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Md Mijanur Rahman, Fedwa El-Mellouhi, Othmane Bouhali, Charlotte Becquart, Normand Mousseau. Pressure effect on diffusion of carbon at the $85.91\circ 100$ symmetric tilt grain boundary of α -iron. Physical Review Materials, 2021, Physical Review Materials, 5 (4), pp.043605. 10.1103/physrevmaterials.5.043605. hal-03256509

HAL Id: hal-03256509 https://hal.univ-lille.fr/hal-03256509v1

Submitted on 10 Jun2021

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Pressure effect on diffusion of carbon at the $85.91^{\circ} < 100 >$ symmetric tilt grain boundary of α -iron

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(Dated: March 31, 2021)

The diffusion mechanism of carbon in iron plays a vital role in carburization processes, steel fabrication and metal dusting corrosion. In this work, using the kinetic activation-relaxation technique (k-ART), an off-lattice kinetic Monte Carlo algorithm with on-the-fly catalog building that allows to obtain diffusion properties over large time scales taking full account of chemical and elastic effects coupled with an EAM potential, we investigate the effect of pressure on the diffusion properties of carbon in $85.91^{\circ} < 100 >$ symmetric tilt grain boundaries (GB) of α -iron up to a pressure of 12 kbar at a single temperature of 600K. We find that while the effect of pressure can strongly modify the C stability and diffusivity in the GB in ways that depend closely on the local environment and the nature of the deformation: isotropic and uniaxial pressure can lead to opposite and non-monotonous effects regarding segregation energy and activation barriers. These observations are relevant to understanding of the evolution of heterogeneous materials, where variations of local pressure can alter the carbon diffusion across the material.

INTRODUCTION I.

Metal dusting is a severe form of corrosion causing a catastrophic degradation of metals and alloys at high temperature in carbon-supersaturated gaseous environments. Engineering alloys such as low-alloy and stainless steels, as well as heat-resisting iron-, nickel-, and cobaltbase alloys catalyse gas processes that release carbon, leading to the formation of deposits at the surface of the alloy and, over time, to failure. In petrochemical plants and gas-to-liquid (GTL) production facilities, metal dusting has been observed in steel equipment. The so-called Fischer-Tropsch synthesis deposits carbon not only on the catalysts but also on the steel equipment, leading to dusting corrosion. The onset of metal dusting for ironbase alloys is initiated with saturation of the alloy matrix with carbon/carbides, usually in a localized manner, a process called carburisation. In our earlier works we focused on carburisation by characterising reaction of iron with CO as well as the diffusion and segregation of carbon within the grain boundaries of the metal.¹⁻¹⁰

In this work we aim to turn our attention towards realistic scenarios that occur at various stages of dusting corrosion, drawing comparisons with experiments whenever possible, focusing more particularly on the effect of pressure on C diffusion in Fe, particularly at grain boundaries (GBs). The diffusion of carbon interstitial atoms around GBs in iron has long been an area of interest for researchers due to its impact on various mechanical properties of steel such as corrosion resistance, and embrittlement, but also at it serves as a rich system for modelling and simulation.^{11,12} Over the years, these mechanisms have been computationally studied at ambient pressure,

i.e. the lattice and atomic positions are both relaxed to obtain the atomistic parameters at zero pressure.^{13–16} While limited information is available regarding the effect of pressure on carbon diffusion in iron through GBs, internal or external pressures can build up in the grains due to local deformation. These may reach a magnitude of a few GPa,^{17–19} inducing problems such as structural instability, stress corrosion cracking, distortion, etc.^{20–22} It is thus necessary to have a clear description of the effects of pressure on defect diffusion in order to understand the behavior of heterogeneous materials such as carburized steels at the atomic scale.

Because of the microscopic nature of these processes, atomistic simulations are needed to provide an understanding of the details controlling these segregation and diffusion mechanisms that are difficult to obtain from experiments. While standard simulation tools, such as molecular dynamics, offer significant information on atomistic details of these diffusion mechanisms, they are often unable to reach the extended time scale over which many of these processes take place. Standard kinetic Monte Carlo (KMC) simulations provide a solution to reaching long-time dynamics, but, they are constrained to lattice-based displacements that fail to capture the full diversity of diffusion mechanisms as well as elastic deformations, especially around extended defects such as GB.²³ Over the years, numerous improvements have been proposed to standard KMC simulations, in order to overcome these limitations leading to a new interest in the fundamental mechanisms associated with defect diffusion in materials. The recent development of onthe-fly off-lattice kinetic Monte Carlo methods such as the kinetic activation-relaxation technique (k-ART)²⁴⁻²⁶

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lifts those limitations and allows us to map these processes in details. K-ART is an off-lattice kinetic Monte Carlo method with on-the fly catalog building capabilities which enables to properly describe diffusion mechanism over experimentally-relevant time scales. It enables to reach simulated times many orders of magnitude longer those reachable by MD simulation while incorporating exact elastic effects and identifying the atomistic details of diffusion mechanisms at every step.

