Periodized discrete elasticity models for defects in graphene

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Abstract

The cores of edge dislocations, edge dislocation dipoles and edge dislocation loops in planar graphene have been studied by means of periodized discrete elasticity models. To build these models, we have found a way to discretize linear elasticity on a planar hexagonal lattice using combinations of difference operators that do not involve symmetrically all the neighbors of an atom. At zero temperature, dynamically stable cores of edge dislocations may be heptagon-pentagon pairs (glide dislocations) or octagons (shuffle dislocations) depending on the choice of initial configuration. Possible cores of edge dislocation dipoles are vacancies, pentagon-octagon-pentagon divacancies, Stone-Wales defects and 7-5-5-7 defects. While symmetric vacancies, divacancies and 7-5-5-7 defects are dynamically stable, asymmetric vacancies and 5-7-7-5 Stone-Wales defects seem to be unstable.

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I. INTRODUCTION

Graphene\textsuperscript{1,2} and other two dimensional (2D) crystals\textsuperscript{3} have been experimentally observed quite recently. This discovery has led to new physics where quantum relativistic phenomena can be mimicked and tested experimentally in condensed matter physics\textsuperscript{4}. Among striking electronic properties of graphene due to its quantum electrodynamics-like spectrum, there are two chiral quantum Hall effects, minimum quantum conductivity in the limit of vanishing concentration of charge carriers and strong suppression of quantum interference effects. Ballistic transport on submicron distances at room temperature makes graphene a promising material for nanoelectronics\textsuperscript{4}.

Defects in graphene strongly affect its electronic and magnetic properties\textsuperscript{5,6,7}, which may be described by the Dirac equation on curved space\textsuperscript{8}. Irradiation experiments show that pentagon-heptagon pairs (5-7 defects), vacancies, divacancies (5-8-5 defects comprising an octagon and two adjacent pentagons) and adatoms are commonly obtained, but seemingly not Stone-Wales (SW) defects (two adjacent 5-7 defects with opposite Burgers vectors and whose heptagons share one side, briefly 5-7-7-5 defects)\textsuperscript{9}. The far field of 5-7 defects corresponds to that of edge dislocations in elasticity, while the far field of vacancies and divacancies is that of an edge dislocation dipole. Quite recently, experimental observations of edge dislocations on high-quality graphene grown on Ir(111) have been reported\textsuperscript{10}. Studies of defects and their motion are important to assess the mechanical response of graphene at the atomic scale and, as indicated above, its electronic properties.

One common way to describe defects in graphene is to use ab initio calculations. Density functional theory (DFT) has been used to ascertain the magnetic properties of graphene sheets and single-wall nanotubes with vacancies\textsuperscript{5}. Local spin DFT has been used to describe glide and shuffle dislocations in irradiated graphitic structures\textsuperscript{11}. Molecular dynamics (MD) has been used to discuss the stability of nanotubes under tension\textsuperscript{12} and also in the presence of different 5-7 pairs such as 5-7-7-5 and 7-5-5-7 (similar to SW but now the pentagons share one side) defects\textsuperscript{13}. Atomistic Monte Carlo simulations are less costly than MD and have been used in studies of the stability of single graphene sheets\textsuperscript{14}. Classical structural models of graphene may account for chirality effects in nanotubes and allow to assess the impact of the lattice structure on some elastic properties\textsuperscript{15}. However, these classical models lack the ability to generate and move defects. The approach presented in this paper is different.
Relevant defects in graphene are the cores of different edge dislocation and edge dislocation dipoles. Thus we will regularize appropriately linear elasticity on the graphene honeycomb lattice and describe which are the stable cores of different edge dislocations and dipoles. It is well known that cores of dislocations in crystals with covalent bonds are very narrow, so that the elastic field decays quite fast to that given by linear elasticity as the distance to the dislocation point (in 2D the dislocation line is a point) increases. This means that we can regularize linear elasticity on a relatively small hexagonal lattice and impose boundary conditions corresponding to the elastic field of an edge dislocation (or a dislocation dipole) on a boundary which is sufficiently far from the dislocation point for the differences of the displacement vector to be well approximated by their corresponding differentials. The result will match seamlessly a calculation on a much larger lattice provided the far field of a dislocation is the same as that given by linear elasticity as we depart from the dislocation point. Despite the slow decay of the elastic strain away from the dislocation point, differences and differentials of the elastic displacement become indistinguishable a few atoms away from the dislocation point.

Recently, we have developed periodized discrete elasticity models of dislocations in cubic crystals that describe their motion and interaction. These models appear to provide the simplest correction to the equations of elasticity allowing nucleation and motion of defects and have two main ingredients. Firstly, by discretizing elasticity, a linear lattice model involving nearest and next-nearest neighbors is found. In the continuum limit, this lattice model provides the equations of elasticity with the appropriate crystal symmetry. Secondly, we need to account for the fact that certain atoms at the core of a dislocation change their next neighbors as the dislocation moves. Thus to describe dislocation motion we need an algorithm to update neighbors. Alternatively, we can periodize discrete differences along the primitive directions of the crystal by using an appropriate periodic function, thereby restoring crystal periodicity. This periodic function is selected in such a way that elasticity equations are recovered far from defect cores and stable static defects can be generated using their known elastic far fields at zero applied stress. For applied stresses surpassing the Peierls stress of the material, the defects should move and the value of the Peierls stress can be used to calibrate the periodic function.

