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Local curvature and stability of two-dimensional systems

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We propose a fast method to determine the local curvature in two-dimensional (2D) systems with arbitrary shape. The curvature information, combined with elastic constants obtained for a planar system, provides an accurate estimate of the local stability in the framework of continuum elasticity theory. Relative stabilities of graphic structures including fullerenes, nanotubes, and schwarzites, as well as phosphorene nanotubes, calculated using this approach, agree closely with ab initio density functional calculations. The continuum elasticity approach can be applied to all 2D structures and is particularly attractive in complex systems with known structure, where the quality of parameterized force fields has not been established.

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I. INTRODUCTION

Layered structures including graphite, hexagonal boron nitride, black phosphorus, transition-metal dichalcogenides such as MoS₂, and oxides including V₂O₅ are very common in nature. The possibility to form stable two-dimensional (2D) structures by mechanical exfoliation of these structures appears very attractive for a variety of applications [1,2]. The most prominent example of such 2D systems, graphic carbon, is the structural basis not only of graphene [1] but also of fullerenes, nanotubes, tori, and schwarzites [3–7]. Even though the structural motif in all of these systems may be the same, their mechanical and electronic properties depend sensitively on the local morphology [8–10]. Not only does the natural abundance of structural allotropes and isomers reflect their net energetic stability, but also the relative chemical reactivity of specific sites in a given structure correlates well with the local curvature and local stability [8–10]. This relationship has been well established for the reactive sites in the C₂₅₀ fullerene [8], used to induce structural collapse leading to chemical unzipping of carbon nanotubes [11–13], and to destroy collapsed carbon nanotubes [10].

For very large structures, estimating the global or local stability using ab initio calculations has proven impracticable. There, the stability has often been estimated using empirical rules or parameterized force fields including the Tersoff potential and molecular mechanics [14–17], with sometimes unsatisfactory results. Application of continuum elasticity theory, which can describe stability changes due to deviation from planarity, has been successful, but limited to systems with a well-defined, constant curvature [18,19]. Since strain energy is local and independent of the global morphology, it is intriguing to explore whether the local deformation energy may be accurately determined from local morphology estimates using the atomic geometry. If so, then the local stability in even arbitrarily shaped structures could be estimated accurately.

Here we propose a fast method to determine the local curvature in 2D systems with a complex morphology using the local atomic geometry. Curvature information alone, combined with elastic constants obtained for a planar system, provides accurate stability estimates in the framework of continuum elasticity theory. We find that relative stabilities of graphic structures including fullerenes, nanotubes, and schwarzites, as well as phosphorene nanotubes, calculated using this approach, agree closely with ab initio density functional calculations. The continuum elasticity approach can be applied to all 2D structures and is particularly attractive in complex systems with known structure, where the quality of parameterized force fields has not been established.

II. LOCAL CURVATURE AND CURVATURE ENERGY

The local curvature at a particular location on a surface is given by the two principal radii of curvature \( R_1 \) and \( R_2 \), as shown in Fig. 1. On a spherical surface, \( R_1 = R_2 \). On a cylindrical surface, \( R_1 \) is the cylinder radius and \( R_2 \to \infty \). Finally, a saddle point on a surface is characterized by opposite signs of \( R_1 \) and \( R_2 \). Knowing the principal radii of curvature everywhere, we may use continuum elasticity theory to determine the curvature energy \( \Delta E_C \) with respect to a planar layer using [20]

\[
\Delta E_C = \frac{1}{2} D \int_{\text{surface}} dA \left( \frac{1}{R_1^2} + \frac{1}{R_2^2} + \frac{2\alpha}{R_1 R_2} \right).
\]  

(1)

Here, the integral extends across the entire closed surface, \( D \) is the flexural rigidity, and \( \alpha \) is the Poisson ratio. Simple expressions for \( \Delta E_C \) can be obtained for simple morphologies such as a sphere or a cylinder, where \( R_1 \) and \( R_2 \) are constant everywhere [18]. This is, however, not the case in general.

