls stress of the effective homogeneous representation of the alloy. Such a representation is important at non-dilute concentrations where the identities of solute and matrix become less clear [10]. In the dilute limit, the explicit dominant effects of the individual solute additions are clear and provide perturbations to the double-kink nucleation barrier of the underlying matrix material. These effects change the apparent Peierls stress, as discussed below, with no additional considerations.

Here, in light of the limitations of previous analyses as sketched above, we re-consider double-kink nucleation in the presence of low concentrations (a few percent or below) of solutes. We present and validate a general analytical statistical model for the effects of solutes on the double-kink nucleation barrier as a function of concentration and applied stress. This model is a significant improvement over the simplified model of Suzuki, and reveals the limitations of other models. It should be noted that all screw-dislocation-based theories to date including the present model assume that the screw dislocation core is the well-defined compact symmetric core. This is true for dilute alloys [8, 11, 12, 13] of interest here, but first-principles studies by Rao et al. [14], Romaner et al.

[10], and Li et al. [15] show that screw cores may not always remain symmetric and compact in some non-dilute alloys.

In applications of the model, we obtain reasonable agreement with experiments at low concentrations where double-kink nucleation is expected to dominate. With increasing concentration, we then show a cross-over from softening, 87 controlled by the present theory, to hardening, controlled by the recent nondilute theory of Maresca and Curtin [16]. The Maresca-Curtin theory argues that the low-energy state of a screw dislocation is intrinsically kinked over a char-90 acteristic length scale, making double-kink nucleation irrelevant. The cross-over 91 from softening to hardening is then mainly related to whether the spacing of 92 the intrinsic kinks is smaller than the typical dislocation length between dislocation junctions. If not, then the dilute theory (softening) applies. If so, then the non-dilute theory (hardening) applies. The strength in the non-dilute domain of the Maresca-Curtin theory is still controlled by some combination of kink-migration, cross-kink formation/pinning of screws, and, at very low temperatures, a Peierls-like motion. These features are similar to, although different in detail than, those postulated by Suzuki [6]. So, both new [16] and old 99 [6] theories, while different, show that double-kink nucleation does not control 100 strengthening at higher concentrations. The present theory and that of Maresca 101 et al. thus bridge across the full composition range of BCC alloys from very 102 dilute up to multicomponent non-dilute High Entropy Alloys.

The remainder of this paper is organized as follows. In section 2, we discuss 104 the effect of multiple solutes on the double-kink nucleation barrier. A statisti-105 cal model to predict the double-kink nucleation barrier as a function of solute 106 concentration, dislocation length, and applied stress is presented. The effect of 107 single solutes, relevant in the very dilute limit, is addressed in Section 3. In Section 4, we validate our analytic statistical model using atomistic simulations 109 on a model Fe-Si alloy. In Section 5, we discuss the general application of our 110 results, make specific comparisons for Fe-Si, W-Ta, and W-Re, show the cross-111 over from dilute to non-dilute in Fe-Si, and discuss further implications of our work. Section 6 summarizes the paper. 113

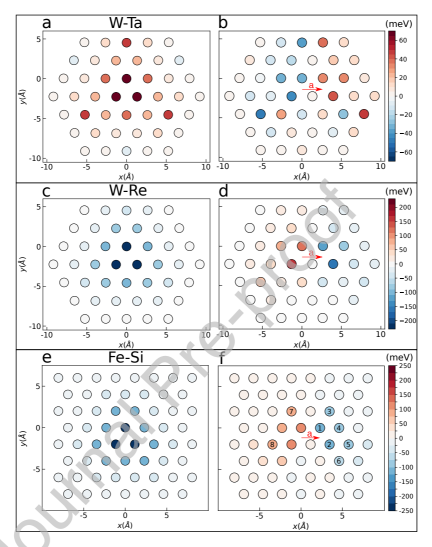


Figure 2. Solute/screw dislocation interaction energy versus solute position for $\bf a$ Ta [8, 17] and $\bf c$ Re [8, 17] in single crystal BCC W as calculated by DFT and $\bf e$ Si in Fe as computed using the EAM potential of [2]. Interaction energy difference as the screw dislocation core moves by one Peierls valley distance a (red arrow) for $\bf b$ Ta and $\bf d$ Re solutes in W and $\bf f$ Si in Fe. Sites marked with 1-8 in $\bf f$ are substituted with Si atoms for subsequent NEB calculations, and are referred to in Figs. 5 and 6.

2. Solute effects on the double-kink nucleation barrier

2.1. Basic quantities

We consider a binary dilute alloy in which there is only one type of substi-116 tutional solute at concentration c and the other component has a high concen-117 tration and serves as the "matrix". The model is easily generalized to multi-118 ple components. The solute concentration c is low (below a few percent) but 119 above a "very dilute limit" defined later by a critical concentration c^* . We con-120 sider solutes for which the maximum interaction energy change as a dislocation 121 moves past an individual solute is smaller than the single dislocation kink en-122 ergy. Then, an individual solute does not create a kink migration barrier that is 123 larger than the solute-reduced kink nucleation barrier. Since kink energies are estimated to be 0.45 eV or larger [18] , this limit holds for many substitutional 125 solutes in most BCC metals although there are exceptions such as Co, Rh, Ir, 126 Ni, Pd, and Pt in W according to DFT results of Hu et al. [8]. Interstitial 127 solutes may also have large interaction energies, and so are not considered here. Underpinning double-kink nucleation in the alloy is the stress-dependent 129 double-kink nucleation barrier and transition state in the pure matrix. Fig. 3 130 shows the transition state configurations for pure Fe at various applied stresses, 131 as obtained using a reasonable EAM potential for Fe [19] (see [20] for a study 132 using a machine-learning potential and [18] for configurations obtained using a 133 line tension model). At zero stress, the transition state is symmetric, has two 134 fully-developed kinks, and spans 1/2 the simulation cell size. With increasing 135 applied stress, the initial position of the dislocation moves away from the zero-136 stress Peierls valley to position $x_{eq}(\tau)$. More importantly, the critical double-137 kink configuration becomes progressively smaller: the kinks are closer together and the kink is not fully-developed. Only at low stresses does the kink extend 139 into the next Peierls energy minimum at distance $a = 0.943 \, b$ from the original 140 minimum. An individual kink typical spans a length > 10 b so that the character 141 angle θ of the kink segment is $\theta < \tan^{-1}(0.943/10) \approx 5^{\circ}$, i.e. very close to pure screw character.

The dislocation shape $x_{\rm disl}(z,\tau)$ of the double-kink transition state versus applied stress can be expressed generally as

$$x_{\text{disl}}(z,\tau)/a = g(z,\tau) + g_{\text{eq}}(\tau) \tag{1}$$

where $g_{\rm eq}(\tau) = x_{\rm eq}(\tau)/a$ is the dimensionless initial straight screw position at stress τ and $g(z,\tau)$ is the dimensionless kink shape, or deviation from the initial straight dislocation, as a function of τ , At low stresses (e.g. 100 MPa in Figure 3), the nucleation occurs over a length l_{dk} approximately equal to that of the two kink widths. Across a range of systems, this length is $l_{dk} \approx 20b-25b$. The geometry of the transition-state double-kink structure versus stress serves as one main input for our analysis of solute effects on nucleation.

Associated with each stress and transition state is the nucleation enthalpy barrier $\Delta H^0(\tau)$ of the pure matrix. The functional form of $\Delta H^0(\tau)$ is not directly relevant for most of our analysis below, but is needed for predictions. This barrier can be obtained by simulations, models, or fitting. The latter often takes the form of a Kocks law [21] $\Delta H^0(\tau) = \Delta H^0(0) \left[1 - (\tau/\tau_p)^p\right]^q$ where $0 \le p \le 1$ and $1 \le q \le 2$ are fitting parameters and $\Delta H^0(0)$ and τ_p are the zero-stress enthalpy barrier and Peierls stress, respectively.

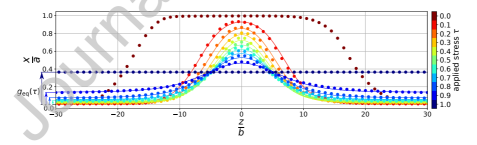


Figure 3: Transition state configurations $x_{\rm disl}(z,\tau)/a$ versus applied shear stress τ determined using pure Fe EAM potential [19]. The points are the direct outcome of the disregistry method [22]. The solid lines are the fits to the functional form for $g(z,\tau)$ given by Eq. (19) with parameteres shown in Table 1. On the lower left, the equilibrium position $g_{\rm eq}(\tau)$ of the initial straight dislocation is indicated by the arrows for several applied stresses.

The second major quantity of importance is the interaction energy between a

type-n solute at position (x_i, y_j, z_k) and a straight screw dislocation positioned 161 in the Peierls valley minimum at the origin and aligned along the z axis. This interaction energy is denoted as U_{ijk}^n . Fig. 2(e) shows U_{ijk}^n for many solute 163 positions ij around the core within the periodic length b along the line direction 164 z for Si solutes in Fe as computed using a model Fe-Si alloy described by EAM 165 potentials [2]. The solute/dislocation interaction energies for Ta and Re solutes in single crystal BCC W as computed by Density Functional Theory (DFT) [8, 17] are shown in Figs. 2(a) and (c); quantitatively similar results are reported 168 in other works [11]. With this projection, the interaction energy is independent 169 of the z coordinate. For the model Fe-Si and W-Re systems all solute/dislocation 170 interactions are negative (attractive) while the interactions for Ta in W are all positive (repulsive). The interaction energies are not localized to the first or 172 second neighbors around the core, as commonly assumed in theories that use 173 these computed energies to predict strengths [6, 1, 8]. 174

Dislocation motion is determined by the *changes* in solute/dislocation energy 175 as a dislocation glides. Fig. 2(f) shows the change in energy of a Si solute in Fe upon glide of the screw dislocation by one Peierls valley distance, a. Figs. 2(b) 177 and (d) show the corresponding DFT energy changes for Ta and Re solutes in 178 W matrix, respectively. Non-negligible changes in energy can exist for solutes 179 located at initial positions up to sixth neighbors from the center line of the 180 core. As discussed extensively in Ref. [23], and important in the present work, it is necessary to consider solutes out to such distant neighbors to capture the 182 relevant energy changes accurately. These interaction energies and changes upon 183 glide are the fundamental quantities driving double-kink nucleation in a dilute 184 alloy.

2.2. Solute/double-kink interaction energy

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Direct computation of the solute/double-kink interaction energies for all possible solute positions around the (transition state) double-kink structure is not possible. Computation of an interaction energy requires full relaxation of the system containing the solute and dislocation but the double-kink structure is a

constrained state and is not a low-energy relaxed equilibrium structure. Here, we thus make a model assumption and then validate that assumption via direct simulations in Section 4.2.

We assume that the solute/double-kink interaction energy at each stress and for a single solute at position (x_i, y_j, z_k) is

where $\Delta U_{ij}(a) = U(x_i - a, y_j) - U(x_i, y_j)$ is the solute/screw dislocation energy

change as the dislocation moves by one Peierls valley distance a (Figures 2b,d,f).

