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Diffusion and interaction of prismatic dislocation loops simulated by stochastic discrete dislocation dynamics

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Body-centered cubic metals and alloys irradiated by energetic particles form highly mobile prismatic dislocation loops with a/2⟨111⟩-type Burgers vectors. We show how to simulate thermal diffusion of prismatic dislocation loops using a discrete dislocation dynamics approach that explicitly includes the stochastic forces associated with ambient thermal fluctuations. We find that the interplay between stochastic thermal forces and internal degrees of freedom of loops, in particular the reorientation of the loop habit planes, strongly influences the observed loop dynamics. The loops exhibit three fundamental types of reactions: coalescence, repulsion, and confinement by elastic forces. The confinement reactions are highly sensitive to the internal degrees of freedom of the loops. Depending on the orientation of the loop habit planes, the barrier to enter an elastically confined bound state is lowered substantially, whereas the lifetime of the bound state increases by many orders of magnitude.

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I. INTRODUCTION

Metals exposed to irradiation develop a highly complex microstructure, involving a mixture of mobile and immobile defects of both interstitial and vacancy type. The defect and dislocation network develops under the effect of internal and external stresses, and temperature, generating its own fluctuating stress field. This leads to a variety of changes in mechanical properties, such as hardening and the loss of ductility, having a detrimental effect on the longevity of structural reactor components in a radiation environment.

Predicting the dynamics of evolution of microstructure is a major challenge to computer modeling because of the broad spectrum of activation energies characterizing defect and dislocation networks. Defect cluster migration barriers vary from meVs to eVs. The binding energy of elastically confined defect structures spans a similar range of energy scales [1], and the magnitude of elastic interaction depends on the size of defects and their spatial distribution. Simulating the temperature dependent dynamics of microstructure requires the treatment of intrinsic thermally activated Brownian motion of defects and dislocations, as well as correlated motion of defects and dislocations mediated by elastic interactions.

Highly glissile prismatic dislocation loops are produced by irradiation [2,3] together with sessile cavities, as evidenced by in situ transmission electron microscopy (TEM) observations [4,5]. The correlated motion of dislocation loops, often observed experimentally, is an elementary process leading to the formation of rafts of defects and their eventual coalescence [6–9]. In other words, the spatial ordering of dislocation loops stems from their elastic interaction, whereas the loop motion itself is a thermally activated process, fundamentally the same as stochastic Brownian motion of individual defects [9–14]. The subject of this paper is the simulation of stochastic glide motion of prismatic a/2⟨111⟩ dislocation loops in body-centered cubic (bcc) iron, with a particular emphasis on the analysis of elementary reactions between the loops, treated as dislocation line objects, and modeled using discrete dislocation dynamics.

Molecular dynamics and lattice type simulations performed over the past two decades investigated the stochastic diffusion of prismatic loops over a range of sizes and temperatures [10–12,15–18], elementary loop and dislocation reactions [1,9], as well as energies of binding of loops to other defects [19]. However, a direct atomistic simulation of an ensemble of interacting dislocation loops still remains a challenge because of the constraint imposed by the simulation cell size accessible to a molecular dynamics simulation, and the relatively short timescale of such a simulation. While the more recent atomistic approaches involving the use of kinetic Monte Carlo [20–23] have reached the experimentally relevant timescale when exploring the relaxation of radiation cascade damage in thin films, the identification of pathways of migration and reaction between interacting dislocation loops containing more than a few dozen interstitials remains a largely unexplored problem.

Discrete dislocation dynamics (DDD) provides a compelling alternative approach to modeling complex dislocation microstructures, offering highly efficient computation of long-range elastic interactions. Furthermore, dislocation dynamics enables the treatment of dislocation reactions, simultaneously simulating internal and collective dislocation loop dynamics, and enabling the investigation of complex networks and junctions within the same methodological framework. We note that bringing dislocation core properties on par with atomistic
simulations remains a challenge, and there have been recent extensive developments addressing this issue [24–32].

The objective of this work is to include thermal stochastic forces in DDD through the Langevin stochastic formalism, to enable modeling the Brownian motion and diffusion of dislocations. The stochastic dislocation dynamics [33,34] approach is formulated and applied to the treatment of diffusion of loops and elementary reactions between interacting loops as an essential step towards modeling thermal evolution of complex dislocation ensembles.

Langevin dynamics has been applied earlier to the treatment of collective dynamics of dislocation loops on a coarse-grained level, where the loops were treated as pointlike objects interacting through long-range elastic fields described in the elastic dipole tensor approximation [1,9,35,36]. Extending the treatment to the case where loop dynamics involves also the relaxation of their internal degrees of freedom, such as tilting of the loop habit plane, we find that this strongly increases the lifetime of configurations where pairs of loops are bound together by their attractive elastic fields. Furthermore, the barriers to entering such bound states are strongly reduced, explaining why dislocation loop rafts are able to form so easily in many materials, as confirmed by in situ TEM observations [1,9,37].