In this study, using k-ART we investigate the effect of pressure on the diffusion mechanism C atom within a GB of α -iron. More precisely, we explore the energy landscape and kinetic pathways. Then, diffusion coefficients are calculated to quantify the effect of stress on diffusivity. The paper is organized as follows: the methodology, including a brief overview of k-ART, the model employed and the computational details, is presented in section II. Then, simulation results are presented and discussed in section III and IV respectively. Finally, the conclusion is given in section V.

II. METHODOLOGY

A. Kinetic-Activation Relaxation Technique (k-ART)

The kinetic activation relaxation technique (k-ART)²⁴⁻²⁶ is an off-lattice kinetic Monte Carlo (KMC) method with on-the-fly catalog building. K-ART is built around three tools: (i) NAUTY,²⁷ which allows a topological characterization of the local environment surrounding each active atom; (ii) the activation-relaxation technique (ART nouveau or ARTn),²⁸⁻³⁰ an open-ended saddle-search methods for exploring the local energy landscape and (iii) a KMC algorithm,³¹ that evaluates the time between two steps and selects the event. The basic principles are briefly described below:

At the beginning of each KMC step, the local environment surrounding each atom is characterized by their local topology evaluated with NAUTY, a topological analysis library developed by McKay.²⁷ Graphs are generated by considering a sphere with a predefined radius (6 Å for the current system) around each atom in the system that contains around 75 atoms and drawing edges between all atoms within a cut-off distance of 2.7 Å of each other, a value that ensure that mostly only first-neighbor atoms are included. This connectivity graph is sent to NAUTY which returns a unique identifier characteristic of its automorphic group.

If the topology is known, events associated with it are recovered from the catalog and placed in a KMC tree; otherwise, 100 ARTn searches are launched to identify the events associated with this topology. The ARTn search takes place in three steps: (i) first, the system is pushed in an arbitrary direction until the lowest eigenvalue of the Hessian matrix, determined using the Lanczos algorithm, becomes negative to confirm that the system is out of the harmonic well; (ii) then the system is pushed along the direction of the negative curvature and the force is minimized in the hyperplane perpendicular to this direction after each push until the total force becomes lower then a predefined threshold value, indicate that a first order saddle point is reached; and (iii) the system is then push over the saddle point and relaxed into a new minimum.

Once the catalog is fully updated and the tree is completed for the current atomistic configuration, generic events are ordered according to their rate, defined as:

$$\Gamma_i = \nu_0 e^{-\frac{E_b}{k_B T}} \tag{1}$$

where ν_0 is a fixed prefactor which is fixed at 10^{13} Hz and E_b , the activation energy for event *i* defined as the energy difference between the transition state and the initial minimum.^{32–34} Once the event tree is completed, the lowest-energy barrier events that make up to 99.99 % of the rate are fully reconstructed and reconverged into specific events to ensure that elastic and configurational effects are exactly taken into consideration. When this is done, specific rates and the overall KMC timestep are evaluated again, with the refined barriers, and an event is chosen according to standard KMC algorithm.³¹ The elapsed time *t* is computed according to a Poisson distribution as:

$$t = -\frac{\ln \mu}{\sum_{i} \Gamma_{i}} \tag{2}$$

where μ is the random number uniformly distributed between 0 and 1.

To sum up, k-ART makes use of a unique topology classification coupled with an unbiased, open-ended search for saddle points, while considering short- and long-range interactions due to elastic effects. An extensive catalog of the events are created on-the-fly which speeds up the simulation as the system evolves. In the past, k-ART has been used in various systems to describe diffusion of point defects in metals and semiconductors and more complex systems with grain boundaries and amorphous silicon.^{9,35,36}

B. Handling flickering states

KMC simulations can become inefficient in the presence of low energy barriers that dominate the kinetics, as each step is then associated with a very small clock increase. In some cases, these events are flickers, a set of states, separated by low-energy barriers deep in a energy basin and that do not contribute to the evolution of the system. To avoid getting trapped by flickers, we use the basin auto constructing mean rate method (bac-MRM)^{26,37} to solve analytically the average residence time for the in-basin states. This method computes onthe-fly a statistically exact analytic solution of the con-



FIG. 1: Atomic arrangement representation of the 85.91° < 100 > tilt GB in BCC Fe after 1 ns of initial relaxation using molecular dynamics (MD) at zero pressure and temperature 600 K prior to starting k-ART simulations. Black circles represent Fe atoms having a crystalline BCC local environment, while green circles represent atoms having a non-bcc environment

surrounding the GB. The GB rotation axes < 100 > are parallel to the x axis out of planes, while GB planes are perpendicular to the z axis

nected flickering states and their escape rate as the energy landscape is explored, at the cost of specific trajectories. The bac-MRM is applied to all states connected by barriers and energy difference lower than a predefined threshold (0.5 eV is used in this study).