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$$E_{dk}(x_i, y_j, z_k, \tau) = \frac{x_{disl}(z_k, \tau)}{a} \Delta U_{ij}(a) + U(x_i, y_j),$$
(2)

That is, the solute/double-kink interaction energy of a solute in plane z is equal to the full energy change upon glide by a scaled by the distance x of 199 the kink in the same z plane. This approximation is reasonable because, as 200 shown above, the character of the kink is very near screw. Therefore, a solute 201 experiences a surrounding region that is very nearly a straight screw dislocation at position $x_{\text{disl}}(z_k, \tau)$. This approximation also satisfies the limiting cases where the dislocation is in the initial and final Peierls valleys, i.e. $x_{\rm disl}(z_k,\tau) \to 0$ and 204 $x_{\text{disl}}(z_k, \tau) \to a$. 205 In the presence of any specific arrangement of solutes, the change in nu-206 cleation barrier is the sum of the changes, whether positive or negative, con-207 tributed by each individual solute. In the dilute limit, we can safely neglect any effects of multi-solute interactions because the average solute-solute sepa-209 ration is large ($\approx b/c^{\frac{1}{3}}$); each solute thus has an effect independent of any other 210 solutes. A specific random arrangement of solutes is represented by a set of 21 occupation variables s_{ijk} where $s_{ijk} = 1$ if a solute is at position (x_i, y_j, z_k) and = 0 otherwise. Overall all sites $N \gg 1$ in the alloy, $1/N \sum_{i,j,k} s_{ijk} = c$. 213 The double-kink nucleation barrier is then the energy difference between all so-214 lutes interacting with the transition state structure $g(z,\tau) + g_{eq}(\tau)$ and with 215 the initial straight dislocation at $g_{eq}(\tau)$, and so involves only the non-straight 216 portion $g(z,\tau) = x_{\text{disl}}(z_k,\tau)/a$ of the transition state shape. Using Eq. (2) for the single solute/double-kink interaction energy, the solute contribution to the 218

nucleation barrier is computed by summing the contributions of all the solutes 219 in the specific realization of the random alloy as

$$\Delta H^{\text{sol}}(\tau) = \sum_{i,j,k} s_{ijk} g(z_k, \tau) \Delta U_{ij}(a)$$
(3)

2.3. Model for solute softening in dilute alloys

For a dislocation segment of length l_{dk} , the mean energy change over all possible fluctuations in the positions of the solutes is zero. This is shown by averaging Eq. (3) over all realizations s_{ijk} . However, the local fluctuations in the spatial arrangement of solutes give rise to a statistical distribution of energy changes of the nucleated dislocation segment. The standard deviation of this energy change is

$$\sigma_{\Delta H^{\text{sol}}} = \left[\left\langle \Delta H^{\text{sol}}(l_{dk}, x_{\text{disl}})^2 \right\rangle - \left\langle \Delta H^{\text{sol}}(l_{dk}, x_{\text{disl}}) \right\rangle^2 \right]^{\frac{1}{2}}, \tag{4}$$

where brackets denote averaging over the random occupation variables s_{ijk} . Carrying out the averaging and algebra leads to (Appendix A)

$$\sigma_{\Delta H^{\text{sol}}} = \Delta \tilde{E}_p(a) G(\tau), \tag{5}$$

where

$$\sigma_{\Delta H^{\text{sol}}} = \Delta \tilde{E}_p(a) G(\tau), \tag{5}$$

$$G(\tau) = \left(\sum_{k=-N_s}^{N_s} g(k \, b, \tau)^2\right)^{\frac{1}{2}}, \tag{6}$$

is a geometrical shape factor and $N_s = l_{dk}/(2b)$ is the total number of atomic sites along the dislocation line z within one kink length $l_{dk}/2$. The standard deviation of the statistical distribution of nucleation barriers at length l_{dk} thus scales with the quantity 227

$$\Delta \tilde{E}_p(a) = \left(\sum_{ij} c \,\Delta U_{ij}(a)^2\right)^{\frac{1}{2}};\tag{7}$$

this is precisely the solute/screw dislocation interaction energy parameter that enters in the theory for yield strength of a BCC alloy [16, 23], but reduced to the

dilute binary alloy limit. This feature will later enable a bridging between dilute and non-dilute alloys. The shape of the double-kink transition state enters only as an overall stress-dependent numerical factor $G(\tau)$. Considering one nucleation length l_{dk} , the random fluctuations in solutes

Considering one nucleation length l_{dk} , the random fluctuations in solutes leads to a Gaussian probability distribution of the solute contributions to the double-kink nucleation enthalpy $\Delta H^{\rm sol}$ as

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$$P[\Delta H^{\text{sol}}] = \frac{1}{\sqrt{2\pi}\sigma_{\Delta H^{\text{sol}}}} \exp\left[-\frac{1}{2} \left(\frac{\Delta H^{\text{sol}}}{\sigma_{\Delta H^{\text{sol}}}}\right)^{2}\right]. \tag{8}$$

The above result is for a dislocation of length l_{dk} only. A longer dislocation line of length L can be considered approximately as a set of $N=L/l_{dk}$ statistically-independent segments of length l_{dk} , and double-kink nucleation could initiate at any one of these segments. Each segment has a distinct local solute configuration and hence has a local double-kink nucleation barrier that is sampled randomly from the above statistical distribution. The total rate of double-kink nucleation due to nucleation at all sites is the sum of the rates of all N segments, which can be written as (see Appendix B)

$$R = \nu_0 N \exp\left(-\frac{\Delta H^0(\tau)}{kT}\right) \exp\left(\frac{\sigma_{\Delta H^{\text{sol}}}(c,\tau)}{\sqrt{2}kT}\right)^2 \times \frac{1}{2} \operatorname{erfc}\left(\frac{\Delta \bar{H}^{\text{sol}}(N,c,\tau)}{\sqrt{2}\sigma_{\Delta H^{\text{sol}}}(c,\tau)} + \frac{\sigma_{\Delta H^{\text{sol}}}(c,\tau)}{\sqrt{2}kT}\right), \tag{9}$$

where $\Delta \bar{H}^{\rm sol}$ is the the largest reduction in activation energy among all the N segments (the "weakest-link", see below), k is Boltzmann's constant, ν_0 is an appropriate attempt frequency, and erfc is the complementary error function. In the absence of solutes, or for very weak solute/screw interactions, the nucleation rate is $R = \nu_0 N \exp\left(-\frac{\Delta H^0(\tau)}{kT}\right)$ which is the rate for a dislocation length in the pure matrix.

The temperature- and stress-dependent plastic shear strain rate $\dot{\epsilon}$ at applied stress τ then follows from Orowan's law as $\dot{\epsilon} \approx \rho baR$, where ρ is the mobile dislocation density and R is the nucleation rate. Using this relationship, the

yield stress as a function of temperature and strain rate can be determined, as described in more detail in the context of the applications presented in Section 5.

At sufficiently low temperatures, the rate of double-kink nucleation is controlled by that one segment having the greatest reduction of the kink-pair nucleation barrier; i.e. nucleation is an extreme-value (weak-link) problem. For a dislocation of length Nl_{dk} , the mean of the lowest activation energy is

$$\Delta \bar{H}^{\text{sol}} = \frac{\int\limits_{-\infty}^{\infty} \dots \int\limits_{-\infty}^{X_3} \int\limits_{-\infty}^{X_2} X_1 \prod\limits_{n=1}^{N} P(X_n) \, \mathrm{d}X_n}{\int\limits_{-\infty}^{\infty} \dots \int\limits_{-\infty}^{\infty} \int\limits_{-\infty}^{X_3} \prod\limits_{n=1}^{X_2} P(X_n) \, \mathrm{d}X_n},$$
(10)

The standard deviation of activation energies around this mean lowest value is

$$\bar{\sigma}_{\Delta H^{\text{sol}}}^{2} = \frac{\int_{-\infty}^{\infty} \dots \int_{-\infty}^{X_{3}} \int_{-\infty}^{X_{2}} (X_{1} - \Delta \bar{H}^{\text{sol}})^{2} \prod_{n=1}^{N} P(X_{n}) dX_{n}}{\int_{-\infty}^{\infty} \dots \int_{-\infty}^{X_{3}} \int_{-\infty}^{X_{2}} \prod_{n=1}^{N} P(X_{n}) dX_{n}},$$
(11)

For long lengths L, corresponding to large N and relevant for real physical systems, the above distributions are asymptotically approximated by the extremevalue Gumbel distribution having the cumulative probability

$$C_N(\Delta H^{\text{sol}}) \approx 1 - e^{-e^{\frac{\Delta H^{\text{sol}} - \sigma_{\Delta H^{\text{sol}}} b_N}{\sigma_{\Delta H^{\text{sol}}} a_N}}}$$
 (12)

where the Gumbel scale a_N and location b_N parameters are

$$a_{N} = \frac{1}{\sqrt{2\log(N)}},$$

$$b_{N} = \frac{\log(\log(N)) + \log(4\pi)}{\sqrt{8\log(N)}} - \sqrt{2\log(N)},$$
(13)

respectively. In terms of $\sigma_{\Delta H^{\rm sol}}$, the mean and standard deviation at length Nl_{dk} are

$$\Delta \bar{H}^{\text{sol}}(c, N, \tau) = \sigma_{\Delta H^{\text{sol}}}(c, \tau) (b_N - \gamma a_N),$$

$$\bar{\sigma}_{\Delta H^{\text{sol}}}(c, N, \tau) = \sigma_{\Delta H^{\text{sol}}}(c, \tau) \frac{\pi}{\sqrt{6}} a_N,$$
(14)

where $\gamma \approx 0.5772$. Numerical tests show these asymptotic results to be quite accurate for N>15 while exact results from Eqs. (10) and (11) can be computed for N<15. One feature of the weak-link analysis is that the mean contribution of solutes increases with increasing N but the standard deviation decreases. Also, the temperature T_0 below which the weak-link dominates can be determined (see Appendix B) and the double-kink nucleation rate can be expressed as

$$R = \nu_0 \exp\left(-\frac{\Delta H^0(\tau) + \Delta \bar{H}^{\text{sol}}(c, N, \tau)}{kT}\right),\tag{15}$$

The weak-link limit provides an easy-to-use analytic form.

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Eqs. (5), (6), and (8) together with Eq. (10) (for N < 15), and Eqs. (13) and (14) (for N > 15) constitute our analytic statistical model for the reduction in double-kink nucleation barrier due to solutes in the dilute limit. The only inputs to the theory are the matrix transition state shape versus stress, the solute/dislocation interaction energies, and the dislocation line length Nl_{dk} .

Lastly, the solute/double-kink interaction energies are modeled using the 262 transition state configurations of the pure matrix at the relevant applied stress. 263 However, a careful treatment is required at zero stress since the transition state 264 in the pure metal is strictly equal to 1/2 the total simulated dislocation length. 265 As shown below, even in the very dilute limit where the transition is affected 266 only by one isolated solute, the transition state at zero stress is always localized around the position of a single favorable solute. The transition state structure is never that found in the pure material (spanning 1/2 the cell size). There-269 fore, here we use the transition state configuration at the lowest non-zero stress 270 where the kinks are well-formed but only separated enough such that the lead-271 ing part of the transition state is very nearly in the final-state Peierls valley. For our model system, this corresponds to the configuration at 0.1 GPa in Fig-273 ure 3. In all subsequent analyses, the transition state configuration at zero 274 stress $x_{\text{disl}}(z_k,0)$ refers to this configuration. This assumption enters the model 275 only through the structure-dependent factor G(0); any other approximation to 276 the zero-stress transition state structure would simply correspond to a (small)

change in the value of G(0). We validate our zero-stress transition state structure via direct simulations for specific solute configurations in a model Fe-Si alloy in Section 4.2.

3. Solute effects in the very dilute limit

The previous analysis is valid for concentrations above a critical value in which multiple solutes are present across the double-kink nucleation length l_{dk} so that a statistical analysis is appropriate. In the very dilute limit, however, at most only a single solute will exist around the dislocation over length l_{dk} and its affect on the double-kink nucleation requires special treatment as discussed here.