The paper is organized as follows. In Sec. II we derive an expression for thermal stochastic forces acting on a dislocation line. In Sec. III the diffusion coefficient of a single prismatic loop is evaluated and examined as a function of temperature, and the DDD analysis is benchmarked against molecular dynamics simulations. We also discuss the fluctuation-dissipation theorem, relating the amplitude of stochastic thermal forces to the magnitude of dissipative forces, the motion for a dislocation line is one dimensional:

\[
B \frac{\partial z(l,t)}{\partial t} = f^\text{tot}(l,t) + f^s(l,t),
\]

where \( B \) is the viscous drag coefficient for the given slip system and the stochastic force \( f^s \) is assumed to be uncorrelated in time and space:

\[
\langle f^s(l,t) \rangle = 0,
\]

\[
\langle f^s(l,t)f^s(l',t') \rangle = \sigma_s^2 \delta(l-l')\delta(t-t'),
\]

where \( \sigma_s \) sets the scale of the stochastic force, and \( \delta(x) \) is the Dirac delta-function

\[
\delta(x) = 0, \quad \forall x \neq 0, \quad \int_{-\infty}^{\infty} dx \, \delta(x) = 1.
\]

To find the amplitude of stochastic force, the effective diffusion coefficient for the center-of-position (COP) of the prismatic loop is matched to a 1D Einstein diffusion law [18]. The projection of the COP on the Burgers vector direction is given by \( z_{\text{COP}} = L^{-1} \int_0^L dl \, z(l,t) \). After a rearrangement, Eq. (4) becomes

\[
\frac{\partial z_{\text{COP}}}{\partial t} = \frac{1}{BL} \int_0^L dl \, f^s(l,t).
\]

The effect of internal elastic force on the COP vanishes due to the boundary condition \( z(l+L,t) = z(l,t) \). Assuming that the initial position of the loop center is \( z_{\text{COP}}(0) = 0 \), the solution to Eq. (7) at time \( \tau \) is

\[
z_{\text{COP}}(\tau) = \frac{1}{BL} \int_0^L dl \int_0^\tau dt \, f^s(l,t). \tag{8}
\]
Since the stochastic force $f_s(l, t)$ is defined in terms of its correlation function, the mean square displacement of the COP can be expressed as

$$\langle z_{\text{COP}}^2(\tau) \rangle = \left( \frac{1}{BL} \right)^2 \int_0^{l_N} dl \int_0^{l_N} dl' \int_0^{\tau} dt \int_0^{\tau} dt' \langle f_s(l, t) f_s(l', t') \rangle.$$

(9)

Substituting Eq. (5) into Eq. (9) yields

$$\langle z_{\text{COP}}^2(\tau) \rangle = \left( \frac{\sigma_t}{BL} \right)^2 \int_0^{l_N} dl \int_0^{l_N} dl' \int_0^{\tau} dt \int_0^{\tau} dt' \delta(l - l') \delta(t - t').$$

(10)

Evaluating the above integral, we arrive at

$$\langle z_{\text{COP}}^2(\tau) \rangle = \left( \frac{\sigma_t}{BL} \right)^2 \tau \equiv 2D_{\text{COP}} \tau.$$

(11)

This equation is a mere corollary of the 1D Einstein diffusion law, where $D_{\text{COP}}$ is the corresponding diffusion coefficient of the center-of-position [43]. Substituting the fluctuation-dissipation theorem [44]

$$D_{\text{COP}} = \frac{k_B T}{BL}$$

(12)

into Eq. (11), which holds under the assumption that the dislocation loop is in thermodynamic equilibrium with the thermostat, we find the amplitude of the stochastic force

$$\sigma_t = \sqrt{2k_B T B},$$

(13)

where $k_B$ is the Boltzmann constant and $T$ is absolute temperature.

As NUMODIS is a nodal dislocation dynamics code, the total force per unit length is converted into an effective nodal force by integrating over the neighboring segments [40]. The same rule is applied to convert the stochastic force per unit length into a stochastic force acting on a node. However, care must be taken when rescaling the force, as the randomly applied force must remain consistent with the choice of the segment length and the integration time step. For a straight segment of length $\Delta l$ indexed by $n$, the scaled stochastic force per unit length is found using the stochastic average:

$$f_{s,n}^a(t) = \frac{1}{\Delta l} \int_{l_{n-\Delta l/2}}^{l_{n+\Delta l/2}} dl f_s(l, t).$$

(14)

Using Eq. (5), for the $\delta$-correlated force generated by thermal fluctuations, the nodal correlation function acquires the form

$$\langle f_{s,n}^a(t) f_{s,n}^a(t') \rangle = \frac{\sigma_t^2}{\Delta l^2} \int_{l_{n-\Delta l/2}}^{l_{n+\Delta l/2}} dl \int_{l_{n-\Delta l/2}}^{l_{n+\Delta l/2}} dl' \times \delta(l - l') \delta(t - t').$$

(15)

Similarly, assuming an integration time step of $\Delta t$, the scaled stochastic force per unit length can be finally expressed as

$$f_{s,n}^a = \sqrt{\frac{2k_B T B}{\Delta l \Delta t}} N(0, 1),$$

(16)

where $N(0, 1)$ is a random number sampled from the standard normal distribution, and the direction of the force is collinear with the Burgers vector of the dislocation loop.

**B. Simulation setup, parameters, and statistics**

All the dislocation dynamics (DD) simulations were performed assuming an infinite elastic medium. The coordinate system is chosen as orthogonal with axes parallel to $x = [\overline{T}T2]$, $y = [\overline{T}0]$, and $z = [111]$ directions. Initially, a hexagonal prismatic (111) dislocation loop is positioned at the origin. The loop radius is chosen as $\rho = 4.5$ nm, corresponding to the loop perimeter of $L = 27$ nm. The hexagonal loop shape was chosen out of convenience as this has an almost negligible effect on its dynamics. A circular loop of equivalent size would have the radius of $4.09$ nm, representing an inclusion containing the same amount of matter.

The three parameters included in the stochastic force (16) require further clarification.