C. Interatomic potential

In order to access relevant length and time scales, we employ an empirical force-field to describe the system. The Fe-Fe interactions are handled by the Embedded Atom Method (EAM) interatomic potential developed by Ackland and Mendelev³⁸ while the Fe-C interaction part was developed by Becquart and collaborators.^{39,40} The combined potential provides a good agreement with DFT calculations in bulk system.⁸ This potential has been used with success to model the effect of the stress field of an edge dislocation on carbon diffusion, the formation of carbon Cottrell atmospheres in bcc-iron as well as the elastic constants of the martensite.^{41–43} For all calculations, k-ART uses the LAMMPS library to compute the forces and energies.^{44,45}

D. Simulation details

Two simulation boxes are used in this work. The first one consists in a perfect cubic BCC bulk crystal containing 2000 iron atoms. The second box is a BCC system containing 25537 atoms featuring a $85.91^{\circ} < 100 >$ tilt GB with dimensions of 34.332 Å $\times 113.090$ Å $\times 76.595$ Å. To allow periodic boundary conditions in all directions,

the box contains two GBs separated by a distance equal to half the box size in the z direction. The $85.91^{\circ} < 100 >$ structure is a general GB with GB energy per unit area, $E_{GB} = 37.118 \ meV/Å^2$. This GB Fe system was chosen because of its structural stability, and the presence of a number of sites where C can be trapped, allowing us to focus on the effect of deformation impurity diffusion. An atomistic representation of the GB is shown in Fig.1. To determine the appropriate simulation volume, the system boxes, with a C interstitial, are equilibrated at the targeted pressure at 600 K temperature by perfoming NPT ensemble with LAMMPS. The volume is then fixed for k-ART simulations.

For each simulation, the ground state (GS) energy is defined as the lowest energy minimum identified during the run, E_{GS} . The activation energy for an event is defined as the energy difference between the initial minimum E_{min} and the saddle E_{sad} (i.e., the barrier crossed between two adjacent minima) or

$$E_b = E_{sad} - E_{min}.\tag{3}$$

The effective barrier is defined as the total barrier crossed when jumping from one GS to another GS.

The square displacement (SD) is computed as usual,

$$SD = \sum_{i}^{N} (r_i(t_n) - r_i(0))^2, \qquad (4)$$

where N is the number of particles and $r_i(t_n)$, the position of atom i at KMC step n.

To ensure sufficient statistics, the diffusion coefficient of carbon interstitial atom calculated using the embedded discrete time Markov chain (EDTMC) described as follows:

- 1. choose an initial state i;
- 2. generate a uniform random number u ϵ (0,1];
- 3. select the next event j probabilistically such that $\sum_{k}^{j-1} p_{ik} < u \leq \sum_{k}^{j} p_{ik}$, where, p_{ik} is the probability to jump from state i to state k;
- 4. update the total selection number of state j by adding 1;
- 5. update the current state $j \rightarrow i$;
- 6. return to step ii.

EDTMC uses the transition matrix obtained from k-ART and that includes all the structurally different sites in which C visited and the mapped connectivity between the sites and their associated relative transition rates. For all coefficient diffusion results presented here, we run 600 million EDMC steps to obtain the relative frequency distribution $\overline{\Pi}$:

$$\Pi = \begin{bmatrix} \pi_1 & \pi_2 \dots & \pi_m \end{bmatrix}$$
(5)

where, π_i represents the proportion of times that site i is selected and m is the total number of structurally different sites in the system, for instance, in the case of perfect bulk crystal there are structurally three different sites directed along x, y or z (shown in supplementary Fig. S1).

After obtaining the relative frequency distribution, we then calculate the average time < t > and average square displacement < SD >:

$$\langle t \rangle = \sum_{i=1}^{m} \pi_i \tau_i \tag{6}$$

$$\langle SD_l \rangle = \sum_{i=1}^{m} \sum_{j=1}^{m} \pi_i p_{ij} (\lambda_{ij}^l)^2$$
 (7)

with l = x, y, z and,

$$\langle SD \rangle = \langle SD_x \rangle + \langle SD_y \rangle + \langle SD_z \rangle \quad (8)$$

where, τ_i is the mean residence time at site i, p_{ij} is the transition probability from site i to site j and $p_{ij} = 0$; $\forall i=j$. λ_{ij}^l correspond to the l = x, y, z components of the displacement vectors for the jump form site i to site j respectively.