Far from the screw dislocation core, the solute/dislocation interaction energy differences are vanishingly small and so do not affect the double-kink nucleation 289 barrier. There are thus only some finite number of distinct atomic sites that 290 contribute to the double-kink nucleation energy difference. We denote the num-29 ber of atomic sites with appreciable negative interaction energy differences per unit b of line length as N_T . Due to symmetry, there are then also N_T sites 293 with positive appreciable energy difference. A critical concentration is then de-294 fined as $c^* = b/(2N_T l_{dk})$, at which there exists, on average, only one solute among all the $2N_T$ -type sites over the entire double-kink nucleation length l_{dk} . For concentrations above this critical concentration, multiple solutes are found 297 within the double-kink nucleation length and the analysis of the previous sec-298 tion quickly becomes relevant. When $c < c^*$, there will be only one solute or no 299 solute within each double-kink nucleation length. The concentration c^* is thus the boundary between the very dilute limit and the dilute limit cases (see Fig. 4). 302

In the very dilute limit, the change in interaction energy of a solute initially at the *i*-th distinct site $(i = 1, ...N_T)$ due to motion of the straight screw dislocation from the initial to the final is now labeled simply as ΔU_i . A dislocation of length L with $N = L/l_{dk}$ double-kink nucleation segments will have a fraction

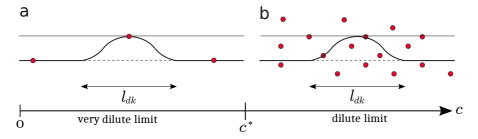


Figure 4: Schematic of the solute (red dots) distribution surrounding a double-kink nucleation structure with length l_{dk} for (a) very dilute ($c < c^*$) and (b) dilute limit ($c > c^*$) cases. When $c < c^*$ there will be at most one solute within the length l_{dk} . Whereas, multiple solutes may be found within double kink nucleation length when $c > c^*$.

 c/c^* of segments containing one solute randomly distributed among the $2N_T$ possibilities while the remaining fraction $(1-c/c^*)$ of segments are solute-free. The latter fraction have a double-kink activation barrier ΔH^0 equal to that of the pure matrix. Nucleation will be preferred in segments containing solutes having favorable interaction energies, and nucleation will be centered on the solute position. Therefore, the barrier change due to a single favorable solute at site i is

$$\Delta H_i^{\text{sol}}(\tau) = \Delta U_i g(0, \tau). \tag{16}$$

The nucleation rate is slower in segments containing solutes with unfavorable interaction energies. As an approximation, we treat these segments as having zero solutes, which slightly overestimates the nucleation rate in these segments. The total double-kink nucleation rate R_{dk} is then the sum of the rates over all segments, with or without solutes,

$$R_{dk} = \sum_{k=1}^{\left(1 - \frac{c}{c^*}\right)N} \underbrace{\nu_0 \exp\left(-\frac{\Delta H_0(\tau)}{kT}\right)}_{\text{pure matrix}} + \underbrace{\sum_{k=1}^{\left(\frac{c}{c^*}N\right)/2}}_{\text{segments containing one unfavorable solute}} \underbrace{\nu_0 \exp\left(-\frac{\Delta H_0(\tau)}{kT}\right)}_{\text{segments containing one unfavorable solute}} + \underbrace{\sum_{k=1}^{\left(\frac{c}{c^*}N\right)/2}}_{i \in N_T} \underbrace{\nu_0 \, s_i \, \exp\left(-\frac{\Delta H_0(\tau) + \Delta H_i^{\text{sol}}(\tau)}{kT}\right)}_{\text{segments containing one forwardle solute}},$$

$$(17)$$

where the summation over i extends over all N_T favorable sites and $s_i = 1$ for

site i containing a favorable solute and $s_i = 0$ for all other sites. Noting that $\langle s_i \rangle = 1/N_T$ the above equation can be reduced to

$$R_{dk} = N\nu_0 \exp\left(-\frac{\Delta H_0(\tau)}{kT}\right) \left[1 + c\frac{l_{dk}}{b} \sum_{i \in N_T} \left(\exp\left(-\frac{\Delta H_i^{\text{sol}}(\tau)}{kT}\right) - 1\right)\right]$$
(18)

Recall that N_T is the number of sites with appreciable negative interaction 303 energy changes. The above relation shows that the inclusion of more-distant sites, i.e. those having comparatively low interaction energy changes compared 305 to those within the N_T sites, will not influence the double-kink nucleation rate 306 since $\exp(-\Delta U_i g(0,\tau)/kT) - 1 \approx 0$ for such sites. Thus, the choice of the N_T 307 sites mainly affects the definition of the critical concentration c^* rather than the actual rate of nucleation in this domain. 309 The transition from the very dilute to the dilute limit is not fully developed. 310 Neither result is quite correct at concentrations $c \sim c^*$: there will often be more 311 than one solute with l_{dk} but not quite enough such that the asymptotic random 312 Gaussian statistics is applicable. However, the critical concentration c^* may be 313 below commonly-studied solute concentrations of dilute BCC binary alloys (1 314 at.\(\%). Therefore, a lack of precision in the definition of c^* and the behavior in 315 the vicinity of c^* is not a major issue.

4. Validation against atomistic simulations using a model Fe-Si alloy

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We now compare predictions of our analytical model for double-kink nucle-318 ation enthalpy barrier against direct atomistic simulations using a model Fe-Si 319 alloy. The only input parameters to the model are the solute/screw-dislocation 320 interaction energies $U(x_i, y_j)$ at all distinct solute positions ij around the screw core, the length L of the dislocation, and the stress-dependent enthalpy barrier and transition state dislocation configurations of the pure matrix material. 323 There are no fitting parameters. 324 We use a model Fe-Si alloy system as represented by a recent EAM potential 325 [2]. The Fe-Fe interactions are based on the Fe EAM potential developed by Proville et al. [19], which has all the key aspects for Fe screw dislocations

such as the non-degenerate compact core structure, the single-hump Peierls 328 potential, and {110} slip-plane. We further set the Si-Si interaction energies to those of Si-Fe, which eliminates direct solute-solute interactions. Since we are interested in dilute alloys and since the theory does not include solute-331 solute interactions, this treatment of solute-solute interactions is convenient 332 and very useful. We note clearly that this Fe-Si potential is not an accurate 333 representation of real Fe or real Fe-Si. However, it is a well-defined model system with no pathological behaviors that often plague atomistic modeling of 335 BCC screw dislocations. Thus, this model Fe-Si system is well-suited to validate 336 our general analytic model in detail. 337

The solute/screw dislocation interaction energy parameter $\Delta \tilde{E}_p(a)$ is computed using the individual solute/dislocation interaction energies shown in Figure 2b. The individual energies were computed as discussed carefully in [23]. The resulting solute/dislocation interaction energy parameter for this model Fe-Si alloy is then computed as $\Delta \tilde{E}_p(a) = 344\sqrt{c}\,\mathrm{meV}$.

To compute the transition path from the initial to the final state, we use the nudged elastic band (NEB) method as implemented in LAMMPS. To find the minimum energy path (MEP) under applied shear stress, we followed the procedure outlined in [20]. Details of the NEB simulation can be found in Appendix C. NEB calculations on the pure Fe matrix at various stresses up to the Peierls stress $\tau_p \approx 1000$ were performed. The transition state configurations, determined using the disregistry method [22], are shown in Figure 3. These configurations were fit to a functional form for $g(z, \tau)$ as

$$g(z,\tau) = \frac{1}{2} \left(\tanh \left[\frac{z/b + \zeta(\tau)}{\alpha(\tau)} \right] - \tanh \left[\frac{z/b - \zeta(\tau)}{\alpha(\tau)} \right] \right), \tag{19}$$

where $\zeta(\tau)$ and $\alpha(\tau)$ are two fitting parameters shown in Table 1. The kink-pair nucleation length was estimated as $l_{\rm dk}=20b$, consistent with the result of [2]. The initial positions $g_{\rm eq}(\tau)$ and the geometrical shape factor $G(\tau)$ are shown in Table 1. The maximum energy along each energy path is the double-kink nucleation energy barrier $\Delta H^0(\tau)$ for pure Fe as described by this potential, which was fit to the empirical form $\Delta H^0(\tau)=(0.62\,{\rm eV})(1-(\tau/1000\,{\rm MPa})^{0.62})^{1.02}$ for

later use.

Table 1: Parameters describing the transition state configurations for double-kink nucleation of pure Fe at various normalized applied stresses, τ/τ_p . The geometrical factor, $G(\tau/\tau_p)$ is obtained by Eq. (6) with $g(z,\tau)$ given by Eq. (19) and $N_s=10$.

τ/ au_p	$\alpha(\tau/\tau_p)$	$\zeta(\tau/\tau_p)$	$g_{\rm eq}(au/ au_p)$	$G(\tau/\tau_p)$
0.0	4.347	6.987	0.00	3.07
0.1	4.347	6.987	0.00	3.07
0.2	4.381	5.367	0.01	2.55
0.3	4.444	4.487	0.02	2.20
0.4	4.571	3.918	0.03	1.94
0.5	4.792	3.584	0.04	1.77
0.6	5.184	3.419	0.05	1.64
0.7	5.755	3.311	0.06	1.52
0.8	6.725	3.241	0.08	1.39
0.9	7.886	2.666	0.14	1.06
1	-	0	0.365	0

To study the (weak-link) scaling of the nucleation barrier with total dislo-350 cation length, NEB simulations are performed for lengths 40b, 70b, 100b, and 351 200b. The shortest length of 40b is the shortest length at which the double-kink 352 nucleation barrier in pure Fe becomes essentially length-independent, i.e. at 353 which there is very limited interaction between the two kinks formed during 354 kink-pair nucleation. Specifically, the pure Fe enthalpy barrier at 40b is 572 355 meV, only 28 meV smaller than the value 600 meV at 200b. NEB calculations are then performed to find the double-kink nucleation barrier in random Fe-Si dilute alloys for a range of stresses and solute concentrations of 1, 2, and 4%. At zero stress 200 different realizations of random Si are considered at each 359 concentration while 120 cases are studied at non-zero stresses.

4.1. Very dilute limit: the single solute

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NEB calculations of the double-kink nucleation process in the model Fe-Si alloy containing a single Si atom at 6 different atomic sites (numbered in Fig. 363 2f) were performed. The interaction energy changes upon glide range from -364 113 meV to -69 meV for these 6 sites. Fig. 5 shows the enthalpy barriers for 365 double-kink nucleation barrier versus the applied stress τ for each solute position. Similar studies were performed by [2], which served as the only input into their subsequent Monte Carlo studies at all concentrations. All of these fa-368 vorable sites reduce the pure double-kink nucleation barrier, with a decreasing 369 effect with increasing applied stress. Also shown is the prediction for one partic-370 ular case, site #4 with interaction energy change -89 meV. For each solute, the 37 trends follow the model predictions (not shown except for site #4) in general, 372 but the precise results do not agree exactly. The nucleation path adjusts very 373 subtly to the precise solute position, affecting the energy barrier slightly, and 374 this cannot be captured in the model. Across all these cases, the differences 375 between NEB and model are roughly +/- 20 meV, which is comparable to the differences among the 6 solute interaction energies. This result sets a level of 377 agreement that can be expected for subsequent results. 378

379 4.2. Validation of solute/double-kink interaction energies

In our model, the collective interaction energies of the solutes with the double-kink transition state (Eq. (3)) were modeled using the transition state configuration of the pure matrix at the relevant applied stress (Fig. 3) and an approximation for the solute/transition state interaction energies (Eq. (2)). Here, we test the model by comparing the predicted and NEB-computed *changes* in nucleation barrier $\Delta H^{\rm sol}(0)$ due to *specific* arrangements of solutes at zero stress.