The viscous drag coefficient $B$ characterizes the drag force acting on a dislocation line. In bcc metals it is generally assumed that $B(T) = B_0 + B_1 T$, where $B_0$ and $B_1$ are independent of temperature [24,45,46]. MD simulations of glissile prismatic loops and self-interstitial clusters in bcc metals show that $B(T) = B_0$ and is independent of $T$ over a wide temperature range. Given that the simulations performed in this study address prismatic dislocation loops of very small size, it is appropriate to treat $B$ as a temperature-independent constant. The numerical value of $B$ used in this work has been evaluated from the atomistic study by Derlet et al. [18] using the fluctuation-dissipation relation (12). The resulting value of the drag coefficient $B = 0.08$ MPa ns describes the effective mobility of edge dislocations at temperatures above $T = 200$ K, but underestimates the magnitude of drag at lower temperatures, as shown in Fig. 1. At low temperatures, the Peierls barrier [47,48] and quantum effects [49,50] play an important part, affecting dislocation mobility, but are not considered in this study. The chosen value of $B = 0.08$ MPa ns agrees well with previous parametrizations derived from the analysis of dislocation-defect interactions in bcc iron [51,52].

Since the simulations were performed by splitting dislocation loops into straight segments, and involved solving the equations of motion by means of a finite difference time integration algorithm, it would be appropriate to assess the effect of discretization length $\Delta l$ and time step $\Delta t$ on the computed diffusion coefficient. Thermal diffusion of a single prismatic loop at 300 K was simulated using three discretization lengths, $\Delta l = 5, 10,$ and $15$ Å, and three time steps, $\Delta t = 0.2, 0.5,$ and $1.0$ fs. Simulations were run over the interval of 6 ns, with the loop configuration data recorded every 0.6 ps. Loop diffusion coefficients were computed using the drift diffusion correction method [18], in which the diffusion trajectory was split into multiple uncorrelated sub-trajectories. The velocity autocorrelation function $\langle v_{\text{COP}}(t) v_{\text{COP}}(t + \tau) \rangle$ yields the correlation time of $\tau \approx 2$ ps, in broad agreement with atomistic estimates [13,29]. The velocity correlation time is longer.
anisotropic moduli used as input were chosen following dislocation loop shapes and stress fields. In this work, the that this method leads to highly accurate predictions of μ and ν (dots). The viscosity is well described by a non-Arrhenius relation B functional expression. The dashed line corresponds to the constant above 200 K.

than the stochastic force correlation time [45] derived from atomistic simulations, and represents the low limit for the time length of a subtrajectory, which here was chosen as 6 ps. The diffusion coefficient is then found by ensemble averaging over the subtrajectories, with the uncertainty characterized by the standard error of the mean.

Figure 2 shows a selection of simulated COP trajectories, which are similar in terms of their statistical properties. The values of diffusion coefficient derived from these trajectories remain within their respective error bounds, independent of the selected values of Δl and Δt, in agreement with the theoretical analysis by Derlet et al. [18].

Following Scattergood and Bacon [53], the elastic moduli μ and ν are chosen by matching the isotropic and anisotropic elasticity energies of infinite a/2[111][110] edge and screw dislocations. Earlier comprehensive studies [54–56] confirm that this method leads to highly accurate predictions of dislocation loop shapes and stress fields. In this work, the anisotropic moduli used as input were chosen following Ackland et al. [57] as C_{11} = 225 GPa, C_{12} = 124 GPa, and C_{44} = 101 GPa, leading to the corresponding isotropic moduli of μ = 63 GPa and ν = 0.43.

The dislocation core radius R_c, which here has the same meaning as the delocalization parameter of the nonsingular elasticity theory [39], and the core strength parameter ξ were evaluated in earlier studies [51,58] from atomistic simulations.

Isotropic elasticity theory represents an effective approximation for bcc iron, which is an elastically anisotropic material. The numerical results presented in this work should therefore be interpreted qualitatively. This is a foregone conclusion if we acknowledge the fact that the commonly accepted models for the dislocation core energy do not capture the complex anisotropic configurational energy landscape found in atomistic simulations. The stochastic forces formalism itself is directly applicable to DDD simulations in elastically anisotropic materials.

All the further simulations presented in Sec. III were carried out using the simulation parameters given in Table I, unless specified otherwise. The integration time step scales approximately as a cube of the dislocation discretization length, see Appendix A. We have opted to use a fine time step and small discretization length, as computational efficiency and performance was not a concern in the simulations presented here.

### III. RESULTS

A. Stochastic dynamics of an individual dislocation loop

Using stochastic dislocation dynamics, we performed a series of simulations, investigating the dynamics of a single prismatic loop at temperatures ranging from 100 to 800 K, with temperature increments of 100 K. No external stress was applied.

Consider first the internal degrees of freedom of the prismatic loop. It is readily seen from simulations that the initially purely prismatic [111] loop with its Burgers vector normal to its habit plane, within a few picosecond adopts a tilted configuration, see Fig. 3(a). If the shape of the loop is defined by its dislocation countour C, the vector area of the loop is given by [36,59] 

\[ A = \frac{1}{2} \oint_C r \times dl, \] 

(17)
FIG. 3. (a) Snapshots from a stochastic dislocation dynamics simulation of a hexagonal initially pure prismatic loop of 4.5 nm radius at 100 K show that the loop habit plane becomes tilted within a few picoseconds. (b) The tilt angle \( \theta \) is defined as the angle between the normal vector (red arrow) and the Burgers vector (black arrow). (c) The prismatic loop adopts a tilted configuration on the glide cylinder to minimize its potential energy.

and the effective loop normal unit vector is

\[
\hat{n} = \frac{1}{2||A||} \oint_C r \times dl.
\]  

(18)

The angle between the Burgers vector and the effective loop normal shall be referred to as the tilt angle \( \theta \), with the azimuthal angle \( \phi \) defined in full analogy with the spherical system of coordinates, see Fig. 3(b) for illustration. Following this definition and depending on the nature of the loop (vacancy or interstitial), the loop is pure prismatic if \( \hat{n} \cdot \hat{b} = \pm 1 \) corresponding to \( \theta = 0 \) or 180°. We note that the elastic relaxation volume of a loop is given by the scalar product of the Burgers vector and the loop vector area \( \Omega_{rel} = \hat{b} \cdot A \) [36].