The total diffusion coefficient is calculated as:

$$D = \frac{\langle SD \rangle}{2n_d \langle t \rangle} \tag{9}$$

where, n_d is the dimensionality of the motion (1-3).

III. RESULTS

A. Carbon in bulk iron

First we analyze the effect of pressure on the evolution of a single C atom in crystalline α -Fe. In the ground state, the C atom occupies the octahedral interstitial position, located in the middle of one of the three edges of the cubic unit cell (shown in the supplementary Fig.S1); it migrates, following its dominant pathway, by going through an adjacent tetrahedral site that represents the transition state. With respect to the orientation of the two closest Fe atoms closest, three octahedral variants directed along x, y or z can be identified. These are energetically equivalent in the perfect BCC lattice, but split into two non-equivalent types under uniaxial deformation: a distortion along x axis divides the octahedral sites into O_x and $O_y \equiv O_z$ (shown in supplementary Fig. S2).

Recently, Tchitchekova *et al.*, using a method called Linear Combination of Stress States (LinCoSS), evaluated the effects of simple-heterogeneous uniaxial and shear stresses on the migration energies of C in bulk Fe.⁴² In this section, we reproduce this result for a C in bulk Fe



FIG. 2: (a) Enthalpy variation for a single carbon in 2000-atom Fe bulk crystal (blue circles) and crystal lattice parameter (red circles) as a function of isotropic pressure. (b) Variation of the activation energies (red circles) and diffusion coefficients (blue squares) as a function of pressure for a single carbon atom in the same system. Diffusion coefficient is calculated with EDTMC and Eq. 9.

with a thermally-adjusted density corresponding to 600 K, but also consider isotropic deformations and extract the diffusion coefficient for each of these deformations.

The variation of the crystalline lattice parameter and of the system's enthalpy as a function of isotropic pressure are shown in Fig.2a while the evolution of the activation energy for an interstitial C atom as it jumps from one octahedral site to a nearest octahedral site and the diffusivity of C as a function of pressure are plotted in Fig.2b. Both the diffusion activation energy and the total enthalpy show a positive correlation with isotropic pressure over the full range considered here. The variation of activation energies E_b and that of the total enthalphy follows a quadratic form with the values for the constant, linear and quadratic parameters listed in the supplementary Table S1.

The effects of uniaxial pressure (along x axis) and biaxial pressure (along y and z axes) are shown in Fig.(3-4). Fig. 3a and Fig. 3b present the activation energy of for an interstitial C atom as it jumps from one octahedral site to a nearest octahedral site as a function of uniaxial pressure and biaxial pressure respectively. Activation energies from O_x to $O_y (\equiv O_z)$, from $O_y (\equiv O_z)$ to O_x and from O_y to O_z follow a quadratic curve (values of the linear parameters and quadratic parameter listed in



FIG. 3: Variation of the activation energy for C diffusion in the bulk iron at 600 K as a function of (a) uniaxial and (b) under biaxial stress. The red square, blue circle, and green triangle represent the carbon diffusion between octahedral variants $O_x \to O_y, O_y \to O_x$ and $O_y \to O_z$ respectively.

the supplementary Table S1). Fig. 4a and Fig. 4b show the C diffusion coefficient at temperature 600K at each directions as a function of uniaxial and biaxial pressure respectively. Calculated activation volume V^{act} , activation barrier E^{act} and diffusion prefactor D_0 are listed in the supplementary Table S2. We find that $D_y (\equiv D_z) < D_x$ under uniaxial compression and $D_y (\equiv D_z) > D_x$ under uniaxial expansion with the the ratio D_x/D_y is approximately inversed when the pressure changes sign (see Fig.4c).

Under uniaxial compression along the x-axis, the O_y and O_z octahedral sites are more energetically stable sites than the O_x octahedral sites by as much as 0.132 eV at 24 kbar (shown in supplementary Fig. S2), with barriers going from O_y (O_z) to O_z (O_y) lower by as much as 0.051 eV as compared with O_y (O_z) to O_x . Consequently the lowest-energy diffusion pathway for C interstitial goes from O_y to O_z to O_y , channeling the C atom along the contracted axis. Therefore, the axial compression promotes axial diffusion. On the contrary, axial expansion promotes transverse in-plane diffusion along the chain (O_y or O_z) $\rightarrow O_x \rightarrow (O_y$ or O_z) $\rightarrow O_x$ (detailed C diffusion trajectories obtained with k-ART under uniaxial deformation are shown in supplementary Fig.S3).