Si solute configurations were created by placing multiple Si solutes at selected atomic sites around the dislocation core. The configurations were selected so as to sample various local energy environments (combinations of solute positions,

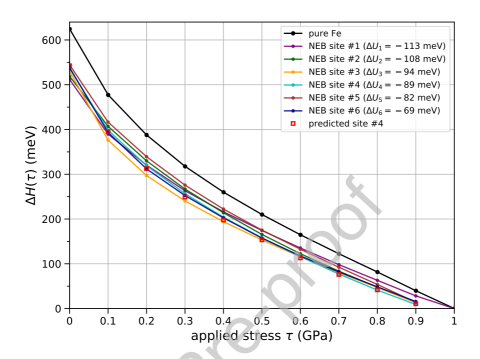


Figure 5: The double-kink nucleation enthalpy barriers, $\Delta H(\tau)$ versus the applied stresses τ for various single Si positions in Fe matrix. The solute/screw interaction energy change associated with each site, ΔU_i is also shown.

including favorable and unfavorable sites, at different distances). The configurations used are shown schematically in see Figure 6(a), with solutes labeled by the atomic row (1-8 as indicated in Figure 2e) and with an additional overbar notation for multiple solutes spaced along the same row. The spacings of solutes along the same row are indicated by either w or d, as shown in the figure. For each specific set of solute positions, we calculated the double-kink nucleation barrier change at zero stress using Eq. (3) and the transition state shape parameter G(0) = 3.07 shown in Table 1. Direct NEB calculations were performed on the exact same solute configurations at zero applied stress.

The atomistic and predicted results for the change in nucleation barrier are shown in Fig. 6(b). The trends in the simulation are well-predicted by the analytic model with no adjustable parameters. The generally good agreement

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between the predicted and simulated energy changes for most of the cases (de-402 viating by only 20-30 meV except for one extreme case discussed below) shows 403 that the use of the zero stress transition state structure to describe the collective effect of multiple solute/double-kink interaction energies is quite accurate. The 405 maximum deviation in Fig. 6(b) is for the case $1 + \overline{1} + 5 + \overline{\overline{5}}, d = 3b$. However, 406 this configuration consists of solutes in four energetically-favorable positions, 407 two of which are in the most favorable positions, and no other solutes. This is thus an extreme case - all the solutes are strongly attracting the double-kink 409 - that is very unlikely to occur in the dilute alloys. Evidently, it distorts the 410 transition state shape away from the assumed shape sufficiently to reduce the 411 effect of the solutes. Overall, however, while the precise double-kink transition 412 state structure in the NEB varies with each distinct solute configuration, with 413 solutes pulling/pushing the dislocation line away from the pure Fe shape, our 414 model remains quite good, and so is a quantitative parameter-free analytical 415 model. 416

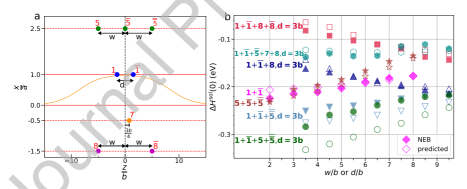


Figure 6: (a) Definition of solute positions relative to a double-kink nucleation shape (orange solid line). Same color and number is used for solutes in a same atomic row position (x_i, y_j) consistent with sites marked in Fig. 2f. (b) The zero stress double-kink nucleation enthalpy barrier changes $\Delta H^{\rm sol}(0)$ due to different solute combinations. The numbers indicate the numbered solutes in (a); e.g. $1 + \overline{1} + 8, d = 3b$ denotes solutes at sites #1 and # $\overline{1}$ at fixed distance d = 3b plus a solute at site #8 at various w distances.

4.3. Double-kink nucleation in model dilute Fe-Si random alloys

Finally, we compare our predictions of the double-kink nucleation barrier for the model binary random Fe-Si dilute alloys to direct NEB simulations for a range of solute concentrations and dislocation lengths and under zero and non-zero stresses. The NEB selects the lowest barrier in any particular realization, and so our predictions use the weakest-link limit (Eq. 10) appropriate for the NEB simulations.

424 4.3.1. zero stress

Figures 7(a)-(d) show the mean, statistical distribution, and standard deviation of the NEB double-kink nucleation barrier ΔH as a function of solute concentration for dislocation lengths L=40b,70b,100b, and 200b, respectively, at zero stress. The corresponding predictions of the theory for the mean and standard deviation, with no adjustable parameters, are also shown. As predicted, the changes in nucleation energy barrier are statistically-distributed in random dilute alloys and are a function of both solute concentration and dislocation length.

At the shortest length of 40b, there is essentially no weak-link scaling effect. The distribution of barriers is centered on the single-solute result, not the zero-solute result, because the nucleation will typically occur around the single most-favorable solute. The standard deviation, which is representative of the statistical variations from sample to sample, increases as the concentration increases. The theory predicts both the mean and standard deviation in good agreement with the simulations, with no adjustable parameters. The results at 40b represent the "unit" statistical process, with weak-link length effects entering for longer lengths.

For the longer lengths 70b - 200b, the mean barrier decreases because the system can find the weakest among several possible nucleation environments along the dislocation line, as predicted by the theory. The standard deviation also narrows slowly, as predicted. All the trends predicted by the theory agree quantitatively with the simulations. At length 200b, essentially none of the sim-

ulations at 1, 2 or 4% has a barrier that exceeds that of pure Fe. The system always finds some favorable solute fluctuations that reduce the double-kink nucleation barrier, i.e. "softening" the alloy, even at this rather small dislocation length. Note that these results are for double-kink nucleation only; kink migration is not investigated even though it may become the rate-controlling factor at the higher concentrations.

4.3.2. Non-zero stress

The predictive model is further verified by comparing to NEB simulations under applied stresses in the range 100-900 MPa. Simulations were performed for a single dislocation length of 70b and concentration of 2%. Figure 8 shows the mean enthalpy barrier as simulated and as predicted, along with the enthalpy barrier for pure Fe. The statistical distribution of barriers as simulated is shown as a histogram and is important because nucleation in realistic systems will occur along lengths much larger than 70b.

Very good agreement is found between the predicted and NEB results. The 461 barrier reduces as the stress increases, but the difference between pure Fe and 462 the alloy decreases with increasing stress. At a stress of 70% of the Peierls stress, 463 the spectrum of barriers extends to zero (and below). That is, for some cases, the initial dislocation position is unstable (zero barrier) against gliding forward into the lower-energy state in the next adjacent Peierls valley. These cases are 466 not shown and the predictions for the mean barrier at the higher stresses must 467 use a lower limit of $-\Delta H^0(\tau)$ in the integrals in Eq. (10). More importantly, for longer lengths, the mean barrier at any stress will decrease even further, with the standard deviation decreasing only slowly, enabling zero-energy barriers to 470 occur with increasing frequency. 471

The results in Figure 8 fully validate our analytic model for double-kink nucleation in random alloys as a function of both concentration and stress. For any length dislocation beyond more than a few segments, and at all stresses below the Peierls stress, the presence of solutes reduces the double-kink nucleation barrier, i.e. solutes always soften double-kink nucleation. This validation is

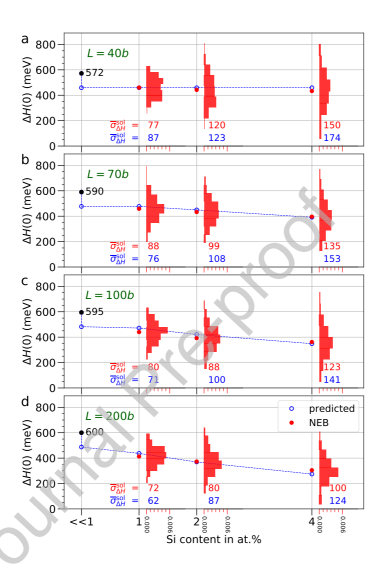


Figure 7: Zero stress double-kink nucleation barrier $\Delta H(0)$ versus solute concentration c for dislocations with length L (a) 40b, (b) 70b, (c) 100b, and (d) 200b. The histograms show the distribution of double-kink nucleation barriers obtained by NEB calculations. The black numbers are the enthalpy barriers for double-kink nucleation in no solute pure Fe. The red and blue numbers are the standard deviations of the distributions of double-kink nucleation barrier obtained by NEB simulations and theory, respectively.

performed on a model Fe-Si alloy where all material parameters are known, but

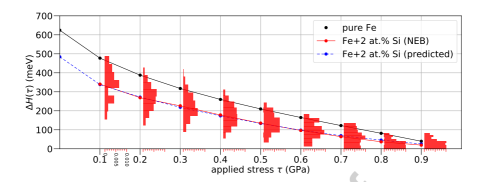


Figure 8: The double-kink nucleation barrier $\Delta H(\tau)$ versus applied stress τ for dislocations with length L=70b in Fe-2at%Si dilute alloys. The histograms are the distribution of double-kink nucleation barriers obtained by NEB atomistic calculations.

now enables application of the analytic model to be applied to realistic alloys using input material parameters relevant for the real alloys.

480 5. Applications and implications

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The agreement between theory and simulation shown in the previous section
on a model random dilute Fe-Si alloy validates all of the major features of the
theory, and demonstrates the quantitative accuracy of the approximations that
render the theory analytic. The theory can thus be applied with confidence to
any bcc alloy system in the dilute limit. Challenges in applying the theory lie in
obtaining accurate input data and also accurate experimental results. Nonetheless, below we achieve some insights in making comparisons to experiments at
a level of detail well beyond previous literature.

We first address how the theory can be used to estimate the yield strength of a dilute alloy. The material parameters in the theory are reiterated: (i) the double-kink nucleation barrier and transition state configurations of the pure matrix; (ii) the double-kink nucleation length and total dislocation length; and (iii) the solute/screw dislocation interaction energies. The interaction energies can in principle be computed using first-principles (e.g. Figures 2(a) and (c)), although this is computationally challenging and still not necessarily precise for

the real material. The double-kink nucleation length can be estimated reasonably by various means [20, 18]. The stress-dependent double-kink nucleation barrier for the pure matrix is derived from experiments of the flow stress versus temperature as explained below.

Obtaining $\Delta H^0(\tau)$ from experiments requires accounting for the number of nucleation sites N along each dislocation segment of length $L=1/\sqrt{\rho}$ between dislocation junctions. When deformed at a constant strain rate $\dot{\epsilon}$, the thermally-activated Arrhenius model plus the Orowan model leads to a strain rate $\dot{\epsilon} = N\rho ba\nu_0 \exp(-\Delta H^0(\tau)/kT)$ where $N \sim 1/(l_{dk}\sqrt{\rho})$ is the number of possible nucleation sites along dislocations in between dislocation junctions. A reference strain rate $\dot{\epsilon}_0$ often replaces the combination $\rho ba\nu_0$. The enthalpy barrier at stress τ is then obtained as

$$\Delta H^0(\tau) = \Delta H^0_{\text{exp}}(\tau) + k T \log(N). \tag{20}$$

where $\Delta H^0_{\rm exp}(\tau) = k T \log(\dot{\epsilon}_0/\dot{\epsilon})$ is the experimental enthalpy barrier obtained from the experimental flow stress versus temperature experimental data.

With the above details, softening by solutes is then determined as shown schematically in Fig. 9. Starting with the experimental enthalpy barrier versus stress, the correction is made for N, and then the contribution of the solutes is added, where the figure shows the case where the low-temperature weak-link result is applicable. An experiment carried out at a particular $\Delta H^{\rm exp}=kT\log(\dot{\epsilon}_0/\dot{\epsilon})$ determined by the specified temperature and strain-rate will then measure the strengths indicated in Fig. 9.

While the weak-link analysis can only be applied for low temperatures $T < T_0$, the total nucleation rate due to all nucleation sites, Eq. (9), can be applied for the whole temperature range of interest. In this case, considering Eq. (20) along with Orowan's law, Eq. (9) is solved numerically to determine the yield stress at a given strain rate and temperature. In the following, the full rate theory has been used to predict the yield stress of real dilute Fe-Si, W-Re, and W-Ta alloys.