We find it convenient to introduce the local mean curvature

\[
k = \frac{1}{2} \left( \frac{1}{R_1} + \frac{1}{R_2} \right)
\]  

(2)

and the local Gaussian curvature

\[
G = \frac{1}{R_1 R_2}.
\]  

(3)

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Using these quantities, we can rewrite Eq. (1) as

$$\Delta E_C = D \int_{\text{surface}} dA\left[2k^2 - (1 - \alpha)G\right].$$

In the following, we will consider the equilibrium arrangement of atoms in a planar 2D structure as the reference structure and will determine the local curvature from changes in the local morphology. The discrete counterpart of Eq. (4) for the curvature energy $\Delta E_C$ is a sum over atomic sites $i$:

$$\Delta E_C \approx DA \sum_i \left[2k_i^2 - (1 - \alpha)G_i\right],$$

where $A$ is the area per atom.

To use Eq. (5) for curvature energy estimates, we need to know the local curvatures $k$ and $G$ at all atomic sites. Our approach to estimate these values at a given site $P$ is illustrated in Fig. 1(d). According to Eq. (2), the local mean curvature $k$ should be close to the average inverse radius of curvature at that point:

$$k \approx \frac{1}{R}.$$

Since the atomic site $P$ and its nearest three neighbors $F_1$, $F_2$, and $F_3$ define the surface of a sphere of radius $R$, we take $k = 1/R$.

The positions of four atoms do not allow us to distinguish whether $P$ is on a plane, on a sphere, on a cylinder, or in a saddle point. We may obtain this additional information using the concept of angular defect. On any surface, which can be triangulated as shown in Fig. 1(d), the angular defect at a representative vertex $V_i$ is defined by $\Delta(V_i) = 2\pi - \sum_i \varphi_i$ in radian units. The local Gaussian curvature at $V_i$ is then given by [21]

$$G(V_i) = \Delta(V_i)/A_i = \left(2\pi - \sum_i \varphi_i\right)/A_i,$$

where $A_i$ is the total area of the triangulated surface divided by the number of vertices.

For trivalent molecular graphs containing five-, six-, and seven-membered rings found in fullerenes, carbon nanotubes, and schwarzites, a unique triangulation may be obtained by connecting the centers of adjacent polygons. This method is referred to as the dual graph in graph theory [22] and its use is illustrated in Fig. 1(d). Since $P$ is not a vertex in the dual graph, but rather the center of the triangle $\Delta V_1V_2V_3$, we must infer the local Gaussian curvature at $P$ from the angular defects at $V_1$, $V_2$, and $V_3$. If vertex $V_j$ is surrounded by $n_j$ triangles, we may assign to point $P$ the angular defect $\Delta(P) = \Delta(V_1)/n_1 + \Delta(V_2)/n_2 + \Delta(V_3)/n_3$. Then, we can estimate the local Gaussian curvature at $P$ as

$$G(P) = \Delta(P)/A,$$

where $A$ is the average area per atom. We use $A = 2.62$ Å², the value found in the honeycomb lattice of graphene, for all graphitic structures.

The above definition of the local Gaussian curvature satisfies exactly the equality

$$A \sum_{\text{atoms}} G(P_j) = A_1 \sum_{\text{vertices}} G(V_j) = 2\pi \chi.$$

Here, $\chi$ is the Euler characteristic of the surface, given by $\chi = 2 - 2g$, where $g$ is the genus, meaning the number of holes. Of interest here is the fact that $\chi = 2$ for spherical objects like fullerenes and $\chi = 0$ for cylindrical objects such as nanotubes.

Equation (9) is the discretized version of the Gauss-Bonnet theorem [23] regarding the integral of the Gaussian curvature over an entire closed surface, called the sum of the defect, which is usually formulated as $\int_{\text{surface}} GdA = 2\pi \chi$.

The variation of the local Gaussian curvature $G$ and the local curvature energy $\Delta E_C/A$ across the surface of carbon polymorphs, including two fullerene isomers discussed in Sec. III B, a nanotube and a schwarzite structure, is displayed in Fig. 2. The local curvature energy in these $sp^2$ bonded structures has been evaluated using the elastic constants of graphene [18] $D = 1.41$ eV and $\alpha = 0.165$. The higher stability of the C$_{38}(17)$ isomer in Fig. 2(b) is reflected in a rather uniform local curvature energy and Gaussian curvature distribution. The low stability of the C$_{38}(2)$ isomer in Fig. 2(a) is reflected in a large variation of local curvature energy and Gaussian curvature, clearly indicating the most reactive sites. Cylindrical carbon nanotubes, such as the (10,10) nanotube displayed in Fig. 2(c), have zero Gaussian curvature and a constant local curvature energy caused by the mean curvature. Schwarzites such as the C$_{152}$ structure, displayed in Fig. 2(d), have only negative Gaussian curvature that may vary across the surface, causing variations in the local curvature energy.