Here we extend this idea to the study of defects and their impact on the mechanical response of graphene sheets at zero temperature. Mostly planar graphene is very different
from cylindrical carbon nanotubes having very small radii and large curvature and from 3D graphite whose descriptions necessarily require studies different from the present one. Two facts complicate the task of designing lattice models of graphene. One is that graphene has a two-atom basis. The other is that a planar hexagonal lattice is intrinsically anisotropic even if its continuum limit is isotropic elasticity. We have dealt with these issues by using difference operators whose continuum limits are linearly independent combinations of the partial derivatives entering the Navier equations of linear elasticity. The key idea to choose our difference operators is that they do not have to involve all neighbors of an atom on an equal footing. Once we have an appropriate discretization of linear elasticity, we periodize the resulting lattice model in a way that allows dislocation gliding. Adding thermal effects and local curvature effects due to ripples increases the complexity of models and we have omitted these effects in the present paper, which is organized as follows. Section II recalls some basic details on the structure of graphene lattices and presents the stable defects we have obtained by solving the periodized discrete elasticity models. The basis of these models are lattice models obtained by discretizing elasticity, as indicated in Section III. Periodized discrete elasticity is discussed in Section IV. Section V describes numerical tests of defects in graphene carried out with the full 2D model and with a scalar reduced version thereof. Among solutions of our models, we have found a stable octagon defect. Known defects such as pentagon-heptagon pairs and 5-7-7-5 SW defects move and interact as expected. In particular, the two 5-7 defects comprising the two edge dislocations with opposite Burgers vectors of a SW glide to each other on their common gliding line and annihilate. This is not the case if we have a 7-5-5-7 defect (similar to the SW, but now the pentagons share a common side), which is stable because the dislocation centers of the component edge dislocations are displaced one atomic distance from each other in different glide lines. As observed in experiments, we have also obtained stable 5-8-5 divacancies and symmetric vacancies. We find that a symmetry-breaking vacancy evolves toward a simple threefold symmetric vacancy. Lastly, Section VI contains our conclusions.

II. STRUCTURAL CHARACTERISTICS OF GRAPHENE AND DEFECTS

Figure illustrates the structure of a graphene sheet comprising a single-layer of graphite. This 2D hexagonal lattice is equivalent to a cubic lattice with a two-atom basis generated
FIG. 1: (Color online) Graphene hexagonal lattice. The unit cell vectors are $\mathbf{a} = a(1, 0)$ and $\mathbf{b} = a(\cos(\frac{\pi}{3}), \sin(\frac{\pi}{3}))$, with a lattice constant $a = 2.461$ angstroms. The unit cell contains two carbon atoms $A = (0, 0)$ and $B = a(\cos(\frac{\pi}{6}), \sin(\frac{\pi}{6}))$ belonging to two sublattices. An atom $A$ has three nearest neighbors, six next-nearest neighbors and three second-nearest neighbors.

FIG. 2: (Color online) Numerically generated single pentagon-heptagon defect and its motion. The pentagon marks the lower end of an extra column of hexagons and its basis forms the top of the heptagon. (a) Under sufficiently large applied shear stress, a dark atom moves to the right and a light one to the left as indicated by the arrows. (b) The result is that the pentagon-heptagon defect moves one step to the right.

by two non orthogonal unit cell vectors, $\mathbf{a} = (1, 0)a$ and $\mathbf{b} = (1, \sqrt{3})a/2$, where $a$ is the lattice constant. The length of a hexagon side is $l = a/\sqrt{3}$. Dark and light colors are used to distinguish the two sublattices: dark atoms belong to sublattice 1 and light ones to sublattice 2.

Graphene layers often contain defects: an experimental study of defects generated by
irradiation and images thereof is reported in Ref. 9. Several defects have been obtained by using the periodized discrete vectorial model presented in the next Section. They appear in Figures 2, 3, 4 and 5. In these figures and successive ones in the paper, only the positions of the atoms have been calculated numerically. The bonds between neighboring atoms are only aids to visualize the structure of the lattice.

A simple defect in a graphene sheet is the single pentagon-heptagon (5-7) pair, depicted in Figures 2(a) and (b). As we will show later, this defect represents an edge dislocation and its gliding motion can be characterized by atom motion together with breakup and attachment of bonds between atoms. Note that link breakup and union occurs only between atoms in the direction of defect motion. As a result of this motion, the neighbors of the moving atoms in the rows immediately above and below them change. If we recall that bonds between neighboring atoms are only visualization aids, saying that “one defect moves by breaking bonds between some atoms and creating bonds with others” is only a figure of speech, not a consequence of the model.

Our models show that edge dislocations can have cores different from those depicted in Figure 2. In Fig. 3, eight atoms, one with a dangling bond, form the core of an edge dislocation. When the dark atom moves to the left as indicated by the arrow and is attached to the light atom with the dangling bond, the light atom formerly connected to it moves to the right and one of its bonds is left dangling. The overall result is the one-step motion of the octagon defect to the right.

Figure 4 depicts a vacancy defect which is a possible core of an edge dislocation dipole, as it will be shown later. Fig. 4(a) is the initial configuration obtained by discretization of
FIG. 4: (Color online) Numerically generated edge dislocation dipoles. Arrangement of atoms in a vacancy: (a) initial configuration as given by linear elasticity; (b) final configuration after time relaxation in the unstressed lattice.

FIG. 5: (Color online) Numerically generated edge dislocation dipoles. (a) 5-8-5 divacancy. (b) 5-7-7-5 Stone-Wales defect.

the elastic field of an edge dislocation dipole. Fig. 4(b) is the final configuration obtained by evolution of (a).

Figure 5(a) depicts another possible core of an edge dislocation dipole: a divacancy formed by an octagon with two adjacent pentagons. This 5-8-5 defect is dynamically stable. Yet another different core of the dipole is the Stone-Wales (SW) defect formed by a pair of heptagon-pentagons, as depicted in Fig. 5(b). This 5-7-7-5 defect is created by the SW bond rotation of one light atom forming the lowest vertex of an hexagon and bond reattachment to another light atom to form the top of a pentagon and, at the same time, the basis of an heptagon. This bond rotation leaves an oppositely oriented 5-7 defect next to the other one. This SW defect has been obtained by superposition of the initial guesses for the elastic displacement fields of two opposite edge dislocations with heptagon-pentagon cores that share the same glide line. In elasticity, we would expect that the two component edge dislocations of this dipole glide towards each other and annihilate, leaving the undisturbed lattice as a result. This is in fact what the numerical solution of our model shows. Thus the SW defect represents an edge dislocation dipole and it seems to be dynamically unstable\cite{16}. 