The elastic potential energy of a prismatic loop is minimized for configurations tilted away from the perfect prismatic loop orientation, with the resulting tilt angle \( \theta \) determined by the competition between the elastic self-energy associated with interaction between dislocation segments and the core energy proportional to the length of the perimeter of the loop, see Fig. 3(c). The potential energy is invariant with respect to rotations around the Burgers vector, allowing the loop to rotate freely with respect to \( \phi \) in a DD simulation.

The mean value of the tilting angle \( \langle \theta \rangle \) decreases at higher temperatures, reflecting the anharmonicity of the potential self-energy of the loop. Indeed, it takes comparatively less energy for the loop normal to tilt towards the Burgers vector than away from it, hence on average smaller values of \( \theta \) are favored at higher temperature.

In addition to the tilting degrees of freedom, the loop shape also develops transient fluctuations on a smaller scale. However, any part of the loop is constrained to remain on the glide cylinder, as the relaxation volume of the loop \( \Omega_{rel} = \hat{b} \cdot A \) is conserved throughout the simulation.

Consider next the diffusion behavior of the entire loop. The prismatic loop trajectories exhibit a characteristic pattern of Brownian motion, with higher temperature inducing a more pronounced loop displacement per unit time. The single loop COP trajectories for 200 and 600 K, and the diffusion coefficients calculated with the drift diffusion correction [18], are given in Fig. 4. Globally, the temperature dependence of the diffusion coefficient is found to be consistent with the fluctuation-dissipation theorem, regardless of the loop radius \( \rho \).

Moreover, for \( \rho = 4.5 \) nm and \( T < 400 \) K the diffusion coefficients derived from simulations are consistently lower than expected from the linear interpolation from higher temperature (dashed line) because the tilting of the loop results in the elongation of its perimeter, see Fig. 3(b). According to the fluctuation-dissipation theorem, \( D_{COP} \propto 1/L \), and therefore the reorientation of the habit plane gives rise to a lower value of the diffusion coefficient. This effect is found to become less pronounced at higher temperature as the mean tilt angle \( \langle \theta \rangle \) decreases with temperature.

The stochastic DD simulations performed in this work describe thermally induced Brownian motion of prismatic loops, which for \( T > 200 \) K is consistent with molecular dynamics. The simulations further reveal that the prismatic loop habit plane becomes tilted with respect to the Burgers vector, while
the Burgers vector direction, see Fig. 5(a). In the absence of attractive elastic interactions, particularly where the loops form bound configurations confined by barriers in relation to its rotation around the Burgers vector. Symmetry of the system, subsequently introducing energy combination with atomic discreteness would break the cylindrical nature of the core energy. Considering that the core energy scales linearly with the loop radius \( \propto \rho \) and \( \rho \log \rho \) [61], one would expect the core energy to become less significant for larger loops. However, the singular nature of the core energy in combination with atomic discreteness would break the cylindrical symmetry of the system, subsequently introducing energy barriers in relation to its rotation around the Burgers vector.

B. Diffusion of interacting dislocation loops

The question about thermal evolution of interacting dislocation loops has recently attracted attention in the context of dipole tensor formalism as an efficient approximation for the core energy [60]. Considering that the core energy scales linearly with the loop radius \( \propto \rho \), whereas the elastic self-energy varies superlinearly as \( \propto \rho \log \rho \) [61], one would expect the core energy to become less significant for larger loops. However, the singular nature of the core energy in combination with atomic discreteness would break the cylindrical symmetry of the system, subsequently introducing energy barriers in relation to its rotation around the Burgers vector.

![Diagram of two interacting loops](image)

**FIG. 5.** (a) Two prismatic loops defined using the coordinate system introduced in Sec. III B, with identical Burgers vectors \( \mathbf{b} \) parallel to \( z \) direction. Loop configuration (line) is free to deviate from the pure prismatic form (dashed) on the glide cylinder, as indicated by the loop normal vectors \( \hat{n} \). (b) A selection of representative metastable configurations of interacting loops extracted from dislocation dynamics simulations, also showing the loop normal and Burgers vectors. Configurations are ordered from top to bottom in the order of increasing stability, and hence in the order of descending total potential energy. Configuration 3 is the most stable configuration.

remaining highly mobile with respect to rotations around the Burgers vector.

The tilting behavior of prismatic loops in atomistic simulations is possibly dominated by singular orientation effects in the core energy [60]. Considering that the core energy scales linearly with the loop radius \( \propto \rho \), whereas the elastic self-energy varies superlinearly as \( \propto \rho \log \rho \) [61], one would expect the core energy to become less significant for larger loops. However, the singular nature of the core energy in combination with atomic discreteness would break the cylindrical symmetry of the system, subsequently introducing energy barriers in relation to its rotation around the Burgers vector.

Energies of elastic interaction are compared in Fig. 6 for the various loop separations \( \Delta x \) using an example of two round loops with radii \( \rho = 4.09 \) nm. Note that the choice of radii is consistent with loops being hexagonal and having the same area, as discussed in Sec. II B.

The exact interaction energy trend for pure prismatic pairs of loops broadly follows the PES trend, but does not reflect the full complexity of interaction between internally relaxed loops. The dipole approximation is consistent with the exact treatment, but only for loop separations several times larger than the sum of loop radii. The dipole tensor formalism becomes inaccurate for smaller separations, resulting in a qualitatively incorrect predicted interaction behavior, see the top two panels in Fig. 6.

A major effect of internal relaxation is found when we follow how the loops approach an elastically confined bound state from infinite separation. This reaction is fundamental to the formation of dislocation loop rafts. From Fig. 6 it is
their energy ordering is consistent with the separations $\Delta x$ studied here.