Since the contraction along x direction is equivalent to expansion along y and z direction, biaxial deformation



FIG. 4: C diffusion coefficient in the bulk crystal at 600 K as a function of (a) uniaxial stress, (b) under biaxial and (c) ratio D_x/D_y . All the values are calculated using EDTMC and Eq. 9.

just reverse the tendancy : biaxial compression promotes C diffusion along the y-z in plane while the biaxial expansion promotes x-channeling diffusion.

B. Carbon in the Grain boundary

To relieve the pressure created on the crystalline network, interstitial impurities such as C tend to segregate towards defects, such as grain boundaries (GB), where there is more free volume than on the crystalline bulk side. This segregation propensity depends, of course, on specific binding sites at the GB and is characterized by the segregation energy defined as the difference between the energy of the system with C in a particular site and



FIG. 5: (a) Atomistic representation of three most frequent diffusion mechanisms of carbon atom at the dislocation core in GB at zero pressure. The orange, blue and black lines indicate mechanisms M_1 , M_2 and M_3 respectively. M_1 and M_2 can move the C atom into a new final configuration (S'_0) (equivalent to the initial configuration (S_0) with

the impurity jumping by one lattice spacing along the x direction); M3 brings the C atom back to its initial configuration (S_0) . It does not contribute to the net diffusion. (b) The minimum energy pathway for the different mechanisms is shown, where the circles and crosses represent minimum and saddle points, respectively. E_{seg} is the relative energy with respect to the bulk ground state for C.

 TABLE I: Values of the coordination number, shortest Fe-C bond length and segregation energies of four most stable sites at the GB region at different pressures.

Site		0 kbar		Isotropic pressure				Pressure applied along x-axis				Pressure applied y and z axes			
	Coordi-			12 kbar		-12 kbar		12 kbar		-12 kbar		12 kbar		-12 kbar	
	nation	d_{Fe-C}^{min}	E_{seg}	d_{Fe-C}^{min}	E_{seg}	d_{Fe-C}^{min}	E_{seg}	d_{Fe-C}^{min}	E_{seg}	d_{Fe-C}^{min}	E_{seg}	d_{Fe-C}^{min}	E_{seg}	d_{Fe-C}^{min}	E_{seg}
	No.	(Å)	(eV)	(Å)	(eV)	(Å)	(eV)	(Å)	(eV)	(Å)	(eV)	(Å)	(eV)	(Å)	(eV)
S_0	8	2.029	-0.745	2.028	-0.822	2.031	-0.660	2.023	-0.740	2.033	-0.735	2.032	-0.814	2.010	-0.673
S_1	5	1.764	-0.20	1.763	-0.220	1.765	-0.174	1.768	-0.190	1.763	-0.206	1.763	-0.230	1.766	-0.164
S_2	5	1.775	-0.164	1.774	-0.198	1.775	-0.123	1.776	-0.120	1.773	-0.178	1.772	-0.214	1.775	-0.080
S_3	5	1.819	-0.057	1.818	-0.089	1.820	-0.017	1.813	0.013	1.823	-0.091	1.823	-0.126	1.812	0.028

that in the octahedral site in the crystal side. A negative value of segregation energy means that the grain boundary site is energetically more favorable than the bulk site and signals a general flux from the bulk to these sites at the GB.

As demonstrated in our previous work,⁹ the disorder associated with GB generates a variety of atomic environments, facilitating the formation of sites more energetically favorable to the presence of carbon than interstitial crystalline sites. Depending on the specific nature of the interface between the grains, these favorable sites could be connected to accelerate or to slow down carbon diffusion as compared to the bulk. In the specific case studied here, we add that, segregation energies as well as related C migration barriers are reported in the current work differ from the value found in our previous work⁹ as they are identified differently: sites identified in the previous work were generated kinetically through C migration from the bulk to the GB⁹ while a direct extensive search for C sites, with simulations launched from different GB sites, was performed for the current work to identify the thermodynamically preferential sites. While this approach leads to different diffusion pathways, it does not affect the validity of our previous analysis.

K-ART search for C diffusion pathways at the 85.91° GB finds 6 different connected sites, with a segregation energy ranging from -0.745 eV to 0.180 eV at zero pressure. Fig.5 represents the four most energetically favor-



FIG. 6: Segregation energy as a function of pressure for a C atom at the 85.91° GB:(a) under isotropic pressure,
(b) pressure along the x-direction and (c) pressure applied along the plane perpendicular to the GB plane.