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FIG. 6: (Color online) The two component 5-7 edge dislocations of a SW defect separate under a large applied shear stress. This figure has been calculated using a simple periodized scalar model in a lattice with $18 \times 18$ lattice spacings.

If a sufficiently large shear stress is applied to a lattice that contains one SW defect, its two component dislocations drift apart as shown in Fig. [6].

If we form a different dipole with component dislocations having opposite Burgers vectors and different glide lines, the two component edge dislocations glide towards each other but do not meet. Instead they form a stable dislocation dipole. As in the case of nanotubes\textsuperscript{13}, the simplest realization of this idea is a 7-5-5-7 defect in which the two pentagons (not the two heptagons as in the SW configuration) share a common side. Our results show that the 7-5-5-7 defect is dynamically stable\textsuperscript{23}.

III. DISCRETE ELASTICITY MODELS FOR GRAPHENE SHEETS

We want to produce a lattice model that reduces to the equations of linear isotropic elasticity in the far field of a defect. In the continuum limit, elastic deformations of graphene sheets are described by the Navier equations for the 2D displacement vector $(u, v)$:

\begin{align}
\rho \frac{\partial^2 u}{\partial t^2} &= C_{11} \frac{\partial^2 u}{\partial x^2} + C_{66} \frac{\partial^2 u}{\partial y^2} + (C_{66} + C_{12}) \frac{\partial^2 v}{\partial x \partial y}, \\
\rho \frac{\partial^2 v}{\partial t^2} &= C_{66} \frac{\partial^2 v}{\partial x^2} + C_{11} \frac{\partial^2 v}{\partial y^2} + (C_{66} + C_{12}) \frac{\partial^2 u}{\partial x \partial y},
\end{align}

where $\rho$ is the mass density. In the basal plane, graphite is isotropic, so that $C_{66} = \mu$, $C_{12} = \lambda$ and $C_{11} = \lambda + 2\mu$, in which $\lambda$ and $\mu$ are the Lamé coefficients.
A. Discretization of the Navier equations and linear lattice model

We would like to find a linear lattice model by discretizing the Navier equations (2). This is not an obvious task because any symmetric combination of differences involving either all nearest neighbors, or all nearest and next nearest neighbors, etc. only yields a multiple of the Laplacian in the continuum limit, no matter how many neighbors we use. We have allowed combinations of differences that do not involve all neighbors of the same type symmetrically. The idea is to select three difference operators that yield three independent linear combinations of \( \partial^2 u / \partial x^2 \), \( \partial^2 u / \partial y^2 \) and \( \partial^2 u / \partial x \partial y \) in the continuum limit. Then we replace the partial derivatives of \( u \) in (2) by the combinations of our differences that provide those partial derivatives in the continuum limit.

Let us select assign the coordinates \((x, y)\) to the atom \(A\) in sublattice 1 (see Figure 1). The three nearest neighbors of \(A\) belong to sublattice 2 and their cartesian coordinates are \(n_1, n_2\) and \(n_3\) below. Its six next-nearest neighbors belong to sublattice 1 and their cartesian coordinates are \(n_i, i = 4, \ldots, 9:\)

\[
\begin{align*}
  n_1 &= \left(x - \frac{a}{2}, y - \frac{a}{2\sqrt{3}}\right), \\
  n_2 &= \left(x + \frac{a}{2}, y - \frac{a}{2\sqrt{3}}\right), \\
  n_3 &= \left(x, y + \frac{a}{\sqrt{3}}\right), \\
  n_4 &= \left(x - \frac{a}{2}, y - \frac{a\sqrt{3}}{2}\right), \\
  n_5 &= \left(x + \frac{a}{2}, y - \frac{a\sqrt{3}}{2}\right), \\
  n_6 &= (x - a, y), \\
  n_7 &= (x + a, y), \\
  n_8 &= \left(x - \frac{a}{2}, y + \frac{a\sqrt{3}}{2}\right), \\
  n_9 &= \left(x + \frac{a}{2}, y + \frac{a\sqrt{3}}{2}\right) .
\end{align*}
\]

(3)

Four of these atoms are separated from \(A\) by the primitive vectors \(\pm a\) and \(\pm b\) (see Figure 1).

Let us define four operators acting on functions of the coordinates \((x, y)\) of node \(A:\)

\[
\begin{align*}
  T u &= [u(n_1) - u(A)] + [u(n_2) - u(A)] + [u(n_3) - u(A)], \\
  H u &= [u(n_6) - u(A)] + [u(n_7) - u(A)], \\
  D_1 u &= [u(n_4) - u(A)] + [u(n_9) - u(A)], \\
  D_2 u &= [u(n_5) - u(A)] + [u(n_8) - u(A)].
\end{align*}
\]

(4) (5) (6) (7)

Note that the operator \(T\) involves finite differences with the three next neighbors of \(A\) which belong to sublattice 2, whereas \(H\) and \(D_1\) involve differences between atoms belonging to the same sublattice along the primitive directions \(a\) and \(b\), respectively. See Figure 1. \(D_2\)
FIG. 7: (Color online) Neighbors of a given atom $A$. Only the neighbors labeled 1, 2, 3, 4, 6, 7 and 9 are affected by the difference operators $T$, $H$ and $D_1$ used in our discrete elasticity model with two slip directions.

involves differences between atoms belonging to the same sublattice along a different choice of the basis vectors: $a$ and $c$ (parallel to the line joining atoms $n_5$ and $n_8$ in Fig. 7 which is also another primitive direction). The operator $D_2$ will be important when we want to consider a dislocation motion along slip directions $a$ and $c$. Taylor expansions of these finite difference combinations about $(x, y)$ yield

$$T u \sim \left( \frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2} \right) \frac{a^2}{4},$$

$$H u \sim \frac{\partial^2 u}{\partial x^2} a^2,$$

$$D_1 u \sim \left( \frac{1}{4} \frac{\partial^2 u}{\partial x^2} + \frac{\sqrt{3}}{2} \frac{\partial^2 u}{\partial x \partial y} + \frac{3}{4} \frac{\partial^2 u}{\partial y^2} \right) a^2,$$

$$D_2 u \sim \left( \frac{1}{4} \frac{\partial^2 u}{\partial x^2} - \frac{\sqrt{3}}{2} \frac{\partial^2 u}{\partial x \partial y} + \frac{3}{4} \frac{\partial^2 u}{\partial y^2} \right) a^2,$$

as $a \to 0$.