This comparison demonstrates that the energy of interaction between prismatic loops is strongly affected by the internal degrees of freedom of the loops. Consequently, the competition between the elastic energy and the core energy plays a pivotal role in determining the landscape of binding energies of loops. This subtlety is neglected in any physical approximation where the dislocation loops are treated as being purely prismatic, or where they are treated as pointlike objects defined only by their position in real space and involving no consideration of their internal degrees of freedom.

In what follows, we carry out stochastic dislocation dynamics simulations of interacting pairs of loops. The simulations start from large initial separations $\Delta x$ and $s$ at 200 K in an attempt to emulate various elementary interactions observed in experiment, see Sec. I, namely coalescence, repulsion, and mutual elastic confinement of interacting loops.

**Case A: Coalescence of dislocation loops**

The coalescence of dislocation loops was observed using TEM and was found to involve loops of comparable size [8], with diameters larger than 4 nm. To match experimental observations, two pure prismatic hexagonal (111) loops with $\rho = 4.5$ nm are introduced in a simulation cell with separations of $\Delta x = 8$ nm and $s = 5$ nm, giving rise to a mutually attractive elastic force, see Fig 6. Note that the glide cylinders of the loops overlap slightly. Sequential snapshots taken during simulations are shown in Fig. 7(a). The loops coalesce into a larger prismatic loop, with small debris released and ejected by a strong repulsive elastic force.

The corresponding time evolution of the diffusion coefficient of the resulting large loop is shown in Fig. 7(b). We observe that the diffusion coefficient becomes constant over the interval of a few nanoseconds and converges to a notably smaller value than the diffusion coefficient of a single loop with $\rho = 4.5$ nm. Using the $D_{\text{coarse}} \propto 1/L$ scaling relation, the equivalent loop size of the loop produced by the coalescence of a pair of loops equals $\rho_{\text{eq}} \approx 7$ nm. This is consistent with an estimate of the equivalent loop size obtained by removing a quarter of each loop’s circumference, leading to $\rho_{\text{eq}} \approx 3/2\rho$. While the relaxation volume of the loops is a conserved quantity, the length of the loop circumference is not; this example demonstrates clearly that the effective diffusivity of an ensemble of prismatic loops may reduce over time as a result of coalescence of loops.

**Case B: Repulsion between the loops**

An example of repulsive interaction between diffusing dislocation loops is obtained by placing the loops with separations of $\Delta x = 5$ nm and $s = 7$ nm, using the different initial configurations shown in Fig. 5(b). Note that configuration 1 was placed at a separation of $s = 12$ nm, as its corresponding PES branch vanishes at closer separations. Figure 8 shows the evolution of the corresponding interaction energies during the simulation performed without stochastic forces ($T = 0$ K) and with stochastic forces ($T = 200$ K) included, in comparison with the theoretical prediction derived from examining the corresponding potential energy surface.
FIG. 7. (a) Snapshots taken from stochastic dislocation dynamics simulations of a loop coalescence reaction, for the initial loop-pair separations of $\Delta x = 8\,\text{nm}$ and $s = 5\,\text{nm}$, viewed at an angle from the $-\hat{y}$ direction. Note the occurrence of ejection of debris during loop coalescence. (b) Plot of the effective diffusion coefficient as a function of time. The dotted line is a reference value computed for a single loop with size $\rho = 4.5\,\text{nm}$.

As expected for repulsive configurations, we find that the distance between the loops gradually increases over the interval of time spanned by the simulation. Inspection of the loop-pair configuration shows that the cold ($T = 0\,\text{K}$) systems retain their initial orientation of the habit plane, which is consistent with the energy trajectories propagating along the distinct PES branches. On the other hand, the trajectories of the heated system ($T = 200\,\text{K}$) soon start overlapping, starting from $s \approx 20\,\text{nm}$, eventually becoming indistinguishable. The stochastic force supplies additional thermal energy to the loops, which is evidently sufficient to overcome the energy barrier between the different PES branches, enabling the loops to rotate and thus oscillate between various tilting configurations.

**Case C: Elastic confinement of loops**

Prismatic loops may exhibit strong elastic attraction and form an elastically confined configuration as seen in Fig. 6. Depending on loop size and loop separation, the binding energy can vary from meVs to eVs, potentially surpassing the binding energy of dislocations to substitutional defects. Therefore it can be reasoned that elastic confinement of loops represents the key step leading to the stabilization of experimentally observed rafts of dislocation loops.

We adopt the initial setup corresponding to $\Delta x = 12\,\text{nm}$ and $s = 12\,\text{nm}$, for which the pair of loops exhibit mutual attraction. As in the repulsive case investigated above, the simulations were run for three initial loop configurations shown in Fig. 5(b), corresponding to distinct branches of the potential energy surface. The evolution of the energy of interaction between the two loops as a function of their separation in comparison with the idealized PES is shown in Fig. 9.

In the absence of stochastic forces, the two-loop system is hindered from reaching the lowest energy state because it is unable to overcome the energy barrier associated with the rotation of the loop habit planes. In contrast, the addition of stochastic forces supplies the loops with additional energy, enabling the system to explore the potential energy landscape more freely to the point where it even oscillates around the global energy minimum. As in the loop repulsion case investigated above, the interaction energy derived using simulations involving elevated temperature is found to be shifted upwards by about 1.5 eV compared to the PES, as the Langevin thermostat adds additional energy to the system.