III. VALIDATION OF THE CONTINUUM ELASTICITY APPROACH

We will next test the accuracy of the continuum elasticity approach by calculating the relative stability of nonplanar structures based on graphitic carbon. An infinite number of morphologies including nanotubes, fullerenes, and schwarzites may be produced by deforming a segment of a graphene layer and reconnecting its edges so that all carbon
To validate the continuum elasticity theory results, we calculated the total energy of a graphene monolayer and selected graphitic structures using \textit{ab initio} density functional theory (DFT) as implemented in the SIESTA code [24]. We used the local-density approximation (LDA) [25,26] and Perdew-Burke-Ernzerhof (PBE) [27] exchange-correlation functionals, norm-conserving Troullier-Martins pseudopotentials [28], and a double-$\zeta$ basis including polarization orbitals. The one-dimensional Brillouin zone of nanotubes was sampled by 16 $k$ points and the 2D Brillouin zone of graphene was sampled by $16 \times 16$ $k$ points [29]. The small Brillouin zones of schwarzites with several hundred C atoms per unit cell were sampled by only one $k$ point. We used a mesh cutoff energy of 180 Ry to determine the self-consistent charge density, which provided us with a precision in total energy of $\leq 2$ meV/atom. All geometries have been optimized using the conjugate gradient method [30], until none of the residual Hellmann-Feynman forces exceeded $10^{-2}$ eV/Å.

**A. DFT results for fullerenes**

Our DFT-LDA results for the relative energy $\Delta E_{\text{tot}}^{\text{DFT}}$ of optimized C$_n$ fullerenes [31,32] with respect to graphene are shown in Fig. 3(a). The various data points for one size correspond to different structural isomers, which are increasing fast in number with increasing $n$. If all fullerenes were perfect spheres, Eq. (4) would simplify to \[ \Delta E_C = 4\pi D(1 + \alpha). \] Using the proper elastic constants for graphene [18] \( D = 1.41 \) eV and $\alpha = 0.165$, we would estimate $\Delta E_C = 20.6$ eV for all fullerenes independent of size. The numerical values for the different optimized fullerene isomers in Fig. 3(a) are all larger, indicating that variations in the local curvature and bond lengths cause a significant energy penalty.

**B. Comparison between computational approaches for C$_{38}$ fullerene isomers**

As we show in the following, considering only local curvature variations across the surface (and ignoring precise atomic positions) allows continuum elasticity theory to quantitatively predict the strain energy with a precision competing with...
ab initio calculations. To illustrate this point, we present in Fig. 3(b) the total strain energy $\Delta E$ in 17 isomers of C$_{38}$ obtained using various approaches. The strain energy $\Delta E_{\text{DFT}}$ based on DFT, which is expected to represent closely the experimental results, is not only significantly lower than the predicted values $\Delta E_{\text{Keating}}$ based on the Tersoff potential [14] but also differs from this popular bond-order potential in the prediction of relative stabilities.

Next we demonstrate that accurate energy estimates may be obtained even for geometries optimized using simple potentials with only bond stretching and bond bending terms such as the Keating potential [33,34]:

$$\Delta E_K = \frac{1}{2} \alpha_K \sum_{i<j} \left( \frac{r_{ij}^2 - R_i^2}{R^2} \right)^2 + 2 \frac{1}{2} \beta_K \sum_{i<j} \left( \frac{\mathbf{r}_{ij} \cdot \mathbf{r}_{jk}}{R^2} + \frac{1}{2} R^2 \right)^2. \tag{10}$$

The first term sums over nearest-neighbor pairs and the second term sums over nearest-neighbor triplets, where $j$ and $k$ share the same neighbor $i$. DFT calculations for graphene yield $R = 1.42 \, \text{Å}$ as bond length, $120^\circ$ as bond angle, $\alpha_K = 11.28 \, \text{eV}/\text{Å}^2$, and $\beta_K = 4.14 \, \text{eV}/\text{Å}^3$.