1. Model with three slip directions

Let us assume that we want to allow dislocations to slip along any of the three primitive directions $a$, $b$ or $c$. Then we replace in Equations (1) and (2) $H u/a^2$, $(4T - H)u/a^2$ and
\((D_1 - D_2)u / (\sqrt{3}a^2)\) instead of \(\partial^2 u / \partial x^2\), \(\partial^2 u / \partial y^2\) and \(\partial^2 u / \partial x \partial y\), respectively, with similar substitutions for the derivatives of \(v\). In terms of the Lamé coefficients, we obtain the following equations at each point of the lattice:

\[
\rho a^2 \frac{\partial^2 u}{\partial t^2} = 4\mu T u + (\lambda + \mu) H u + \frac{\mu + \lambda}{\sqrt{3}} (D_1 - D_2)v, \tag{8}
\]

\[
\rho a^2 \frac{\partial^2 v}{\partial t^2} = 4\mu T v + (\lambda + \mu) (4T - H) v + \frac{\mu + \lambda}{\sqrt{3}} (D_1 - D_2)u. \tag{9}
\]

**2. Model with two slip directions**

If we only allow slip along the two primitive directions \(a\) and \(b\) that form our vector basis, we should replace in Equations (1) and (2) \(H u / a^2\), \((4T - H) u / a^2\) and \(2(3D_1 - 3T + H/2)u / (\sqrt{3}a^2)\) instead of \(\partial^2 u / \partial x^2\), \(\partial^2 u / \partial y^2\) and \(\partial^2 u / \partial x \partial y\), respectively, with similar substitutions for the derivatives of \(v\). In terms of the Lamé coefficients, the following equations are then obtained

\[
\rho a^2 \frac{\partial^2 u}{\partial t^2} = 4\mu T u + (\lambda + \mu) H u + 2\left(\frac{\mu + \lambda}{\sqrt{3}}\right) (D_1 - 3T + \frac{1}{2}H) v, \tag{10}
\]

\[
\rho a^2 \frac{\partial^2 v}{\partial t^2} = 4\mu T v + (\lambda + \mu) (4T - H) v + 2\left(\frac{\mu + \lambda}{\sqrt{3}}\right) (D_1 - 3T + \frac{1}{2}H) u, \tag{11}
\]

at every point of the lattice. To have slip directions \(a\) and \(c\), we replace the operator \(-(D_2 - 3T + H/2)\) instead of \((D_1 - 3T + H/2)\) in (10) and (11). The same equations are found if \(B = (x, y)\) is an atom in sublattice 2. In this case, the relevant neighbors of \(B\) entering the definitions of \(T\), \(H\), \(D_1\) and \(D_2\) have coordinates

\[
\begin{align*}
n_1 &= \left( x - \frac{a}{2}, y + \frac{a}{2\sqrt{3}} \right), & n_2 &= \left( x + \frac{a}{2}, y + \frac{a}{2\sqrt{3}} \right), & n_3 &= \left( x, y - \frac{a}{\sqrt{3}} \right), \\
n_4 &= \left( x - \frac{a}{2}, y - \frac{a\sqrt{3}}{2} \right), & n_5 &= \left( x + \frac{a}{2}, y - \frac{a\sqrt{3}}{2} \right), & n_6 &= (x - a, y), \\
n_7 &= (x + a, y), & n_8 &= \left( x - \frac{a}{2}, y + \frac{a\sqrt{3}}{2} \right), & n_9 &= \left( x + \frac{a}{2}, y + \frac{a\sqrt{3}}{2} \right). \tag{12}
\end{align*}
\]

We may see the hexagonal lattice as a set of atoms connected by springs. These springs connect each atom \(A\) with its nearest neighbors \(n_1\), \(n_2\) and \(n_3\), and with its nearest neighbors along the primitive directions, \(n_4\), \(n_6\), \(n_7\) and \(n_9\). If we add symmetrically the missing two next-nearest neighbors \(n_5\) and \(n_8\) in the operator \(D_1\), its Taylor expansion produces the 2D Laplacian. Similarly, adding all the second-nearest neighbors, the Laplacian is found again:
Symmetric choices of neighbors only generate Laplacians, but not the terms involving cross derivatives. It seems reasonable to break the central symmetry about a given atom when defining finite differences by giving preference to the primitive directions. Notice that if we move along the lattice, the hexagonal arrangement itself is a source of anisotropy. Along the \( x \) direction (a primitive direction of the lattice), atoms are arranged in a ‘zig-zag’ pattern. The same arrangement occurs along the other primitive directions. However, along the \( y \) direction atoms are arranged in an ‘arm-chair’ pattern.

**B. Lattice model in primitive coordinates**

Equations (10) and (11) can be written in primitive coordinates \( u'_i \), \( i = 1, 2 \) (with \( u'_1 = u' \), \( u'_2 = v' \)), by means of the transformation \( u_i = T_{ij} u'_j \) (summation over repeated indexes is intended), with

\[
\begin{pmatrix}
  u \\
  v 
\end{pmatrix} = a \begin{pmatrix}
  1 & 1/2 \\
  0 & \sqrt{3}/2 
\end{pmatrix} \begin{pmatrix}
  u' \\
  v' 
\end{pmatrix}.
\]  