The COP trajectories of the two loops corresponding to conf. 1 state are shown in Fig. 10. After a brief initial relaxation time, the loops become mutually trapped in their relative frame by attractive elastic interaction, with their COP trajectories becoming strongly spatially correlated. The loop separation distance in the elastically confined state fluctuates around the global potential energy minimum as a result of the effect of stochastic force, in agreement with experimental observations and simulations reported in Figs. 3–5 of Ref. [1]. Interestingly, the simulated trajectories suggest that the two bound loops oscillate on a $\sim 0.5\,\text{ns}$ timescale, thus evolving significantly slower than the tilt angle of the isolated loop, see Fig. 3.
over the duration of the simulation. In this paper we show that the internal degrees of freedom significantly influence the nature of interaction between the loops, with potentially significant implications for the lifetime of elastically confined loop configurations. We now analyze this effect quantitatively.

Introducing the probability density $P(s, t)$ of finding the two loops at separation $s$ at time $t$, the equation of motion for this probability density, see Appendix C, can be expressed as a Fokker-Planck equation

$$\frac{\partial P(s, t)}{\partial t} = -\frac{\partial J(s, t)}{\partial s},$$

(20)

where $J$ is the flux of the probability density

$$J(s, t) = -\frac{2}{\beta BL} e^{-\beta V(s)} \frac{\partial}{\partial s} \left( e^{\beta V(s)} P(s, t) \right),$$

(21)

where $\beta = 1/(k_B T)$ and $V(s)$ refers to a branch of the potential energy surface. It is sufficient to consider the $s \in [0, +\infty)$ interval of variation of $s$ as the potential energy surface is symmetric. In the following discussion we assume that the potential energy surface includes a confined state that transforms, over an energy barrier, into an unbound state at large separation $s$. While this assumption is valid for a pair of loops, caution needs to be taken in densely populated microstructures, as the potential energy can be strongly distorted by the elastic field of other defects.

Consider now the pair of loops at an energy minimum at $s_{\text{min}}$. At equilibrium steady state the flux vanishes, $J = 0$, leading to the probability acquiring the form of the Gibbs distribution $P(s) \sim \exp \left[ -\beta V(s) \right]$. Similarly, the escape process from the energy minimum at $s_{\text{min}}$ to a very far separation along the glide direction $s_{\text{far}}$ can be considered to proceed slow enough to preserve the steady state, leading to a constant flux $J = J_0$. The steady-state flux is found by solving Eq. (21) for the derivative and subsequently integrating from $s_{\text{min}}$ to $s_{\text{far}}$, namely

$$\left[ e^{\beta V(s)} P(s) \right]_{s_{\text{min}}}^{s_{\text{far}}} = -\frac{J_0 \beta BL}{2} \int_{s_{\text{min}}}^{s_{\text{far}}} ds e^{-\beta V(s)}. \quad (22)$$

Using $P(s_{\text{min}}) \gg P(s_{\text{far}})$, the escape flux can be found as

$$J_0 \approx \frac{2}{\beta BL} \int_{s_{\text{min}}}^{s_{\text{far}}} ds e^{\beta V(s)}. \quad (23)$$

Assuming that the probability density decays rapidly outside the potential well associated with the energy minimum, the probability $p$ of finding the pair of loops in an elastically confined state is derived by integration over the well width $\pm \delta s$, using the method of steepest descent

$$p = \int_{s_{\text{min}} - \delta s}^{s_{\text{min}} + \delta s} ds P(s)$$

$$= P(s_{\text{min}}) \int_{s_{\text{min}} + \delta s}^{s_{\text{min}} - \delta s} ds e^{-\beta(V(s_{\text{min}}) - V(s))}$$

$$\approx \frac{P(s_{\text{min}})}{2} \int_{-\infty}^{+\infty} ds e^{-\beta V'(s_{\text{min}})^2/2}$$

$$= \frac{P(s_{\text{min}})}{2} \left( \frac{2\pi}{\beta V''(s_{\text{min}})} \right)^{1/2} \quad (24),$$

FIG. 9. Dislocation dynamics simulation of a pair of prismatic loops in an attractive arrangement without (a) and with stochastic forces at $T = 200$ K (b) included. The initial configurations (conf.) of loops are taken from Fig. 5. Trajectories in (b) are shifted down by an estimated amount of additional thermal energy supplied by the thermostat $W^\text{th}$ for better comparison. In the absence of stochastic forces, the loop pair is stuck in metastable configurations. In contrast, the heated system escapes from the metastable state, instead fluctuating around the global minimum.

IV. THE LIFETIME OF ELASTICALLY CONFINED LOOP CONFIGURATIONS

In Sec. III B above, we have explored the three types of fundamental reactions between prismatic dislocation loops. These reactions, namely loop coalescence, repulsion, and mutual elastic trapping or confinement, have all been modeled using stochastic dislocation dynamics at 200 K. The simulations enable comparison with models developed earlier for simulating the thermal evolution of multiple loops, which involved the dipole approximation [9] and treat the loops as point objects, assuming that they remain purely prismatic.
where $V''(s_{\text{min}})$ is the second derivative of $V(s)$ evaluated at the stationary point $s_{\text{max}}$. Similarly, the integral term in Eq. (23) peaks at the point $s_{\text{max}}$ corresponding to the maximum barrier height. Applying the same approach as in Eq. (24), the escape rate $\Gamma$ can be finally expressed as

$$
\Gamma = \frac{I_0}{\rho} = \frac{V'(s_{\text{min}})V''(s_{\text{max}})}{2\pi BL} e^{-\beta \Delta V},
$$

where $\Delta V = V(s_{\text{max}}) - V(s_{\text{min}})$ is the energy barrier that the pair of loops has to overcome in order to separate. The inverse of the escape rate equals the lifetime of the confined state of the loops. Under these conditions, a small variation of $\Delta V$ can significantly affect the lifetime.