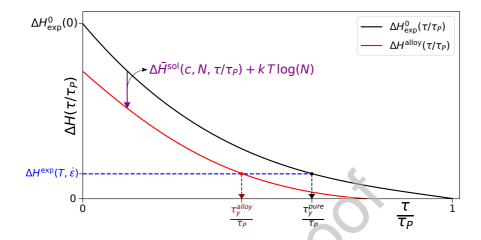


Figure 9: Generic prediction of solute softening by double-kink nucleation starting from double-kink nucleation in the pure metal. The experimental barrier versus stress must first be corrected for the number of nucleation segments N, and then the additional effect of softening is added to obtain the total activation enthalpy. At a given experimentally-imposed activation enthalpy determined by the temperature and strain rate, the alloy has a lower yield stress than the pure metal. The example here uses the weak-link limit for illustration.

5.1. Application to Fe-Si alloys

We now use the theory to make specific predictions for real dilute Fe-Si (0.52 at%, 1.3 at%, 3.2 at%, and 5 at%) alloys over a wide temperature range and compare to the experimental data of [24] performed at a strain rate of $1.7 \times 10^{-4} \, \mathrm{s}^{-1}$. We use a reference strain rate of $10^4 \, \mathrm{s}^{-1}$ and take a dislocation density of $\rho = 10^{12} \mathrm{m}^2$ ($L \sim 1 \mu \mathrm{m}$; $N \sim 170$) based on the TEM measurements of Takeuchi [25].

The same Fe-Si alloys at 5 at% and higher were previously studied by [16] using the non-dilute theory. There is significant strengthening at this concentration, indicating that the non-dilute theory is appropriate. Good agreement in strength versus temperature at these higher concentrations was obtained using a solute/dislocation interaction energy parameter $\Delta \tilde{E}_p = 178\sqrt{c}$ meV, and so we use this value here. The transition state configurations for pure Fe are taken from a study using a machine-learning interatomic potential trained on a very

large DFT dataset [20] with double-kink nucleation length $l_{dk} = 12 \, b$. Both the interaction energy parameter and the double-kink transition state structures differ from those of the model Fe-Si EAM potential precisely because that potential is not quantitative for real Fe-Si. Our predictions are shown in Fig. 10. For 0.52at%Si at low T, the experiments show a very small hardening while the theory predicts a very small

amount of softening. At higher T> 200K, the prediction and experiment are similar, and deviations from pure Fe are quite small. There is a small hump in 537 the data around 150K, widely discussed but unexplained to date [26, 27, 28]. 538 For the 1.3at%Si alloy, there is again a very small strengthening seen at low T 530 and minimal effects at higher T. The predictions show a small softening at low T and then little net effect at higher T. For 3.2at%Si, at low T there is slight softening and then a small but distinct strengthening at higher T> 100K. The 542 theory shows a stronger softening that remains roughly constant relative to pure 543 Fe over the entire temperature range. At 3.2at%Si, the double-kink nucleation theory deviates significantly from experiments throughout the entire temperature regime, suggesting another mechanism controlling deformation (see below). Further significant deviations are obtained at an even higher concentration of 547 5at%Si, where the alloy is significantly strengthened but the double-kink nucle-548 ation theory predicts a large softening, relative to pure Fe, again indicative of 549 another controlling mechanism (see below). The observed strengthening at low T for 0.52at%Si and 1.3at%Si is small and not explained by the present model. 551 However, we show below another set of data at 1.0at%Si that shows distinct 552 softening relative to pure Fe. Thus, the theory is qualitatively consistent with 553 experiments at low concentrations in showing little effect of the solutes while the experiments themselves are in some conflict.

The failure of the present double-kink nucleation theory in comparison to the experiments at 3.2at%Si and higher concentrations is attributed to a transition from nucleation controlled softening at lower concentrations to some strengthening mechanisms at the higher concentrations. The traditional view is a transition to strengthening controlled by kink migration [6, 1, 2, 3]. Here we apply

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the non-dilute yield strength model of Maresca et al. [16] that has successfully captured strengths in a number of bcc alloys. In this theory, an initially-straight screw dislocation in a random alloy lowers its energy by spontaneously forming a multiply-kinked structure at zero stress. The strength is then controlled by a combination of kink glide and cross-kink (also called jog or dipole) pinning, with a Peierls-like mechanism replacing kink glide at very low T. Crucially, there is no role of thermally-activated nucleation of double-kinks in this model, and so it is applicable only in the non-dilute regime.

The transition between dilute (double-kink nucleation controlled softening) 569 and non-dilute (kink glide and cross-kink controlled strengthening) is then 570 mainly dictated by the spacing of the spontaneous kinks. If this length is smaller than the distance L between forest junctions, then the Maresca et al. model 572 should apply and the alloy is in the non-dilute regime. If the kink spacing 573 is predicted to be larger than L, then the spontaneous kinked structure can-574 not actually form, and thermally-activated double-kink nucleation is necessary; 575 such alloys are in the dilute regime where the present theory should apply. Maresca et al. predict the spontaneous energy-minimizing kink-spacing to be 577 $2.5\,\zeta_c$ where $\zeta_c=(1.083E_k/\Delta \tilde{E}_p)^2 b$ is a critical length scale that depends only 578 on the ratio between the kink formation energy $(E_k = 0.5 \Delta H^0(0))$ and the 579 solute/dislocation energy quantity $\Delta \tilde{E}_p$. The length scale 2.5 ζ_c that determines 580 the boundary between dilute and non-dilute domains thus depends on precisely the two major material quantities that enter into the present double-kink nu-582 cleation theory $(\Delta H^0(0), \Delta \tilde{E}_p)$. This enables direct comparisons of the two 583 theories with no additional material parameters. 584

The legend in Figure 10 shows the predictions of Maresca et al. for the length scale $2.5 \zeta_c$ at each Si concentration. At both 0.52at%Si and 1.3at%Si, the length scale $2.5 \zeta_c$ is larger than or comparable to L. According to our criterion, these alloys are thus expected to be in the "dilute" regime where double-kink nucleation controls the strengthening. This conclusion is supported by the reasonable agreement of the strength predictions using the double-kink nucleation theory and the poor agreement of the strength predictions (too low) of the Maresca

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et al. theory as shown in Figure 10. In contrast, at 3.2at%Si, the length scale $2.5 \zeta_c$ is definitely smaller than L and, correspondingly, the strength predictions of Maresca et al. are in fairly good agreement with experiments, aside from a small over-prediction at low T < 75K. At this concentration, the double-kink nucleation prediction is too low at all temperatures. The experiments at 54% with $2.5 \zeta_c \ll L$ are then, as expected, captured well by the Maresca et al. model while the double-kink nucleation model predictions are poor. This analysis supports a transition from the dilute softening regime to the non-dilute strengthening regime between 1.3at% and 3.2at%.

The transition here is based on temperature-independent length scales. The
experiments at 3.2at%Si suggest that there can be both softening at low T and
strengthening at higher T. The trends in strength versus temperature of the
two different models also suggests that at concentrations in between 1.3at%
and 3.2at% there could changes in the strongest mechanism vs. temperature.
Such details are well beyond the scope of the present models.

Our predictions for Fe-Si can be further compared to experiments of Chen et al. [30] who reported the yield stress of dilute Fe-1at%Si and Fe-4at%Si 608 polycrystals. Chen et al. converted the uniaxial polycrystalline yield stresses 609 to critical resolved shear stresses (CRSS) by using the Taylor factor ~ 2.75 for 610 bcc pencil glide. The resulting CRSS for pure Fe was then approximately 20%611 higher than that of Ref. [24], indicating the possibility that the data of Ref. [24] on pure Fe is low (for unknown reasons) which then brings into question the 613 apparent strengthening at 0.52at%Si relative to the pure Fe data of Kitajima 614 More importantly, Chen et al. report distinct softening in Fe-1at%Si 615 relative to pure Fe, in contrast to the results of Ref. [24] at 1.3at%Si. These 616 comparisons demonstrate uncertainty in the pure and dilute Fe-Si data, which 617 then leads to uncertainty in assessing softening versus strengthening. 618

To apply our analysis to the data of Chen et al., we must first shift the results of Chen et al. because our analysis was based on the pure Fe of Ref. [24] as a reference. To do so, we simply rescale all the data of Chen et al. by a single factor of 0.8 to best-match the pure Fe experimental data of Kitajima et al. We

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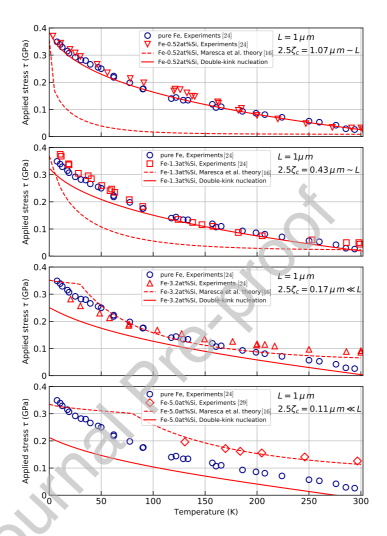


Figure 10: Theory predictions for $Fe_{1-x}Si_x$ alloys.

then make predictions for Fe-1at%Si as shown in Fig. 11. For 1at%Si, both experiments and theory show slight softening up to T=225K. The experiments then show a slight strengthening at T=300K that is not captured by the theory. The length scale $2.5\zeta_c$ and the strengthening predicted by Maresca et al. are also shown in Figure 11. At 1at%Si, the predicted strengthening is well below the experiments at all temperatures and the length scale $2.5\zeta_c$ is comparable to

L. These results indicate that this alloy is in the 'dilute' regime where double-629 kink nucleation controls the screw dislocation motion. This is consistent with our conclusions based on the data of Kitajima et al. Similar predictions for Fe-4at%Si are shown in Fig. 11. At T = 77K, the 632 experiments show a notable softening followed by a transition to strengthening 633 above 200K. The length scale $2.5\zeta_c$ of Maresca et al. indicates this alloy should be in the non-dilute regime. In addition, the non-dilute theory captures the strengthening reasonably well at the higher temperatures. The double-kink 636 nucleation theory is in good agreement with the experiments at low T, which is 637 surprising since we would expect this alloy to be in the non-dilute regime. This 638 suggests some temperature dependence to the dilute/non-dilute regime that is currently beyond the scope of the present analysis. In spite of uncertainties in the experimental data, our analysis of the classic 641 Fe-Si alloy provides a solid conceptual understanding of the nature of the tran-642 sition in strengthening with increasing concentration. At low concentrations, there is generally softening by double-kink nucleation. At high concentrations, double-kinks are pre-existing and strength is controlled by kink migration and cross-kink formation. The transition is formulated in terms of length scales 646 only, however. So, at concentrations in the transition range between 1.3 and 4 647 at%Si, strengthenging may be a complex and temperature-dependent interplay 648 between the two mechanisms. Such an interplay could be revealed by detailed kinetic Monte Carlo studies on long dislocation lengths that account carefully 650 for the origins of kinks (pre-existing versus nucleating) and using the full sta-651 tistical analyses for solute/screw interactions developed here. Existing kMC 652 methods have already shown a transition generically, but have made simplifying assumptions about the solute/screw interactions, the kink formation, and the initial structure of the dislocation [31, 2, 3], that can perhaps be revisited. 655 5.2. Application to W-Ta and W-Re alloys 656

We now examine the yield stress for W-Re and W-Ta alloys at 0 (pure W), 1, and 3 at%, and temperatures $T = 150 \, K$, $T = 300 \, K$, and $T = 590 \, K$. Ex-

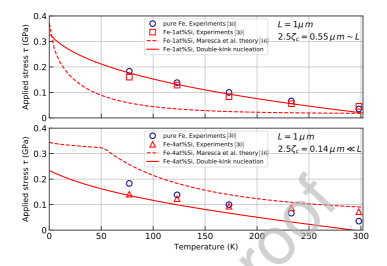


Figure 11: Theory predictions for $Fe_{1-x}Si_x$ alloys.