Writing Equations (8) and (9) or (10) and (11) as

\[
\rho a^2 \frac{\partial^2 u}{\partial t^2} = L_{ij} u_j,
\]

the linear equations of motion in the primitive coordinates are

\[
\rho a^2 \frac{\partial^2 u'_i}{\partial t^2} = L'_{ij} u'_j
\]

with

\[
L'_{ij} = T_{ik} L_{kn} T_{nj},
\]

or equivalently:

\[
\rho a^2 \frac{\partial^2 u'}{\partial t^2} = 4 \mu T u' + \frac{\lambda + \mu}{3} (3H - D_1 + D_2) u' + (\lambda + \mu) \left( H + \frac{D_1 - D_2}{3} - 2T \right) v',
\]  

\[
\rho a^2 \frac{\partial^2 v'}{\partial t^2} = \frac{\lambda + \mu}{3} (D_1 - D_2 - 3H) v' + 4(\lambda + 2\mu) T v' + \frac{2}{3}(\lambda + \mu)(D_1 - D_2) u',
\]

when there are three slip directions, or

\[
\rho a^2 \frac{\partial^2 u'}{\partial t^2} = \frac{\lambda + \mu}{3} [(H - D_1) u' + (2H + D_1) v'] + T[(\lambda + 3\mu) u' - 2(\lambda + \mu) v'],
\]  

\[
\rho a^2 \frac{\partial^2 v'}{\partial t^2} = \frac{\lambda + \mu}{3} [(H + 2D_1) u' + (D_1 - H) v'] + T[(\lambda + 3\mu) v' - 2(\lambda + \mu) u'],
\]

when there are only two slip directions along the basis vectors \( \mathbf{a} \) and \( \mathbf{b} \). Note that \( u' = (u - v/\sqrt{3})/a \) and \( v' = 2v/(a\sqrt{3}) \) are nondimensional. Equations (14) and (15) do not look symmetric in the same way as (16) and (17) do because we have selected a basis along primitive directions \( \mathbf{a} \) and \( \mathbf{b} \), so that \( \mathbf{c} \) (which defines the third slip direction associated to the operator \( D_2 \)) is not a basis vector.
C. Scalar model

When the displacements in the $y$ direction are negligible we may ignore the vertical component and work in cartesian coordinates. The evolution equations for the displacements in the horizontal direction are:

$$\rho a^2 \frac{\partial^2 u}{\partial t^2} = 4\mu Tu + (\lambda + \mu) Hu.$$ \hspace{1cm} (18)

This equation can also be obtained from $u = (u' + v'/2) a$ by adding twice (16) to (17), and then setting $v' = 0$. The only slip direction of the scalar model is along $a$.

IV. PERIODIZED DISCRETE ELASTICITY MODELS FOR GRAPHENE SHEETS

The models described by (8) - (9), (10) - (11) or (18) are linear and do not allow for the changes of neighbors involved in defect motion. An obvious way to achieve this is to update neighbors as a defect moves. Models such as (8) - (9), (10) - (11) or (18) would have the same appearance, but the neighbors $n_i$ would be given by (3) and (12) only at the start. At each time step, we keep track of the position of the different atoms and update the coordinates of the $n_i$. This is commonly done in Molecular Dynamics, as computations are actually carried out with only a certain number of neighbors. Convenient as updating is, its computational cost is high and analytical studies thereof are not easy. Another advantage of periodized discrete elasticity is that boundary conditions can be controlled efficiently to avoid spurious numerical reflections at boundaries.

In simple geometries, we can avoid updating by introducing a periodic function of differences in the primitive directions that automatically describes link breakup and union associated with defect motion. Besides greatly reducing computational cost, the resulting periodized discrete elasticity models allow analytical studies of defect depinning$^{19,20,22}$, motion and nucleation$^{21}$.\footnote{Here we refer to a previous study on the topic.}

To restore crystal periodicity, we replace the linear operators $T$, $H$, $D_1$ and $D_2$ in (14) -
by their periodic versions:

\[
T_p u' = g(u'(n_1) - u'(A)) + g(u'(n_2) - u'(A)) + g(u'(n_3) - u'(A)),
\]

(19)

\[
H_p u' = g(u'(n_6) - u'(A)) + g(u'(n_7) - u'(A)),
\]

(20)

\[
D_{1p} u' = g(u(n_4) - u'(A)) + g(u'(n_9) - u'(A)),
\]

(21)

\[
D_{2p} u' = g(u(n_5) - u'(A)) + g(u'(n_8) - u'(A)),
\]

(22)

where \( g \) is a periodic function, with period one, and such that \( g(x) \approx x \) as \( x \to 0 \). In our tests we have taken \( g \) to be a periodic piecewise linear continuous function:

\[
g_{\alpha}(x) = \begin{cases} 
  x, & -\alpha \leq x \leq \alpha, \\
  -\frac{2\alpha}{1-2\alpha}x + \frac{\alpha}{1-2\alpha}, & \alpha \leq x \leq 1 - \alpha.
\end{cases}
\]

(23)

The parameter \( \alpha \) controls defect stability and mobility under applied stress. \( \alpha \) should be sufficiently large for elementary defects (dislocations, vacancies) to be stable at zero applied stress, and sufficiently small for dislocations to move under reasonable applied stress\(^\text{19}\). The periodic function \( g \) can be replaced by a different type of periodic function to achieve a better fit to available experimental or numerical data.

Periodized discrete elasticity is a Lagrangian model: the atoms are labeled from the start and we track their motion. The periodic functions allow us to simulate dislocation motion without updating neighbors thereby greatly reducing computational cost.

In the simpler case of scalar elasticity\(^\text{18}\), the corresponding periodized discrete elasticity model is:

\[
\rho a^2 \frac{\partial^2 u}{\partial t^2} = 4 \mu T_p u + (\lambda + \mu) H u,
\]

(24)

\[
T_p u = a \left( \frac{u(n_1) - u(A)}{a} \right) + a \left( \frac{u(n_2) - u(A)}{a} \right) + a \left( \frac{u(n_3) - u(A)}{a} \right).
\]

(25)

The nonlinear function \( g \) is only needed for differences between the neighbors that may change due to defect motion. In the direction \( x \), neighbors never change, therefore we use the operator \( H \) of Eq. (5). Horizontal rows may shift, resulting in a shift of neighbors that is taken into account by using \( T_p \) as in (25) with the periodic function \( g_{\alpha} \) defined in (23).