Red square, blue circle, green triangle and violet diamond are correspond to carbon at the site S_0, S_1, S_2

and S_3 respectively.

able sites for carbon segregation (S_0 to S_3), along with their segregation energy, measured with respect to the bulk ground state for C at zero pressure, and activation energies for diffusion between these sites. Further details, including shortest Fe-C bond length, coordination number and segregated energies under various static deformations, are listed in Table I. The coordination number is defined as the number of neighboring Fe atoms within a cut-off distance of 2.47 Å from the C atom. As previously found for this specific grain boundary,⁹ even the highest energy states along the dominant diffusion pathways at the GB are lower than the bulk ground state, which indicates a strong segregation tendency of C atoms



FIG. 7: Variation of the effective barrier of the diffusion mechanism M_1 , M_2 and M_3 as a function of pressure: (a) under isotropic pressure, (b) pressure along the x-direction and (c) pressure applied along the plane perpendicular to the GB plane.

to the GB. At the GB, the lowest-energy state at the GB is labeled S_0 (situated at the center of the dislocation core of the GB), with a segregation energy of -0.745 eV. Among the four lowest-energy states (S_0 to S_3), S_0 shows the highest coordination number, with the longest Fe-C nearest neighbor distance associated with a low-stress environment as compared with the bulk site (Table I).

Fig.6(a-c) represents the variation of the segregation energies for these four lowest-energy sites as a function of isotropic pressure as well as uniaxial (along x-axis: direction of GB) and biaxial (along y and z axes: plane perpendicular to the GB direction) deformations. Within a range of -12 to 12 kbar, the segregation energy of the C at each binding site tends to evolve linearly, although the slope varies with specific sites. The displayed relative



FIG. 8: (a) Variation of the C diffusion coefficient at the GB as a function of pressure: red circle, blue square, and green triangle represent the applied pressure isotropic, along x-axis, and along y and z axes respectively. (b) Arrhenius plot of the diffusivity of C in GB: red, blue and green line represent for the applied pressure isotropic, along x-axis, and along y and z axes respectively. Black line refers to stress-free conditions.

(c) Variation of the diffusion coefficient of C as a function of isotropic pressure: red circle and blue square represent C at the GB and crystalline bulk respectively. Results are presented here are calculated using Eq. 9.

energy shows that all C binding sites at the GB become more stable as compared to the interstitial octahedral sites of the crystal side under isotropic compression. As the pressure is increased from 0 to 12 kbar, the segregation energy of C at the most stable site (S_0) increases in stability by 0.077 eV, whereas the segregation energy of C at sites S_1 , S_2 and S_3 increases in stability by 0.02, 0.034, and 0.032 eV respectively. A similar behavior is observed under biaxial pressure. However, sites S_1 , S_2 , and S_3 are even more favored, as compared to the case of isotropic pressure, with increasing the pressure (values listed in table I). Under biaxial compression, the two Fe nearest-neighbors are displaced along the x direction, leaving more space for the C atom. Interestingly, the segregation energy of sites S_0 remains unaffected by uniaxial deformation, since the radius of the dislocation core is unaffected by the deformation. Hence, the increment (or decrement) in the total energy of the C at site S_0 and the octahedral site in the crystal side are almost same under uniaxial pressure. By contrast to isotropic and biaxial pressure the sites S_1 , S_2 and S_3 are destabilized under uniaxial compression.

After characterizing the change in stability of the dominant segregation sites under pressure, we now turn to the effect of pressure on the C activation energies and diffusion pathways at the 85.91° GB. C diffusion at the GB is complex as compared with that of the crystalline bulk. As demonstrated in a previous work,⁹ at zero pressure, C diffusion occurs through several pathways that move through the most favorable GB sites (the energy details of all available events and the histogram of the activation energy barrier for selected events at 600 K for a single C at GB during the 12000 KMC steps are shown in the supplementary Fig.S3 and Fig.S4 respectively). Fig.5 provides an atomistic representation of the three most frequent C diffusion local pathways at the GB that we label as M_1 , M_2 and M_3 . Each pathways is a two steps process: C diffuses from the ground state (site S_0) to a metastable state $(S_1, S_2 \text{ or } S_3)$, then from this metastable state to a new GS (site $S'_0 = S_0$). While, after these two jumps, M_1 and M_2 can reach a new lower-energy state, dubbed S'_0 , M_3 systematically brings the C back to its initial position when following the lowest-energy pathways and, as such, does not contribute to diffusion. M_1 exhibits the lowest maximal energy barrier, which makes mechanism M_1 the most probable. At zero pressure, the first step of mechanism M_1 requires crossing an energy barrier of 0.764 eV to reach S_1 . From there the C then crosses a 0.216 eV barrier to complete the migration into a new S_0 site. The second (M_2) and third (M_3) lowenergy diffusion mechanism are associated with effective barriers of 0.840 and 0.877 eV respectively compared to 0.810 eV in the bulk.