perimental data was reported by Stephens[7] at a strain rate of $5.5 \times 10^{-4} \,\mathrm{s}^{-1}$. 659 We use the pure W data to estimate the pure W enthalpy barrier versus stress 660 assuming the reference strain rate $\epsilon_0 = 10^4 \, s^{-1}$ and fitting a Kocks law as 663 with $\Delta H_{\rm exp}^0(0) = 1.4$ eV and $\tau_P =$ $\Delta H_{\rm exp}^0(\tau) = \Delta H_{\rm exp}^0(0) \left(1 - \left(\frac{\tau}{\tau_P}\right)\right)$ 662 1.1 GPa. We use the transition state configurations obtained by GAP for pure 663 Fe, since estimates for W are similar [18]. We use the solute/dislocation interaction energies obtained by DFT (Fig. 2) from which we compute the so-665 lute/dislocation interaction energy parameters as $\Delta \tilde{E}_p = 345\sqrt{c}$ meV for W-Re 666 and $\Delta \hat{E}_p = 137\sqrt{c}$ meV for W-Ta. We again use a mobile dislocation density 667 of $\rho = 10^{12} \text{ m}^{-2}$ corresponding to $L = 1 \mu \text{m}$ and $N \sim 150$ segments along the length L when considering the transition to the non-dilute limit. We note clearly that, given the limited data to fit H_{exp}^0 and the uncertainty of other parameters, 670 our predictions below are mainly qualitative and illustrative. 671 Predictions of the strength using the double-kink nucleation theory for the 672 W-Re alloys are shown in Fig. 12 (a). The predicted softening for W-1at%Re is in reasonable agreement with the experiments at all three temperatures. For W-3at%Re, the predicted softening increases, as expected, while the experi-

ments at 150K and 500K show a slight strengthening relative to 1at%Re and 676 the experiments at 300K exhibit continued softening. This suggests a mechanism transition between 1at%Re and 3at%Re. We thus evaluate the length scale $2.5\zeta_c$ of the Maresca model to assess a possible transition 3at%Re. Us-679 ing the kink energy $E_k = 0.905$ eV from the DFT-based GAP potential for W 680 [32], we predicted $2.5\zeta_c = 0.55 \,\mu m$ for W-1at%Re. This is comparable to the 683 (approximate) dislocation length L, indicating that W-1at%Re is in the dilute regime. In contrast, we predict $2.5\zeta_c = 0.18 \,\mu m$ at 3at%Re, which is rather 683 smaller than the (approximate) dislocation length L, suggesting that 3at%Re is 684 in the non-dilute regime. The underpredictions of strength by the double-kink nucleation model at 3at%Re are thus qualitatively understood as being due to a transition to the non-dilute limit. The Maresca et al. model does not predicted a temperature-dependence of $2.5\zeta_c$ and so the observed softening at T=300K in 688 W-3at%Re requires a full application of the Maresca et al. model before further 689 conclusions can be made.

Both trends and uncertainities in the experiments on W-Re are revealed by examining the data of Raffo [33] on very clean polycrystals at 0, 1, and 2at% 692 at temperatures of 77K, 298K, and 589K. Data at 77K (not shown) shows a 693 steady softening, although very small at 1at%; there is no comparable data by 694 Stephens. At T=298K, as shown in Figure 12a scaled by the Taylor factor 2.71, Raffo shows a small strengthening at 1at%Re followed by softening at 2at% (but still stronger than pure W). At T=589K, as shown in Figure 12a, there is 697 softening at 1at% and strengthening at 2at%. If the single data point for pure 698 W at T=298K were higher, the Raffo data would be qualitatively similar to the 699 data of Stephens, and generally consistent with our analysis. Raffo also showed data at 7at% indicating continued softening at 77K but continued strengthening 70 at 298K and 589K, the latter two results consistent with our analysis. 702

Predictions of the strength using the double-kink nucleation theory for the W-Ta alloys are shown in Fig. 12 (b). The experiments on W-Ta show a softening at 1at%Ta for all three temperatures studied. The double-kink nucleation theory predicts only a very slight softening for the 1at%Ta alloys, and so agrees

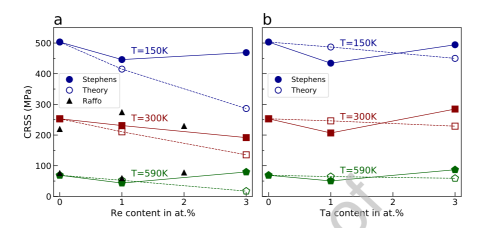


Figure 12: Critical resolved shear stress (CRSS) versus solute concentration at various temperatures as indicated for (a) W-Re (filled symbols: data from Stephens [7]; open symbols: present theory; black triangles: data from Raffo [33]); (b) W-Ta (filled symbols: data from Stephens [7]; open symbols: present theory) dilute alloys.

only qualitatively with the experiments. The experiments at 3at%Ta show a slight strengthening at all three temperatures. The double-kink nucleation theory predicts continued softening, suggesting a transition in mechanism between 1 and 3at%Ta. However, the non-dilute theory of Maresca et al. predicts a length scale of $2.5\zeta_c = 1.1\,\mu m$ at 3at%Ta that remains comparable to the (assumed) L and so does not quite indicate a transition at this composition. A transition would be expected only at slightly higher Ta content.

Overall, softening at 1at% is predicted and observed for both Re and Ta solutes, supporting the general concept that softening arises for all solutes. A transition to hardening at 3at% is captured by our analysis for Re but not for Ta. These findings remain tentative on both theoretical and experimental sides. The material parameters needed for the theory are estimated to the best of current ability but remain uncertain. In addition, it is not expected that there is a distinct transition from softening to hardening at some very specific concentration but rather a delicate and possibly temperature-dependent interplay of mechanisms. Identifying a precise transition is also not of practical

importance - what is important is that the operative physical mechanisms be identified and quantified so that alloy design can proceed on the basis of these mechanisms. Even with these significant caveats, the two theories together (present theory and Maresca et al. theory) remain broadly consistent with the experiments and so provide the desired framework for alloy design.

We conclude with some remarks about previous modeling and simulation on the W-Re and W-Ta alloys. In the Introduction, we already discussed theoretical/conceptual issues with the model of Hu et al., and now focus on predic-730 tions. The model of [8] was applied to W-Re but was compared to hardness 731 data at T=298K. Hardness and yield strength are not directly related, unfor-732 tunately, because hardness depends on the work hardening rate as well as the yield strength. This is clearly seen in the differences in flow stress versus solute type as a function of plastic strain in the uniaxial data of Stephens. Conclusions 735 about relative softening and hardening based on experiments at a few percent 736 plastic strain are thus inconsistent with the actual data at the onset of yielding. 737 Hu et al. nonetheless converted the hardness data of Stephens and Witke at 298K [34] to a CRSS, leading to a CRSS for pure W of \sim 350 MPa significantly 739 higher then the ~ 250 MPa reported by Stephens in direct uniaxial single crystal 740 experiments. Hence, quantitatively, the predictions of Hu et al. do not agree 741 with the uniaxial data of either Stephens or Raffo. The trends in converted 742 hardness data do, however, suggest softening up to 5at%Re at T=298K while the uniaxial data of both Stephens and Raffo also indicate softening up to at least 2-3at%Re at T=298K. The uniaxial data, however, shows hardening at 745 3at%Re at both 150K and 590K, suggesting an underlying transition around 746 3at%Re that might not be predicted by the Hu et al. model. Zhao et al. [3] 747 compared results from Monte Carlo studies to the data of Stephens at 2% plastic strain rather than to the data reported at yield, and these results differ. 749 Zhao et al. also calibrated the pure W parameters in the Monte Carlo model to 750 DFT results that are well-established to greatly overestimate the yield strength. 751 Thus, the quantitative results of Zhao et al. are far higher than the experiments at both yield and 2% plastic strain. Examining trends only, Zhao et al. predict 753

slight softening relative to pure W up to 5at%Re at 150K, in conflict with the
experimental hardening at 3at%Re. They also predict essentially no softening
at T=300K up to 1.5at%Re and very slight strengthening up to 5at%Re, again
in conflict with the distinct softening observed experimentally up to 3at%Re. At
T=590K, Zhao et al. show a trend in good agreement with experiments. Overall, the results of Zhao et al. are quantitatively far from experiments and the
trends are inconsistent with experiments. Finally, as noted in the Introduction,
the model of Hu et al. was also applied to W-Ta and predicted strengthening,
inconsistent with the softening data of Stephens.

⁷⁶³ 5.3. Implications for spontaneous kinking in the non-dilute regime

The present theory is developed for understanding the softening in dilute 764 alloys where double-kink nucleation controls the strengthening. However, the 765 analysis of the effects of solutes on the double-kink nucleation barrier remains 766 applicable to non-dilute alloys even though strength is not controlled by this 767 mechanism. Recall that, in the non-dilute screw strengthening theory of Maresca et al., the initial straight dislocation in the random alloy at zero stress is sponta-769 neously kinked at a characteristic length of $2.5\zeta_c$. Envisioning that the system is 770 somehow prepared with initially-straight dislocations, the present analysis is ap-771 plied to examine what barriers must be overcome to achieve the spontaneously-772 kinked length. This was not addressed in Maresca et al., where it was assumed that there is sufficient time to overcome such barriers in any real systems. Here, 774 we can use the present analysis to examine the typical reduction in double kink 775 nucleation barrier due to solutes over the characteristic length of ζ_c . The reduc-776 tion in barrier then corresponds to a reduction in the average time required for 777 the dislocation to attain the fully-kinked configuration envisioned by Maresca 778 et al. 779

The relevant solute/screw interaction energy is a generalization of the dilute limit of Eq. (7) in which all elements of the alloy are considered to be solutes within an effective or average alloy matrix (see Maresca et al. and Ref. [23] for details). The interaction parameter is then $\Delta \tilde{E}_p = \left(\sum_{i,j} \sum_n c_n \Delta U_{ij}^{n\,2}\right)^{1/2}$

where the ΔU_{ij}^n is the energy change for a solute of type-n at initial position ij784 relative to a screw dislocation in the effective or average alloy material. This interaction parameter has been estimated for various HEAs in Maresca et al. and has been examined for a wide range of alloys described by interatomic 787 potentials by Ghafarollahi et al. [23]. Below, we use the values previously 788 reported by Maresca et al. as shown in Table 2. For simplicity, we assume a single double-kink nucleation length $l_{dk}=20\,b$ for all of these non-dilute complex alloys. Based on the number of possible 791 nucleation segments over each characteristic length $N = \zeta_c/l_{dk}$, the energy 792 reduction at zero stress due to the solutes can be then estimated from Eqs. 793 (10) and (11) as $|\Delta \bar{H}^{\text{sol}}(\sigma_{\Delta H^{\text{sol}}}, N)| + \bar{\sigma}_{\Delta H^{\text{sol}}}(\sigma_{\Delta H^{\text{sol}}}, N)$, where the standard deviation of the solute energy changes $\bar{\sigma}_{\Delta H^{\mathrm{sol}}}$ has been also taken into account 795 and $\sigma_{\Delta H^{\rm sol}} = G(0)\Delta \tilde{E}_p$ is the zero stress standard deviation. For each of the 796 non-dilute complex alloys considered, the energy reduction $\left|\Delta \bar{H}^{\mathrm{sol}}\right| + \bar{\sigma}_{\Delta H^{\mathrm{sol}}}$ 797 normalized with respect to $\sigma_{\Delta H^{\rm sol}}$ has been computed as shown in Table 2. The time required for the dislocation to kink is the inverse of the nucleation rate t = 1/R, where R is given by Eq. (15). We consider room temperature 800 $T = 300 \, K$, an attempt frequency $\nu_0 = 10^{12} \mathrm{s}^{-1}$, and a zero-stress geometrical 801 factor G(0) = 2.8. 802 Fig. (13) shows the double-kink nucleation enthalpy barriers and the time 803 required to move a kink over the length ζ_c at room temperature for various alloys. There is a considerable reduction in the zero stress double-kink nucleation 805 barrier at room temperature for all the non-dilute complex alloys. This leads to 806 a huge reduction in time required for the kink to overcome the nucleation bar-807 rier. For instance, the NbMoTaW high-entropy alloy has the largest estimated double-kink nucleation barrier among all these alloys and the solute fluctuations reduce the barrier by ~ 0.4 eV, which decreases the nucleation time from years 810 to less than 2 hours (note however that Maresca et al. contend that the strength 811 of this alloy is controlled by edge dislocations [35]). The maximum nucleation 812 time across all alloys is 7 hours for the dilute alloy Nb₉₅Mo₅. 813 The present analysis thus shows that attaining the low-energy kinked screw

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Table 2: The kink formation energy E_k , solute/dislocation interaction energy parameter $\Delta \tilde{E}_p$, characteristic length ζ_c , and the typical energy reduction due to solute fluctuations ($|\Delta \tilde{H}^{\rm sol}| + \bar{\sigma}_{\Delta H^{\rm sol}}|$) at characteristic length $\zeta_c = N \, ld_k$, for various non-dilute complex alloys.