V. DEFECTS IN GRAPHENE

In this Section, we discuss the defects obtained with our periodized discrete elasticity models. In our numerical calculations, we use for graphene the elastic constants of graphite
in the basal plain (which is isotropic), $C_{11} = C_{12} + 2C_{66} = 1060$ GPa, $C_{12} = \lambda = 180$ GPa, $C_{66} = \mu = 440$ GPa. This yields $\nu = 0.17$. We have used $\alpha = 0.2$ in the periodic function $g_\alpha(x)$. For this value, the Peierls stress for a 5-7 defect is 0.025 $\mu$, which is of the order of known values for covalent crystals. Larger (smaller) values of $\alpha$ yield larger (smaller) Peierls stresses: for $\alpha$ between 0.15 and 0.3, the Peierls stress varies in the range between $10^{-3}\mu$ and $10^{-1}\mu$. We have not calibrated our model to a precise value but this can be done when Peierls stresses for single graphene sheets are measured in experiments. All defects have been calculated using the vectorial model (16) - (17), periodized by means of (19) - (21). Analogous results can be found using appropriate computationally cheaper scalar models such as (24) - (25). Scalar models are more convenient to analyze defect motion, interaction and even nucleation of defects constrained to move along a given primitive direction. In more complex geometries a combination of periodized discrete elasticity models and neighbor updating could be useful.

In all cases, the construction of defects is similar. We use the displacement field of a given defect in continuum linear elasticity at zero applied stress both as initial and boundary condition for the discrete model. Then the model with overdamped dynamics (replacing second order time derivatives in the model equations by first order ones) is used to relax the initial field to the stable stationary solutions representing the sought defects. Applied stress can be implemented via the boundary conditions. To study defect motion and interaction, we can use the models with inertia.

How large should our computational lattice be? We can have an idea by using results by Zhang et al. As the far field of defects, they considered linear elasticity with a strain energy that was a quadratic functional of the strain tensor and also of its first and second derivatives. They compared results given by this theory with those of classical linear elasticity and with results from atomistic simulations. In their figure 11, it can be seen that four lattice spacings away from the core of a SW defect or of a divacancy in graphene, linear elasticity already approximates very well MD results (they use only 132 carbon atoms in their simulations). Thus a lattice with $18 \times 18$ lattice spacings (and therefore with $36 \times 36 = 1296$ atoms) should be sufficiently large to obtain good results for a centrally located defect by using our discrete elasticity models. The figures presented in this paper have been obtained using such a lattice and we have checked that our results do not change significantly by decreasing slightly the lattice size or by increasing it.
As we have said before, our model is more efficient than MD. In MD, one has two options when computing the force acting on each atom, either to include the interactions of one atom with all its neighbors, or to include only the neighbors within a certain distance of the atom and update them as time evolves. The cost of computing the force on each atom at each time step according to the first option is huge due to the large number of atoms involved, and therefore we need a very long time to compute just a few time steps. The second option (few neighbors and updating) is less computationally costly than the first one, but its cost is still large. In contrast to this, our model involves a few nearest neighbors and the periodic function used in the governing equations avoids the need of updating neighbors at each step. Thus computation of the forces at each step is cheap. We may compute the evolution of atoms in much larger lattices in a much shorter time. Since our forces are cheaper to compute, we may use higher order solvers to integrate the resulting system of differential equations. Since we are able to handle larger lattices, our trouble with numerical artifacts due to reflections on the walls of the lattice is smaller. In MD, reflections of waves at the boundaries of the computational domain limit the time spans over which simulations are reliable and there is a trade off between lattice size and the total time of a simulation. In fact, it is not yet clear how to prevent spurious reflections in MD simulations in an efficient manner\textsuperscript{28}. The simplicity of our model equations allows us to introduce simple non reflecting boundary conditions, thereby suppressing numerical size effects due to reflections at the boundaries.

A. Edge dislocations

Static edge dislocations can be generated using the overdamped version of (16)-(17) periodized by means of (19) - (21) and the elastic field of edge dislocations for (10)-(11). To find the stationary edge dislocation at zero stress, we first write the corresponding stationary edge dislocation of isotropic continuum elasticity. The displacement vector \( \mathbf{u} = (u(x, y), v(x, y), 0) \) of an edge dislocation directed along the \( z \) axis (dislocation line), and having Burgers vector \((b, 0, 0)\) is

\[
\begin{align*}
  u &= \frac{b}{2\pi} \left[ \tan^{-1} \left( \frac{y}{x} \right) + \frac{xy}{2(1-\nu)(x^2 + y^2)} \right], \\
  v &= \frac{b}{2\pi} \left[ -\frac{1 - 2\nu}{4(1-\nu)} \ln \left( \frac{x^2 + y^2}{b^2} \right) + \frac{y^2}{2(1-\nu)(x^2 + y^2)} \right], 
\end{align*}
\]

(26)
cf. Ref. 29, pag. 57. (26) has a singularity \( \propto r^{-1} \) at the core and it satisfies \( \int_C (d\mathbf{x} \cdot \nabla) \mathbf{u} = -(b, 0, 0) \), for any closed curve \( C \) encircling the \( z \) axis. It is a solution of the planar stationary Navier equations with a singular source term:

\[
\Delta \mathbf{u} + \frac{1}{1 - 2\nu} \nabla (\nabla \cdot \mathbf{u}) = -(0, b, 0) \delta(r).
\] (27)

Here \( r = \sqrt{x^2 + y^2} \) and \( \nu = \lambda/[2(\lambda + \mu)] \) is the Poisson ratio; cf. page 114 of Ref. 30.

The continuum displacement (26) yields the nondimensional static displacement vector in primitive coordinates \( U'(l, m) = [u(x, y) - v(x, y)/\sqrt{3}]/a \), \( V'(l, m) = 2v(x, y)/(a\sqrt{3}) \), where \( x = a(x' + y'/2), y = a\sqrt{3}y'/2 \). The primitive coordinates \( x' = x'_0 + l, y' = y'_0 + m \) are centered in an appropriate point \( (x'_0, y'_0) \) which is different from the origin to avoid the singularity in (26) to coincide with a lattice point.