Consider now the choice of the potential barrier $V(s)$. We use three different tilting configurations to investigate the effect of internal degrees of freedom on the confinement lifetime. First, the pure prismatic pair of loops is a reference configuration to models involving no internal degrees of freedom. Next, the freely tilting loop pair is represented by the lowest energy curve of the PES. Finally, the tilting of each loop is fixed ad-hoc at an angle of $30^\circ$ each [unfavorable in energy, see conf. 1 in Fig. 5(b)], in an attempt to mimic the habit plane locking observed in molecular dynamics. The corresponding lifetimes of elastically confined loop configurations are listed in Table II for circular loops with $\rho = 4.09$ nm and $\rho = 1.8$ nm, which are equivalent to hexagonal loops with $\rho = 4.5$ nm and $\rho = 2$ nm. In either case the separation between the two loops in the plane perpendicular to the glide cylinders is chosen as $\Delta x = 12$ nm.

Table II shows that the lifetime of the elastically confined configuration depends strongly on the loop size and temperature. For $\rho = 4.5$ nm the pair of loops is effectively unable to escape from the elastically confined state, as the lifetime is dominated by the escape barrier of $\approx 10$ eV. On the other hand, the lifetime of an elastically confined configuration involving smaller loops $\rho = 2$ nm is comparable with experimental timescales even at low temperatures.

The specific form of configuration of interacting dislocation loops is found to have a most significant effect on its lifetime. An approximation where the loops are treated as pure prismatic objects underestimates the escape time in comparison with the case of freely rotating loops by several orders of magnitude. In contrast to that, the pair of loop with the orientation of their habit planes fixed at $30^\circ$ has a significantly longer lifetime in comparison with a freely rotating pair of loops, and it only breaks apart at relatively high temperatures. This offers a possible explanation for why the experimentally observed rafts of loops remain stable over an appreciable temperature range, while a simple estimate based on the purely prismatic picture of interacting loops predicts much shorter lifetimes [1]. The loop habit plane reorientation not only changes the barrier that the system needs to overcome in order to escape, but most importantly it strongly lowers the curvature of the potential energy barrier, see Fig. 6, hence further increasing the lifetime of an elastically confined configuration by several orders of magnitude.

| Conditions | Pure prismatic | Lowest PES | 30° fixed tilt |
|------------|---------------|------------|----------------|
| $\rho = 4.5$ nm, $T = 200$ K | $\sim 10^{59}$ yr | $\sim 10^{38}$ yr | $\sim 10^{38}$ yr |
| $\rho = 4.5$ nm, $T = 600$ K | $\sim 10^{57}$ yr | $\sim 10^{37}$ yr | $\sim 10^{37}$ yr |
| $\rho = 2.0$ nm, $T = 200$ K | 4 s | 18 min | 3 yr |
| $\rho = 2.0$ nm, $T = 300$ K | 4 s | 0.5 s | 5 min |
| $\rho = 2.0$ nm, $T = 400$ K | 0.2 s | 6 ms | 0.5 s |
| $\rho = 2.0$ nm, $T = 500$ K | 0.02 ms | 0.5 ms | 10 ms |
| $\rho = 2.0$ nm, $T = 600$ K | 0.005 ms | 0.1 ms | 0.8 ms |

The stochastic motion of prismatic dislocation loops dif- fusing in the glide direction is successfully simulated using dislocation dynamics that also includes the stochastic thermal forces treated using the Langevin equation formalism. The dependence of the diffusion coefficient of a dislocation loop on temperature is consistent with molecular dynamics simulations.

Reactions involving interacting dislocation loops, including loop coalescence, repulsion, and the formation of an elastically confined pairs of loops, are well reproduced using the stochastic dislocation dynamics framework proposed above. The internal degrees of freedom of interacting loops result in the formation of complex potential energy landscape of states with distinctly tilted loop habit planes, separated by potential barriers. The addition of thermal energy through stochastic Langevin forces acting on dislocation lines enables interacting loops to switch between the tilted configurations, allowing the system to explore the entire energy landscape of excited states.

In comparison to the purely prismatic case of interacting loops first explored by Foreman and Eshelby [64], the reorienta- tion of the habit plane of interacting loops is found to strongly affect the rates of reactions resulting in the elastic confinement of loops. For one, the potential barrier for the elastic trapping a loop approaching another loop from a distance is strongly reduced, making it much more likely for loops to form elastically trapped configurations. Second, the lifetime of the elastically confined state increases by several orders of magnitude, bringing the estimated lifetime into broad agreement with experimental observations. The habit plane reorientation effect highlights the pivotal significance of including internal degrees of freedom of loops in the treatment of microstructural evolution, to achieve a physically consistent description of dynamics of complex dislocation microstructures.

V. CONCLUSION

The stochastic motion of prismatic dislocation loops dif- fusing in the glide direction is successfully simulated using dislocation dynamics that also includes the stochastic thermal forces treated using the Langevin equation formalism. The dependence of the diffusion coefficient of a dislocation loop on temperature is consistent with molecular dynamics simulations.

Reactions involving interacting dislocation loops, including loop coalescence, repulsion, and the formation of an elastically confined pairs of loops, are well reproduced using the stochastic dislocation dynamics framework proposed above. The internal degrees of freedom of interacting loops result in the formation of complex potential energy landscape of states with distinctly tilted loop habit planes, separated by potential barriers. The addition of thermal energy through stochastic Langevin forces acting on dislocation lines enables interacting loops to switch between the tilted configurations, allowing the system to explore the entire energy landscape of excited states.