Alloy	$E_k \text{ (eV)}$	$\Delta \tilde{E}_p \; (\text{meV})$	ζ_c/b	$\left(\left \Delta \bar{H}^{\mathrm{sol}}\right + \bar{\sigma}_{\Delta H^{\mathrm{sol}}}\right) / \sigma_{\Delta H^{\mathrm{sol}}}$
$\mathrm{Nb_{95}Mo_{5}}$	0.634	48.8	198	2.13
$\mathrm{Nb_{75}Mo_{25}}$	0.611	84.7	61	1.59
Fe _{91.7} Si _{8.3}	0.493	51.4	108	1.83
NbMoTaW	0.6695	86.2	71	1.66
VNbMoTaW	0.6214	90.6	55	1.59

structure envisioned by Maresca et al. is facilitated by the reduction of the double-kink nucleation barrier in a complex alloy over the crucial length scale ζ_c . However, at very low temperatures, the time to overcome these barriers remains very large and this may inhibit the transition to the non-dilute low-energy configuration in the Maresca et al. model. This might explain some differences between experiments and the non-dilute theory at T=77K and T=150K at higher solute concentrations where the non-dilute theory is expected to be applicable.

6. Summary

We have developed an analytical theory for the stress-dependent double-kink 824 nucleation barrier in dilute random substitutional BCC alloys as a function of 825 solute concentration and dislocation length, including the range of practical in-826 terest ($\sim 10^3 \, b$) where direct atomistic calculations are infeasible. The material parameters in the theory are only (i) the solute/screw dislocation interaction 828 energies, (ii) the transition state configurations of the pure matrix, and (iii) 829 the double-kink enthalpy barrier versus stress for the pure matrix. The theory 830 has been extensively validated on model random Fe-Si alloys with no adjustable 831 parameters. Comparisons to experiment on Fe-Si, W-Re, and W-Ta, have been made and reasonable agreement is obtained at low solute concentrations where

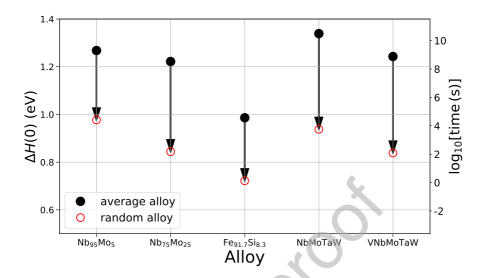


Figure 13: Double-kink nucleation enthalpy barrier and the time required to move a kink over the length ζ_c at room temperature for various non-dilute complex alloys.

double-kink nucleation is expected to control the strength. We further rationalize a transition from softening to hardening with increasing solute concen-835 tration as a change in controlling mechanism from double-kink nucleation to 836 a spontaneously kinked structure with strength controlled by kink migration 837 and cross-kink/jog pinning via the model of Maresca et al. This transition 838 involves the same fundamental material parameters in both regimes, and identifies a critical dislocation length scale as determining whether the alloy is in the dilute or non-dilute limit. The present double-kink nucleation model plus 841 the non-dilute model of Maresca et al. together thus provide a single coherent 842 analytic framework within which the initial yield strength in random BCC substitutional alloys can be quantitatively understood when strength is controlled by screw dislocations. 845

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Appendix A. Standard deviation of the potential energy change

In this appendix, we derive the standard deviation of the potential energy change $\sigma_{\Delta U_{\rm tot}}$ when a dislocation segment nucleates over a distance l_{dk} in a random field of solutes, in a dilute binary alloy. The stress-dependent total change in the nucleation barrier due to the nucleation of dislocation is

$$\Delta H^{\text{sol}}(\tau) = \sum_{i,j,k} s_{ijk} \Delta U_{ij}(a) g(z_k, \tau), \tag{A.1}$$

where the site occupation variable $s_{ijk}=1$ if a solute is at position (x_i,y_j,z_k) and 0 otherwise. Positions (x_i,y_j,z_k) refer to the atomis sites along one dislocation segment of length l_{dk} . Let $N_s=l_{dk}/b+1$ denote the number of atomic sites along z direction at each atomic row (x_i,y_j) . Also, let label the atoms in each atomic row (x_i,y_j) by $k=1\ldots N_s$.

The quantity of interest in the nucleation model is the standard deviation of the total energy change as defined in Eq. (4) which requires $\left\langle \Delta H^{\rm sol} \right\rangle^2$ and $\left\langle \Delta H^{\rm sol} \right\rangle$. We can write

$$\left\langle \Delta H^{\text{sol}} \right\rangle^{2} = \sum_{i,j} \sum_{k} \left\langle s_{ijk} \right\rangle^{2} \Delta U_{ij}^{2} g(z_{k}, \tau)^{2}$$

$$+ \sum_{i,j} \sum_{k,l \neq k} \left\langle s_{ijk} \right\rangle \left\langle s_{ijl} \right\rangle \Delta U_{ij}^{2} g(z_{k}, \tau) g(z_{l}, \tau)$$

$$+ \sum_{\substack{i,j\\m,n \neq i,j}} \sum_{k,l} \left\langle s_{ijk} \right\rangle \left\langle s_{mnl} \right\rangle \Delta U_{ij} g(z_{k}, \tau) \Delta U_{mn} g(z_{l}, \tau)$$
(A.2)

$$\left\langle \Delta H^{\text{sol 2}} \right\rangle = \sum_{i,j} \sum_{k} \left\langle s_{ijk}^2 \right\rangle \Delta U_{ij}^2 g(z_k, \tau)^2$$

$$+ \sum_{i,j} \sum_{k,l \neq k} \left\langle s_{ijk} s_{ijl} \right\rangle \Delta U_{ij}^2 g(z_k, \tau) g(z_l, \tau)$$

$$+ \sum_{\substack{i,j \\ m,n \neq i,j}} \sum_{k,l} \left\langle s_{ijk} \right\rangle \left\langle s_{mnl} \right\rangle \Delta U_{ij} g(z_k, \tau) \Delta U_{mn} g(z_l, \tau)$$
(A.3)

The square of the standard deviation of the potential energy change therefore equals

$$\sigma_{\Delta H^{\text{sol}}}^{2} = \sum_{i,j} \left[\sum_{k} \left[\left\langle s_{ijk}^{2} \right\rangle - \left\langle s_{ijk} \right\rangle^{2} \right] \Delta U_{ij}^{2} g(z_{k}, \tau)^{2} \right] + \sum_{k,l \neq k} \left[\left\langle s_{ijk} s_{ijl} \right\rangle - \left\langle s_{ijk} \right\rangle \left\langle s_{ijl} \right\rangle \right] \Delta U_{ij}^{2} g(z_{k}, \tau) g(z_{l}, \tau) \right].$$
(A.4)

This result involves the variances and co-variances of the occupation variable s_{ijk} . Note that $\langle s_{ijk} \rangle = c$. We have

$$\langle s_{ijk} s_{ijl} \rangle = \sum_{k} \frac{N_s!}{k!(N_s - k)!} e^k (1 - c)^{N_s - k}$$

$$\times \sum_{k_k, k_l} \frac{k!}{k_k! k_l! (k - k_k - k_l)!} \left(\frac{1}{N_s}\right)^{k_k + k_l} \left(1 - \frac{1}{N_s}\right)^{k - k_k - k_l} k_k k_l$$

$$= \sum_{k} \frac{N_s!}{k!(N_s - k)!} e^k (1 - c)^{N_s - k} \left(\frac{1}{N_s}\right)^2 k(k - 1)$$

$$= e^2 \left(1 - \frac{1}{N_s}\right),$$
(A.5)

with $1 \le k \le N_s$ and $0 \le k_k + k_l \le k$, and

$$\begin{split} \left\langle s_{ijk}^{2} \right\rangle &= \sum_{k} \frac{N_{s}!}{k!(N_{s}-k)!} c^{k} (1-c)^{N_{s}-k} \\ &\times \sum_{k_{k}} \frac{k!}{k_{k}!(k-k_{k})!} \left(\frac{1}{N_{s}}\right)^{k_{k}} \left(1-\frac{1}{N_{s}}\right)^{k-k_{k}} k_{k}^{2} \\ &= \sum_{k} \frac{N_{s}!}{k!(N_{s}-k)!} c^{k} (1-c)^{N_{s}-k} \\ &\times \left[\left(\frac{1}{N_{s}}\right)^{2} k^{2} + \frac{1}{N_{s}} \left(1-\frac{1}{N_{s}}\right) k\right] \\ &= \left(\frac{1}{N_{s}}\right)^{2} \left[c^{2}N_{s}^{2} + c(1-c)N_{s}\right] + c\left(1-\frac{1}{N_{s}}\right) \\ &= c + c^{2} \left(1-\frac{1}{N_{s}}\right), \end{split} \tag{A.6}$$

with $1 \le k \le N_s$ and $0 \le k_k \le k$. Inserting Eqs. (A.5) and (A.6) into Eq. (A.4) and neglecting the second order terms in c ($c^2 \ll 1$), we obtain

$$\sigma_{\Delta H^{\text{sol}}}^2 = \sum_{i,j} c \,\Delta U_{ij}^2 \sum_k g(z_k, \tau)^2 \tag{A.7}$$

Finally, by defining the z=0 to be located right at the double-kink nucleus, the above equation can be rewritten as

$$\sigma_{\Delta U_{sol}}^2 = \sum_{i,j} c \, \Delta U_{ij}^2 \sum_{k=-N_s}^{N_s} g(k \, b, \tau)^2. \tag{A.8}$$

Here, $N_s = l_{dk}/(2b)$

Appendix B. Double-kink nucleation rate

The total double-kink nucleation rate is the sum of the rates of all N nucleation sites which can be described as

$$R = \nu_0 N \exp\left(-\frac{\Delta H^0(\tau)}{kT}\right) \int_{\Delta \bar{H}^{sol}(c,N,\tau)}^{\infty} d\Delta H P[\Delta H] \exp\left(-\frac{\Delta H}{kT}\right), \quad (B.1)$$

where $P[\Delta H]$ is the probability of finding the double-kink nucleation segment with solute energy change ΔH given by Eq. (8). After some manupulations

the above equation can be rewritten as Eq. (9). It is of interest to find the temperature T_0 below which the weakest link dominates and thus the rate can be expressed by 15. To this end, we use the following approximated form for $\operatorname{erfc}(x)$ [36]

$$\operatorname{erfc}(x) \approx \frac{(1 - e^{-1.98x})e^{-x^2}}{1.135\sqrt{\pi}x}$$
 (B.2)

Substitution the above relation into Eq. (9) and equating the resultant equation with Eq. (15), gives the following solution for T_0 which is the temperature at which the two representations of double-kink nucleation rate become equal

$$\frac{k T_0}{\sigma_{\Delta H^{\rm sol}}(c,\tau)} = \left(0.35 N e^{-\frac{c_N^2}{2}} + 0.7 W \left[-0.5 N \exp \left(-0.5 N e^{-\frac{c_N^2}{2}} - \frac{c_N^2}{2}\right)\right] - c_N\right)^{-1} \tag{B.3}$$

where $W[\cdot]$ stands for the Lambert W-function and $c_N = b_N - \gamma a_N$. This temperature is approximately $k T_0 = 0.255 \, \sigma_{\Delta H^{\rm sol}}$ over a wide range of N. However, since $\sigma_{\Delta H^{\rm sol}}(c,\tau)$ depends on the applied stress τ which is not known a-priori, the exact computation of the cross-over temperature is not possible. It can be shown that, to a high approximation, T_0 can be obtained by solving the following equation

$$\Delta H^0_{\rm exp}(\tau) = 0.255\,\sigma_{\Delta H^{\rm sol}}(c,\tau)\log\left[250\frac{\dot{\epsilon}_0}{\dot{\epsilon}}\right].$$

The above equation gives the yield stress at temperature T_0 from which $\sigma_{\Delta H^{\rm sol}}$ and thus T_0 can be computed.