\( U'(l, m) \) will be used to find the stationary edge dislocation of the discrete equations of motion. To this end, we replace the inertial terms \( \rho a^2 \partial^2 u'/\partial t^2 \) and \( \rho a^2 \partial^2 v'/\partial t^2 \) in (16) - (17) by \( \beta \partial u'/\partial t \) and \( \beta \partial v'/\partial t \), respectively. The resulting overdamped equations have the same stationary solutions. We use an initial condition \( u'(l, m; 0) = U'(l, m) \) given by Eqs. (26), and boundary conditions \( u'(l, m; t) = U'(l, m) + (Fm, 0) \) at the lattice boundaries \( (F \) is a dimensionless applied shear stress). If \( |F| < F_{cs} \) \( (F_{cs} \) is the static Peierls stress for edge dislocations), the solution relaxes to a static edge dislocation \( (u'(l, m), v'(l, m)) \) with the appropriate continuum far field.

Depending on the location of the singularity \( (x'_0, y'_0) \), there are two possible configurations corresponding to the same edge dislocation in the continuum limit. Figure 2 shows the structure adopted by the deformed lattice \( (l + u'(l, m), m + v'(l, m)) \) when the singularity is placed between two atoms that form any non-vertical side of a given hexagon. The core of the dislocation is a 5-7 defect. If the singularity is placed in any other location different from a lattice point, the core of the singularity forms an octagon having one atom with a dangling bond, as shown in Fig. 3. The dangling bond causes this configuration to be more reactive due to the possibility of attaching impurity atoms to the dangling bond. The octagon can also be seen as a 5-7 defect with an extra atom inserted between heptagon and pentagon. The basin of attraction of the octagon configuration seems to be larger than that of the 5-7 defect. That the same dislocation type may have two different cores is a familiar fact in crystals with diamond structure and covalent bonds, such as silicon; see page 376 in Ref. 17. There it is shown that the 60° edge dislocations may belong to the ‘glide set’ or to
the ‘shuffle set’. Seen from a certain direction, the cores of the glide set look like 5-7 defects whereas the cores of the shuffle set look like octagons with a dangling bond attached to one of their atoms. Ewels et al use the names “glide dislocations” and “shuffle dislocations” for the 5-7 defects and the octagons with a dangling bond (glide + adatom), respectively.

The glide motion of edge dislocations occurs in the direction of their Burgers vector and on the glide plane defined by the Burgers vector and the dislocation line. In the configurations of Figs. 2 and 3, a supercritical applied shear stress will move the dislocation in the $x$ direction on the glide plane $xz$. Our simulations show that the shuffle dislocations move more easily than the glide dislocations, as predicted by Ewels et al. For conservative or damped dynamics, the applied shear stress has to surpass the static Peierls stress to depin a static dislocation, and a moving dislocation propagates provided the applied stress is larger than the dynamic Peierls stress (smaller than the static one). A moving dislocation is a discrete traveling wave advancing along the $x$ axis, and having far field $(u'(l - ct, m) + Fm, v'(l - ct, m))$. The analysis of depinning and motion of planar edge dislocations follows that explained in Ref. with technical complications due to our more complex discrete model.

Similar results are obtained with the simpler scalar model. In this case, the continuum displacement vector of an edge dislocation at zero applied stress is

$$U(x, y) = \frac{b}{2\pi} \theta \left( x - x_0, \frac{(y - y_0)\sqrt{\mu}}{\sqrt{\lambda + 2\mu}} \right),$$

where $b(1, 0)$ is the Burgers vector and $\theta(x, y) = \arctan(y/x)$. Choosing the singularity point $(x_0, y_0)$ as explained above, we obtain configurations similar to those in Figures 2 and 3 except that there is no displacement in the vertical direction. To see the effect of a shear stress applied in the $x$ direction, we select as initial and boundary condition

$$U(x, y) = \frac{b}{2\pi} \theta \left( x - x_0, \frac{(y - y_0)\sqrt{\mu}}{\sqrt{\lambda + 2\mu}} \right) + Fy.$$

### B. Edge dislocation dipoles

An edge dislocation dipole is formed by two edge dislocations with Burgers vectors in opposite directions. Depending on how we place the cores of these dislocations, different dipole configurations result. Let $E(x, y)$ be the displacement vector corresponding to the edge dislocation we have considered before. The static configuration corresponding to a
dipole is either a vacancy (Fig. 4), a divacancy (Fig. 5a) or a Stone-Wales defect (Fig. 5b).

To obtain these different dipole cores, we use the following initial and boundary conditions at zero stress:

- **Vacancy**: \( \mathbf{E}(x - x_0, y - y_0 - l/2) - \mathbf{E}(x - x_0, y - y_0) \). \( l = a/\sqrt{3} \) is the hexagon side in terms of the lattice constant \( a \).

- **Divacancy**: \( \mathbf{E}(x - x_0, y - y_0 - l) - \mathbf{E}(x - x_0, y - y_0) \).

- **Stone-Wales**: \( \mathbf{E}(x - x_0 - a, y - y_0) - \mathbf{E}(x - x_0, y - y_0) \).

We have set \( x_0 = -0.25a, y_0 = -0.4l \) to draw Figures 4 and 5.

The vacancy represented in Fig. 4(b) is similar to that proposed in Ref. 24 using tight-binding calculations in graphite surfaces. The initial configuration represented in Fig. 4(a) (asymmetric vacancy) evolves toward the dynamically stable symmetric vacancy represented in Fig. 4(b) for overdamped dynamics. For conservative dynamics and zero initial velocity, the initial asymmetric vacancy evolves towards a stable oscillation about the symmetric vacancy. The symmetric vacancy has the threefold symmetry observed in experiments. In a recent paper, Telling et al. 31 propose that asymmetric vacancies are stable single vacancy defects in graphite sheets provided atoms are allowed to be displaced from the plane (see Figure 1 in Ref. 25). They further propose that, at room temperature, the displaced atom rotates around the vacancy center which would also explain the threefold symmetry observed in experiments (and possessed by the symmetric vacancy). For the initial conditions we have considered with either conservative or overdamped dynamics and at zero temperature, we have not found stable configurations resembling Fig. 4(a) or oscillations between similarly asymmetric configurations with different orientations. Thus the asymmetric vacancy seems to be unstable for both overdamped and conservative dynamics in our model 16. If this is indeed the case, this configuration is a saddle point with the conservative dynamics. Even if we allow bending modes of the graphene sheet, the instability of this saddle is likely to persist for small bending modulus. Future work will be devoted to study this problem and whether our stability results for different defects change if ripples in graphene are allowed.