Geometries of C$_{38}$ fullerene isomers optimized by DFT and the Keating potential are presented in [32]. Strain energies for Keating optimized fullerenes are shown in Fig. 3(b). Whereas the Keating optimized geometry is close to the DFT optimized geometry, the Keating strain energy $\Delta E_{\text{Keating}}$ clearly underestimates the DFT values and does not correctly represent the relative stabilities of the different isomers. As an alternative, we used the Keating optimized geometry to obtain the curvature strain energy $\Delta E_{\text{Keating}}$ using the continuum approach. We found that this approach represents the relative stabilities of isomers adequately and compares well to $\Delta E_{\text{DFT}}$. The curvature strain energy values are somewhat lower than the DFT values, since energy penalties associated with bond stretching and bending do not appear in the continuum approach. The small value of such corrections reflects the fact that in equilibrated structures bond lengths and angles are near their optimum. The largest errors are expected in frustrated structures, where not all bond lengths and angles can be optimized simultaneously.

One of the key findings of this study is that continuum elasticity theory provides not only a fast but also a relatively robust way to determine relative stabilities that are, to some degree, insensitive to the precise geometry. We illustrate this point in Fig. 3(c), where we compare different ways to determine the total strain energy $\Delta E$ in all C$_{38}$ isomers discussed in Fig. 3(b). $\Delta E_{\text{DFT}}$, shown by the solid line, is the difference between the total energy in DFT of optimized C$_{38}$ isomers and 38 carbon atoms in the graphene structure. $\Delta E_{\text{DFT}}$, given by the dashed line, is the curvature energy based on the DFT optimized geometry. $\Delta E_{\text{Keating}}$, given by the dash-dotted line, is the curvature energy based on the Keating optimized geometry. We note that all expressions provide an accurate representation of relative stabilities. As mentioned above, the fact that $\Delta E_C$ is about 10% lower than $\Delta E_{\text{DFT}}$ is caused by our neglecting the stretching and bending of discrete atomic bonds in the continuum approach.

### Encouraged by the level of agreement for C$_{38}$, we present in Fig. 4(a) the correlation between the curvature energy $\Delta E_{\text{Keating}}$ and $\Delta E_{\text{DFT}}$ based on DFT for all fullerenes discussed in Fig. 3(a). The narrow spread of the data points around the $\Delta E_{\text{Keating}} = \Delta E_{\text{DFT}}$ line confirms that the continuum elasticity approach is competitive in accuracy with computationally much more involved ab initio calculations.

To demonstrate the generality of our approach, we extend it from near-spherical fullerenes to nanotubes with cylindrical symmetry and schwarzites with local negative Gaussian curvature. Since nanotubes and schwarzites are infinitely large, we compare stabilities on a per-atom basis in these structures. Besides results for the fullerenes discussed in Figs. 3 and 4(a), Fig. 4(b) displays results for nanotubes with radii ranging between 2.5 and 9.0 Å and for schwarzite structures with 152, 192, and 200 carbon atoms per unit cell. These results again indicate an excellent agreement between curvature energies in Keating optimized structures and DFT-based strain energies.

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**FIG. 4.** (Color online) Strain energy $\Delta E$ in carbon nanostructures with respect to the graphene reference system. (a) Comparison between DFT-based total energies $\Delta E_{\text{DFT}}$ and the curvature energy $\Delta E_{\text{Keating}}$ based on Keating optimized geometries for all fullerene isomers considered in Fig. 3(a). (b) Comparison between DFT-based strain energies $\Delta E_{\text{DFT}}/n$ and curvature energies per atom $\Delta E_{\text{Keating}}/n$ for Keating optimized geometries of fullerenes, nanotubes, and schwarzites. Dashed lines represent agreement between DFT and continuum elasticity results.
This agreement is particularly impressive, since the spread of atomic binding energies extends over more than 1 eV.