The variation of the effective energy barriers for the three dominant pathways at the GB along with their selection's frequency as a function of pressure is shown in Fig.7. Within a pressure range -12 to 12 kbar, the effective barrier associated with M_1 , M_2 and M_3 varies linearly with pressure, but at different rate. In the case of isotropic deformation, as the pressure is increased from 0 to 12 kbar, the effective barriers associated with the dominant diffusion mechanism (M_1) increase by 0.111 eV, whereas the effective barriers for mechanism M_2 and M_3 increase by 0.055 and 0.045 eV respectively. Since, the energy gap between the pathway's effective barriers narrows as pressure is increased, the less probable mechanisms (such as M_3) become more frequent under isotropic

compression at temperature 600 K. In the case uniaxial deformation, on the contrary, the energy gap between the pathway's effective barriers widen and mechanism M_1 become even more dominant under compression. As the pressure is increased from 0 to 12 kbar the effective barriers associated with the diffusion mechanism M_1 , M_2 and M_3 increase by 0.014, 0.028, and 0.053 eV respectively. By contrast to the isotropic and uniaxial pressure, effective barriers associated with the diffusion mechanism M_3 decreases under biaxial compression. As the pressure is increased from 0 to 12 kbar, the effective barriers associated with the diffusion mechanism (M_3) decrease by 0.003 eV, whereas the effective barriers for mechanism M_1 and M_2 increase by 0.078 and 0.017 eV respectively.

C diffusion at $85.91^{\circ} < 100 >$ tilt GB is largely restricted to the interface and diffusion take place mostly along the x-axis (diffusion trajectories obtained with k-ART at different pressure are shown in supplementary Fig.S6).

To extract a converged diffusion coefficient, we turn, as discussed in the method section, to a Markov chain solution on a transition probability matrix (the size of which ranges from $5 \times 5 - 7 \times 7$ depending of the nature of the deformation) that incorporates the full set of events identified with k-ART, irrespective of their energy barrier. Fig.8a shows the variation of C diffusion coefficient at the GB under isotropic pressure along the x, y and zaxes at temperature 600 K. Fig.8b represents the Arrhenius plot for C diffusivity in GB. The calculated activation volume V^{act} , activation barrier E^{act} and diffusion prefactor D_0 are listed in the supplementary Table S2. We find that it decreases (increases) linearly under lattice compression (expansion), but at different rates. Under lattice compression, as the pressure is increased from 0 to 12 kbar the diffusion coefficient of C at the GB is decreased by $1.39 \times 10^{-13} m^2 s^{-1}$ (net effective barrier increase: 0.068 eV), $5.30 \times 10^{-14} m^2 s^{-1}$ (net effective barrier increase: 0.01 eV) and $1.03 \times 10^{-13} m^2 s^{-1}$ (net effective barrier increase: 0.046 eV) for the isotropic, uniaxial and biaxial pressure respectively. On the other hand, under lattice expansion as the negative value of pressure is increased from 0 to 12 kbar the diffusion coefficient of C at the GB is increased by $4.95 \times 10^{-13} m^2 s^{-1}$ (net effective barrier decrease: 0.067 eV), $1.09 \times 10^{-13} m^2 s^{-1}$ (net effective barrier decrease: 0.020 eV) and 2.15 \times $10^{-13} m^2 s^{-1}$ (net effective barrier decrease: 0.047 eV) for the isotropic, uniaxial and biaxial pressure respectively.

Fig.8c compares the results for a similar set of calculations for the C diffusion coefficient at the GB and in the crystalline bulk under isotropic pressure at 600 K. We find that the applied pressure strongly affects C diffusivity at the GB as compared to bulk. At zero pressure, C diffusion at the GB is faster than in the crystalline bulk: the diffusion coefficient of C at GB is $1.94 \times 10^{-13} m^2 s^{-1}$ (corresponds to an effective barrier 0.772 eV), which is order of magnitude higher as compared with $2.65 \times 10^{-14} m^2 s^{-1}$ (corresponds to a barrier of 0.810 eV) in the crystalline bulk. The diffusivity of C at GB decreases rapidly as compared in bulk when isotropic pressure increases: as the pressure is increased from 0 to 12 kbar the diffusion coefficient of C at GB decreased by $1.39 \times 10^{-14} \ m^2 s^{-1}$ (net effective barrier increase: 0.068 eV) compared with $4.9 \times 10^{-15} \ m^2 s^{-1}$ (net barrier increase: 0.01 eV) in the crystalline bulk.