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APPENDIX A: RELATION BETWEEN THE INTEGRATION TIME STEP AND SEGMENT LENGTH

Here we give an estimate to the maximum time step that can be used for integrating the coupled stochastic equations of motion describing a fluctuating dislocation. We assume that the stochastic force acting on a dislocation node is greater than elastic forces, and that the adjacent segments have length $\Delta l$:

$$B_{\text{adj}}(t) \approx f_n^0 = \sqrt{\frac{2k_B T B}{\Delta l \Delta t}} N(0, 1). \quad (A1)$$

We continue by making a conservative assumption that the nodal positions are updated according to Euler’s scheme, and that the randomly generated number $N(0, 1)$ is equal to three. For the sake of numerical stability in updating the dislocation network, we assert that the node should not move by more than a fraction $q \leq 1$ of the segment length, leading to $\tilde{z}_n(t) \Delta t \leq q \Delta l$. We then arrive at a cubic relation between step time and segment length:

$$\Delta t \leq \frac{q^3(\Delta l)^3 B}{18k_B T}. \quad (A2)$$

Using values of $q = 0.1$, $B = 0.08$ MPa ns, $T = 300$ K, and $\Delta l = 10$ Å, we arrive at a maximum allowed time step $\Delta t = 11$ fs. Similarly, for $\Delta l = 30$ Å we find $\Delta t = 290$ fs. Hence the stochastic simulations of coarsely resolved dislocation networks can proceed with larger time steps than the matching atomistic simulations.

APPENDIX B: POTENTIAL ENERGY SURFACE OF THE LOOP PAIR

Given a vector $z$ containing all the node positions, we define a set of local energy minima that depend parametrically on the distance between the loops:

$$V_S(s) = \{W^{\text{bulk}}(z) \mid z \in Z \text{ is a local minimum}\}, \quad (B1)$$

where

$$Z = \{ z \mid z^{(2)}_{\text{COP}} - s^{(1)}_{\text{COP}} = s \} \quad (B2)$$

refers to the set of nodal positions for which the separation between the loops is $s$. In other words, the loop separation is held constant, whereas the remaining internal degrees of freedom are varied to find local potential energy minima.

The potential energy surfaces defined by Eqs. (B1) and (B2) are found by the numerical minimization of energy of a simplified system of two loops. The internal degrees of freedom are reduced to tilting modes only, leading to the following parametrization of the dislocation loop:

$$r(\psi) = \begin{pmatrix} \rho \cos \psi \\ \rho \sin \psi \\ u \sin \psi + v \cos \psi \end{pmatrix}. \quad (B3)$$

where $\psi \in [0, 2\pi)$ is the parametrization variable, and $u \in \mathbb{R}$ and $v \in \mathbb{R}$ are tilting amplitudes. The normal vector of the parametrization is independent of $\psi$ and is free to point in any direction, while the loop relaxation volume is constant as $\Omega_{rel} = b \cdot A \equiv b \pi \rho^2$. Thus the loop habit plane may tilt freely within the glide cylinder. The above parametrization may also be used to include tilting in the dipole-tensor approximation [35], though for carrying out a dynamic simulation one would also need an approximate analytic expression for the self-energy of a mixed ellipsoidal loop. Some expressions suitable for this purpose are already available in literature [68,69], but they are relatively limited in comparison with the general case addressed here.

The total potential energy of interacting loops is computed using the nonsingular de Wit formula [39], including the core energy (3). The energy has multiple stationary points at a given separation $s$, which are not trivially identified. Here the total energy was minimized over tilt amplitudes of both loops ($u^{(1)}$, $v^{(1)}$, $u^{(2)}$, $v^{(2)}$) using the BFGS [70] implementation in SCIPY [71] for a broad range of initial tilt configurations. While this approach does not consistently identify all the stationary points, it still gives a qualitative overview of the potential energy landscape.

All the energy minima identified in this way are shown in Fig. 6. Multiple potential energy branches belonging to distinct tilt configurations are found. Note that transitions at a crossing may involve a significant change in tilting, and thus would involve a transition over a large energy barrier.

APPENDIX C: LANGEVIN EQUATION OF MOTION FOR THE LOOP-LOOP SEPARATION

Assuming that the relaxation of internal loop degrees of freedom occurs on a much shorter timescale than the COP diffusion, the adiabatic equation of motion for the individual loop COP is derived from the two equations of motion for the loops:

$$B_{\text{COP}}^{(1)} = -\frac{1}{L^{(1)}} \frac{\partial V(s)}{\partial s_{\text{COP}}} + F_s^{(1)},$$
$$B_{\text{COP}}^{(2)} = -\frac{1}{L^{(2)}} \frac{\partial V(s)}{\partial s_{\text{COP}}} + F_s^{(2)},$$

where $V \in V_S$ is a branch of the PES, and the total stochastic force $F_s^{(i)}$ with strength $\sigma_{\text{COP}}^{(i)}$ acting on loop $i$ is derived following Sec. II using

$$\langle F_s^{(i)}(t) F_s^{(i)}(t') \rangle = \frac{\sigma_{\text{COP}}^{(i)}^2}{L^{(i)}} \delta(t - t'),$$

leading to $\sigma_{\text{COP}}^{(i)} = \sigma_{\text{COP}} / \sqrt{L^{(i)}}$. The total energy derivative is evaluated using the chain rule with $s = z^{(2)}_{\text{COP}} - z^{(1)}_{\text{COP}}$, and the two equations of motion (C1) are subtracted, resulting in

$$B_s = -V'(s) \left( \frac{1}{L^{(1)}} + \frac{1}{L^{(2)}} \right) + F_s^{(2)} - F_s^{(1)}.$$ 

The equation of motion (C3) simplifies further for the case $L^{(1)} = L^{(2)} = L$: 

$$BLs = -2V'(s) + F_s,$$ 

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where $F_i$ is the net stochastic force with standard deviation $\sigma = \alpha \sqrt{2L}$, following the sum theorem of Gaussian distributed variables. Note that the expectation value of the loop velocity over independent trajectories is temperature independent as the stochastic force has zero mean:

$$\langle \dot{s} \rangle = -\frac{2}{BL} V'(s).$$

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