Both vacancies and divacancies are dynamically stable in an unstressed lattice. A sufficiently large applied stress along their glide direction (in both cases, the critical stress is about 0.12\( \mu \) for \( \alpha = 0.2 \)) splits these dipoles, thereby originating two edge dislocations with opposite Burgers vectors that move in opposite directions.
Another possible core of an edge dislocation dipole is a 5-7-5 SW defect, as in Fig. 5(b). When introduced as initial condition of Eqs. (16) - (17) periodized with (19) - (21), the SW configuration is likely to be unstable under zero applied stress. In fact, this configuration corresponds to two identical edge dislocations that have opposite Burgers vectors and share the same glide line. The dislocations comprising the dipole attract each other and are annihilated, leaving an undistorted lattice as the final configuration. Thus SW defects seem to be dynamically unstable except if we add external forces that produce the necessary bond rotation and stabilize them. If a shear stress is applied in their glide direction, these defects either continue destroying themselves or, for large enough applied stress (0.15µ for α = 0.2), are split in their two component heptagon-pentagon defects that move in opposite directions as shown in Fig. 6. Note that stability of SW defects may be very different in small-radius nanotubes which, unlike graphene, do not have edges on their lateral surface and have a large curvature.

If we shift the 5-7 defects comprising a SW in such a way that their glide lines are not the same, then we can obtain a configuration which is stable at zero applied stress. The simplest such case is a 7-5-5-7 defect in which the two pentagons share one side. We have checked that this defect is dynamically stable and that it splits in its two component edge dislocations when a shear stress exceeding 0.09µ is applied to the lattice for α = 0.2. Note that in all cases (vacancies, divacancies, 5-7-7-5 and 7-5-5-7 defects) the critical shear stress needed to split the dipole is larger than the Peierls stress for an edge dislocation. The reason is that splitting a dipole requires overcoming the resistance of the lattice to motion (Peierls stress) and the attraction experienced by two edge dislocations of opposite Burgers vectors. The latter depends on the defect configuration which thus determines the critical dipole splitting shear stress.

Instead of a dislocation dipole, our initial configuration may be a dislocation loop, in which two edge dislocations with opposite Burgers vectors are displaced vertically by one hexagon side: \( E(x - x_0 - a, y - y_0) - E(x - x_0, y - y_0 - l) \) \((l = a/\sqrt{3} \) is the length of the hexagon side). In principle, the dislocation loop could evolve to an inverse SW defect (7-5-5-7). Instead, this initial configuration seems to evolve towards a single octagon. If we displace the edge dislocations vertically by \( l/2 \), \( E(x - x_0 - a, y - y_0) - E(x - x_0, y - y_0 - l/2) \), the resulting dislocation loop seems to evolve towards a single heptagon defect.
C. Energetics

We can have an idea about the energy associated to each defect by using the energy of the scalar model (18) (measured with respect to the stress-free undistorted lattice),

\[ E = \sum_n \{ 2\mu \left( [u(n_1) - u(n)]^2 + [u(n_2) - u(n)]^2 + [u(n_3) - u(n)]^2 \right) \]

\[ + \frac{1}{2}(\lambda + \mu) \left( [u(n_6) - u(n)]^2 + [u(n_7) - u(n)]^2 \right) \}, \]  

(28)

where we sum over all points in the hexagonal lattice, belonging to sublattices A or B in Fig. 7, with neighbors given by (3) or (12), respectively. It can be seen that Eq. (18) is equivalent to \( \rho a^2 \partial^2 u/\partial t^2 = -\partial E/\partial u \). For a lattice with 16 × 16 lattice spacings, the energies associated to the defects we have described are 0.658 eV for the apparently unstable 5-7-7-5 SW defect, 2.283 eV for the octagon with a dangling bond (shuffle dislocation), 4.917 eV for the 5-7 defect, and in the case of the dislocation dipoles: 8.825 eV for the vacancy, 12.074 eV for the 5-8-5 divacancy and 9.483 eV for the stable 7-5-5-7 defect. These values are similar to those found by Ewels et al.\textsuperscript{11} for the activation barrier to form a glide dislocation dipole (8.99 eV) and a shuffle dislocation (2.29 eV) in graphene; cf. their Figure 3. Except for the SW defect, all other dislocation and dislocation dipole cores are stable and are obtained by dynamical evolution using the governing equations of the model from the class of initial conditions we mentioned above.

VI. CONCLUSIONS

We have studied edge dislocations and dislocation dipoles in planar graphene at zero temperature by means of periodized discrete elasticity models which seamlessly match the elastic field of dislocations and dipoles as the distance from their core increases. The cores of edge dislocations may be the well-known pentagon-heptagon defects of Fig. 2 (glide dislocations) or octagons with a dangling bond (shuffle dislocations)\textsuperscript{11} as in Fig. 3 depending on how we choose the initial configuration. Similarly, different cores are possible for edge dislocation dipoles: vacancies, 5-8-5 divacancies, Stone-Wales defects and 7-5-5-7 defects. Of these possible cores, symmetric vacancies, divacancies and 7-5-5-7 defects are dynamically stable whereas asymmetric vacancies and 5-7-7-5 SW defects are likely to be unstable. Our results show that regularizing linear elasticity near dislocation cores by periodized discrete
elasticity is a good alternative to computationally intensive atomistic simulations provided defects are sparse.

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