IV. DISCUSSION

To understand the impact of pressure on impurity diffusion at grain boundaries, we first establish a reference case by considering impurity diffusion in the bulk crystal under isotropic, uniaxial and biaxial deformations. In agreement with simulations done on a system equilibrated at T = 0 K,⁴² we find that the activation energy barrier increases under istropic pressure. We show that this leads to a non-linear reduction in the diffusion coefficient, as should be expected. A similar observation was made in experiments of vacancy-mediated In diffusion in Cu,^{46,47} which suggest that this slowing down, in a dense metallic system is associated with the reduced space available along the diffusion pathway between two stable states.

Uniaxial and biaxial compressions, by breaking symmetry, introduce divergence in activation energy as a function of jump directions for the interstitial C, with some barriers decreasing while others increase under increasing pressure, leading to a non-monotonous effect of pressure on the diffusion coefficient, a phenomenon that does not seem to have been reported previously. This non-monotonous effect can be explained by the fact that, in addition to making the lattice stiffer, increasing pressure raises the formation energy for an interstitial. Along some diffusion directions, the enthalpy at the stable site increases faster than that at the transition state, leading to an effective reduction in the diffusion barrier.

These effects are made more complex at the grain boundary due to an increased richness in the energy land-scape. This explains why results on various GBs differ. For example, Lojkowski *et al*, looking at In diffusion at a Cu grain boundaries, find that diffusivity increases with pressure^{46,47} while it decreases by an order of magnitude for Zn diffusing along tilt GBs in Al as the pressure is raised to 12 kbar.⁴⁸

Recently, Karki *et al.*⁴⁹ performed first-principles simulations, based on density functional theory, on the $\{310\}/[001]$ tilt GB in MgO. They showed that native defects and impurities (Ca, Al, and proton) favorably segregate to the GB with the segregation considerably increasing with pressure. They also remarked that the impurity diffusion is faster at the GB as compared to the bulk as the pressure is increased as they showed that migration energies for host ions and impurities at the grain boundary are smaller than the bulk values, more so at higher pressures with their values being as low as 1.5 eV at 100 GPa compared to the bulk values of 4 eV. Using k-ART, we obtain a more complete picture of the energy landscape of our system, C at the $85.91^{\circ} < 100 >$ tilt GB. We find that, while segregation energy tends to become more favorable to the GB with respect to the bulk for a C interstitial with increasing isotropic pressure, the gain depends on specific sites, with some barely changing moving over a 24 kbar pressure change. Moreover, we observe that the orientation of pressure with respect to the GB changes the result : applying a pressure normal to the GB reduces the available free volume at the GB and, therefore, reduces the segregation energy, while a pressure parallel to the GB plane increases the formation enthalpy in the bulk while preserving a larger free volume, and increasing the energy gain of moving to the GB.

Pressure will also affect the various transition states differently as is shown in our detailed analysis of its effect for the three dominant transition states at the GB. While all three barriers increase linearly under increasing isotropic and normal pressure, albeit with a different slope, this is not the case for pressure applied parallel to the GB plane, which could, at high enough pressure, lead to a change in the dominant diffusion mechanisms and the overall diffusion. This suggests that inhomogeneities in materials, that can create large local pressure fluctuations, will increase the complexity of impurity diffusion and could favor mechanisms that are missed in analyses that do not explore the full landscape but could affect the overall properties of the material. k-ART provides a complete picture about evolution of C within a representative grain boundary under the influence of pressure, thus mimicking realistic working conditions and providing unique long-time description of the evolution of the system. The resulting full energy landscape on the impact of pressure on the C diffusion in iron once combined with surface phenomena can be regarded as one of the key parameters to determine the threshold and peak carburization rates of Fe surfaces.

V. CONCLUSIONS

Using the kinetic activation-relaxation technique, an off-lattice kinetic Monte Carlo method with on-the-fly

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show a richness in the diffusion mechanisms as well as a

More work is required to better characterize this behavior and extract general rules. Clearly, additional simulations are needed to assess how multiple C interstial atoms or other defects such as vacancies and substitutional atoms diffuse at GBs under pressure; simulations are underway. The detailed analysis of the energy landscape provided by k-ART, however, shows that it is necessary to take specific mechanisms into account, with the possibility of developing precise control on the evolution of impurities in complex materials under local and global deformations.

VI. ACKNOWLEDGMENTS

This work has been supported by the Qatar National Research Fund (QNRF) through the National Priorities Research Program and the Canada Research Chairs program and by grants from the Natural Sciences and Engineering Research Council of Canada (NSERC). We are grateful to Calcul Québec/Compute Canada (CQ/CC) for generous allocations of computer resources.

VII. CODE AVAILABILITY

Various ART nouveau implementations are available freely for download from http://normandmousseau.com. The most recent packages for k-ART and ART nouveau are available freely from the authors upon request.

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