Appendix C. Simulation details of NEB calculations

To compute the minimum energy path (MEP) connecting the initial and final states, nudged elastic band (NEB) computations [37, 38, 39, 40] are performed as implemented in LAMMPS [41] using the interatomic potential developed by [2]. MEP calculations were started as follows. Using the Fe lattice parameter, a rectangular simulation cell oriented with [112] along the glide direction X, [110] along the glide plane normal direction Y, and [111] along the dislocation line direction Z, was constructed. Periodic boundary conditions are imposed

along X and Z directions, and Y having imposed tractions. The simulation cell 871 has dimensions $L_1 \sim 110A$ and $L_2 \sim 100A$ and L_3 depends on the dislocation length. A screw dislocation of Burgers vector a/2[111] with line direction along 873 Z is then introduced into the center of the cell by imposing a linear displacement 874 $u_z = -bx/l_x$ for $0 < x < l_x$ on all atoms in the upper half of the simulation 875 cell. This generates the initial atomic configuration. The final state has the 876 same structure as the initial state but shifted by a relative to the initial state. Atomic positions are then relaxed by using a combination of the FIRE algorithm 878 [42] and relaxation of the cell dimensions. Convergence is achieved when the 879 norm of the force vector fell below 10^{-6} eV/A and stresses σ_{XX} , σ_{XY} , and σ_{YY} 880 fell below 0.1 MPa. The above initial and final states are used as templates to carry out the MEP calculations. In order to model the Fe-Si dilute alloys, the Fe atoms in initial state configurations were replaced randomly by Si solutes at the 883 desired alloy concentration. Accordingly, the same Fe solutes were replaced by 884 Si atoms in the final state. The energy of the intial and final alloy configurations were then minimized using CG and FIRE with the same tolerences on forces and stresses as specified above. NEB simulations are performed using 60 replicas. An 887 initial path of intermediate configurations (replicas) is constructed by linearly 888 interpolating the atomic positions between the relaxed initial and final states. 889 The NEB inter-replica spring constant is set to 10^{-2} eV/Å² and convergence is assumed when the maximum of the force acting on all of the atoms across all replicas is less than $1 \times 10^{-3} \text{ eV/Å}$. 892

References

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- [1] D. R. Trinkle, The chemistry of deformation: How solutes soften pure
 metals, Science 310 (5754) (2005) 1665–1667.
- [2] S. Shinzato, M. Wakeda, S. Ogata, An atomistically informed kinetic monte
 carlo model for predicting solid solution strengthening of body-centered
 cubic alloys, International Journal of Plasticity 122 (2019) 319–337.

- Y. Zhao, J. Marian, Direct prediction of the solute softening-to-hardening
 transition in w-re alloys using stochastic simulations of screw dislocation
 motion, Modelling and Simulation in Materials Science and Engineering
 26 (4) (2018) 045002.
- J. Weertman, Dislocation model of low-temperature creep, Journal of Applied Physics 29 (12) (1958) 1685–1689.
- [5] E. Pink, Low-temperature softening in body-centered cubic alloys, Progress
 in Materials Science 24 (1) (1979) 1–50.
- [6] H. Suzuki, Solid solution hardening in body-centred cubic alloys, Dislocations in solids 4 (1980) 191–217.
- [7] J. R. Stephens, Dislocation structures in single-crystal tungsten and tung sten alloys, Metallurgical and Materials Transactions B 1 (5) (1970) 1293–
 1301.
- [8] Y.-J. Hu, M. R. Fellinger, B. G. Butler, Y. Wang, K. A. Darling, L. J.
 Kecskes, D. R. Trinkle, Z.-K. Liu, Solute-induced solid-solution softening
 and hardening in bcc tungsten, Acta Materialia 141 (2017) 304–316.
- [9] A. Argon, Strengthening mechanisms in crystal plasticity, Vol. 4, Oxford
 University Press on Demand, 2008.
- [10] L. Romaner, C. Ambrosch-Draxl, R. Pippan, Effect of rhenium on the dis location core structure in tungsten, Physical review letters 104 (19) (2010)
 195503.
- 921 [11] L. Romaner, Unpublished research.
- [12] M. Hossain, J. Marian, Stress-dependent solute energetics in w-re alloys
 from first-principles calculations, Acta materialia 80 (2014) 107–117.
- [13] T. Tsuru, T. Suzudo, First-principles calculations of interaction between solutes and dislocations in tungsten, Nuclear Materials and Energy 16 (2018)
 221–225.

- [14] S. Rao, B. Akdim, E. Antillon, C. Woodward, T. Parthasarathy, O. Senkov,
 Modeling solution hardening in bcc refractory complex concentrated alloys:
 Nbtizr, nb1. 5tizr0. 5 and nb0. 5tizr1. 5, Acta Materialia 168 (2019) 222–
 236.
- [15] H. Li, S. Wurster, C. Motz, L. Romaner, C. Ambrosch-Draxl, R. Pippan,
 Dislocation-core symmetry and slip planes in tungsten alloys: Ab initio cal culations and microcantilever bending experiments, Acta materialia 60 (2)
 (2012) 748–758.
- [16] F. Maresca, W. A. Curtin, Theory of screw dislocation strengthening in
 random bcc alloys from dilute to high-entropy alloys, Acta Materialia 182
 (2020) 144–162.
- 938 [17] Y.-J. Hu, Z.-K. Liu, Personal communication.
- [18] L. Dezerald, L. Proville, L. Ventelon, F. Willaime, D. Rodney, First-principles prediction of kink-pair activation enthalpy on screw dislocations
 in bcc transition metals: V, nb, ta, mo, w, and fe, Physical Review B 91 (9)
 (2015) 094105.
- [19] L. Proville, D. Rodney, M.-C. Marinica, Quantum effect on thermally activated glide of dislocations, Nature materials 11 (10) (2012) 845–849.
- [20] F. Maresca, D. Dragoni, G. Csányi, N. Marzari, W. A. Curtin, Screw dis location structure and mobility in body centered cubic fe predicted by
 a gaussian approximation potential, npj Computational Materials 4 (1)
 (2018) 1–7.
- 949 [21] U. F. Kocks, A. AS, A. MF, Thermodynamics and kinetics of slip.
- [22] L. Ventelon, F. Willaime, E. Clouet, D. Rodney, Ab initio investigation of
 the peierls potential of screw dislocations in bcc fe and w, Acta Materialia
 61 (11) (2013) 3973–3985.

- [23] A. Ghafarollahi, F. Maresca, W. Curtin, Solute/screw dislocation interaction energy parameter for strengthening in bcc dilute to high entropy alloys, Modelling and Simulation in Materials Science and Engineering 27 (8)
 (2019) 085011.
- [24] K. Kitajima, Y. Aono, H. Abe, E. Kuramoto, Solid solution hardening and
 softening in iron alloy single crystals between 4.2 k and 300 k, in: Strength
 of Metals and Alloys, Elsevier, 1979, pp. 965–970.
- [25] S. Takeuchi, Solid-solution strengthening in single crystals of iron alloys,
 Journal of the Physical Society of Japan 27 (4) (1969) 929–940.
- [26] H. Koizumi, H. Kirchner, T. Suzuki, Kink pair nucleation and critical shear
 stress, Acta Metallurgica et Materialia 41 (12) (1993) 3483–3493.
- [27] D. Brunner, J. Diehl, The effect of atomic lattice defects on the softening
 phenomena of high-purity α-iron, physica status solidi (a) 160 (2) (1997)
 355–372.
- [28] D. Caillard, A tem in situ study of alloying effects in iron. isolid solution
 softening caused by low concentrations of ni, si and cr, Acta materialia
 61 (8) (2013) 2793–2807.
- [29] S. Takeuchi, H. Yoshida, T. Taoka, Solid-solution strengthening in iron
 alloys, Transcations of the Japan Institute of Metals, Suppl 9 (1968) 715–
 719.
- [30] Y. Chen, D. Atteridge, W. W. Gerberich, Plastic flow of fe-binary alloysi.
 a description at low temperatures, Acta Metallurgica 29 (6) (1981) 1171–
 1185.
- [31] A. Stukowski, D. Cereceda, T. D. Swinburne, J. Marian, Thermally activated non-schmid glide of screw dislocations in w using atomistically informed kinetic monte carlo simulations, International Journal of Plastic ity 65 (2015) 108–130.

- [32] W. J. Szlachta, A. P. Bartók, G. Csányi, Accuracy and transferability of
 gaussian approximation potential models for tungsten, Physical Review B
 90 (10) (2014) 104108.
- [33] P. L. Raffo, Yielding and fracture in tungsten and tungsten-rhenium alloys,
 Journal of the Less Common Metals 17 (2) (1969) 133–149.
- [34] J. R. Stephens, W. R. Witzke, Alloy softening in group via metals alloyed
 with rhenium, Journal of the Less Common Metals 23 (4) (1971) 325–342.
- [35] F. Maresca, W. A. Curtin, Mechanistic origin of high strength in refractory
 bcc high entropy alloys up to 1900k, Acta Materialia 182 (2020) 235–249.
- [36] G. K. Karagiannidis, A. S. Lioumpas, An improved approximation for the
 gaussian q-function, IEEE Communications Letters 11 (8) (2007) 644–646.
- [37] G. Henkelman, H. Jónsson, Improved tangent estimate in the nudged elastic
 band method for finding minimum energy paths and saddle points, The
 Journal of chemical physics 113 (22) (2000) 9978–9985.
- [38] G. Henkelman, B. P. Uberuaga, H. Jónsson, A climbing image nudged
 elastic band method for finding saddle points and minimum energy paths,
 The Journal of chemical physics 113 (22) (2000) 9901–9904.
- 997 [39] A. Nakano, A space–time-ensemble parallel nudged elastic band algorithm for molecular kinetics simulation, Computer Physics Communications 178 (4) (2008) 280–289.
- [40] E. Maras, O. Trushin, A. Stukowski, T. Ala-Nissila, H. Jonsson, Global
 transition path search for dislocation formation in ge on si (001), Computer
 Physics Communications 205 (2016) 13–21.
- [41] S. Plimpton, Fast parallel algorithms for short-range molecular dynamics, Tech. rep., Sandia National Labs., Albuquerque, NM (United States)
 (1993).

[42] E. Bitzek, P. Koskinen, F. Gähler, M. Moseler, P. Gumbsch, Structural
 relaxation made simple, Physical review letters 97 (17) (2006) 170201.

Declaration of interests

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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