D. Phosphorene nanotubes

As suggested at the outset, our approach to estimate relative stabilities is particularly valuable for unexplored systems such as monolayers of blue phosphorus [35], where model potentials have not yet been proposed. Our DFT-PBE results for a blue phosphorene monolayer indicate \( A = 4.78 \, \text{Å}^2 \) as the projected area per atom, \( D = 0.84 \, \text{eV} \), and \( \alpha = 0.10 \). The monolayer structure, shown in the top panel of Fig. 5(a), has an effective thickness of 1.27 Å. This structure can be rolled up to phosphorene nanotubes with different radii \( R \) using the approach used in the construction of carbon nanotubes [7]. As seen in Fig. 5(b), the strain energy for this geometry, obtained using continuum elasticity theory, agrees very well down to very small radii with results obtained using much more involved DFT calculations [36].

IV. DISCUSSION

Given a set of points in space, such as atomic positions, it is possible to construct a smooth surface that contains all these points in order to characterize its shape everywhere, and to eventually determine the deformation energy using the continuum elasticity approach.

We illustrate this point by tessellating the smooth surface of a graphitic nanocapsule, consisting of a cylinder capped by hemispheres at both ends and representing \( C_{120} \), in different ways. Our results in Fig. 6 show that the average curvature energy \( \langle \Delta \Delta E_C \rangle \) is rather insensitive to the tessellation density. The horizontal dashed line at \( \langle \Delta \Delta E_C \rangle = 0.099 \, \text{eV/Å}^2 \) represents an extrapolation to a dense tessellation, is \( \approx 5\% \) higher than the exact continuum elasticity value of 0.093 eV/Å², obtained for an ideal capsule with cylinder and hemisphere radius \( R = 3.55 \, \text{Å} \). The small difference arises from our approximate way to estimate the mean curvature \( k \) on the cylinder surface and at the interface between the cylinder and the hemisphere. The extrapolated value is also close to the

\[ \langle \Delta \Delta E_C^{\text{DFT}} \rangle = 0.100 \, \text{eV/Å}^2 \] based on the DFT optimized \( C_{120} \) capsule.

The reverse process to determine atomic positions from the shape alone is not unique. An informative example is the structure of a carbon nanotube. Whereas the precise atomic structure within each nanotube is defined by the chiral index, many nanotubes with different chiral indices share essentially the same diameter and the same local curvature. Thus, given only the diameter of a (wide) hollow cylinder representing a nanotube, it is impossible to uniquely identify the chiral index and thus the atomic position. As a matter of fact, identifying the precise atomic positions is not necessary, since according to continuum elasticity theory, supported by experimental evidence, the stability of nanotubes depends only on the tube diameter [7].

From its construction, the continuum elasticity description of local and global stability is best suited for very large structures with small local curvatures. Therefore, the high level of agreement between its predictions and \( ab \) \textit{initio} results in structures with large local curvatures is rather impressive. Among the different allotropes, we find the continuum elasticity description to be most accurate for carbon nanotubes, where all bond lengths are at their equilibrium value. In fullerences and blackazines, the presence of nonhexagonal rings, including pentagons and heptagons, prevents a global optimization of bond lengths and bond angles, reducing the agreement with DFT results.

Our stability results are consistent with the pentagon adjacency rule that provides an energy penalty of 0.7–0.9 eV for each pair of adjacent pentagons [37–39], which causes an increase of the local curvature. While this rule is surely useful, it cannot compare the stability of isomers with isolated pentagons or structures of different size.

What we consider the most significant benefit of our approach to determine local strain [32] is to identify the least stable sites in a structure. Local curvature and in-plane...
strain play the key role in both local stability and local electronic structure [9], which also controls the chemical reactivity [8,10]. Thus, our approach can identify the most reactive and the least stable sites, which control the stability of the entire system.

V. SUMMARY AND CONCLUSIONS

In conclusion, we have introduced a fast method to determine the local curvature in 2D systems with arbitrary shape. The curvature information, combined with elastic constants obtained for a planar system, provides an accurate estimate of the local stability in the framework of continuum elasticity theory. Relative stabilities of graphitic structures including fullerenes, nanotubes, and schwarzites, as well as phosphorene nanotubes calculated using this approach, agree closely with ab initio density functional calculations. The continuum elasticity approach can be applied to all 2D structures and is particularly attractive in complex systems with known structure, where the quality of parameterized force fields has not been established.

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