Interplay between nanometer-scale strain variations and externally
applied strain in graphene

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We present a molecular modeling study analyzing nanometer-scale strain variations in graphene as a function of externally applied tensile strain. We consider two different mechanisms that could underlie nanometer-scale strain variations: static perturbations from lattice imperfections of an underlying substrate and thermal fluctuations. For both cases we observe a decrease in the out-of-plane atomic displacements with increasing strain, which is accompanied by an increase in the in-plane displacements. Reflecting the non-linear elastic properties of graphene, both trends together yield a non-monotonic variation of the total displacements with increasing tensile strain. This variation allows to test the role of nanometer-scale strain variations in limiting the carrier mobility of high-quality graphene samples.

I. INTRODUCTION

Graphene is a promising material with many remarkable properties. However, its experimentally measured characteristics often do not match those theoretically predicted. Prominently, for example, the carrier mobility of graphene samples was found to strongly depend on the fabrication process and type of substrate. On rough SiO$_2$ the mobility of graphene samples is e.g. limited to a few ten thousand cm$^2$/Vs while using atomically flat hexagonal boron nitride (hBN) as a substrate allows mobilities up to several hundred thousands cm$^2$/Vs or more at low temperatures. Notwithstanding, world-record mobility graphene devices consist of a suspended graphene sheet and can even reach one million cm$^2$/Vs. As suspended graphene devices are not very practical, it is thus imperative to understand and tailor the carrier mobility limitations in supported graphene. A key element for this will be the control over the substrate induced interplay between structural and electronic properties. The truly two-dimensional nature of graphene makes this interplay, or generally the electromechanical coupling, very unique and promises tailored electronic properties by so-called strain engineering.

The effect of strain on the carriers in graphene is generally twofold. First, the area of the unit cell is altered resulting in a redistribution of the charge carrier density in a way that strain variations give rise to effective electron-hole puddles at low carrier densities. Second, strain generates a so-called pseudovector potential that is similar to the vector potential of a real magnetic field. It is exactly such a strain-induced randomly varying pseudovector potential, which has recently been identified as the limiting mechanism for the carrier mobility in high-quality graphene devices on a substrate. As Couto and co-workers showed, these strain variations are on a length scale of a few nanometers such that they act as long-range scattering centers allowing for pseudospin flips. Thus nanometer-scale strain variations enable direct backscattering that limits the carrier mobility in high-quality graphene samples. It has been shown that the mobility is inversely proportional to the strength of nanometer-scale strain variations. As these strain variations can be non-invasively monitored by the line-width of the graphene specific Raman 2D-line, fabrication processes have been recently optimized to suppress strain variations leading to significant improvements in carrier mobility. For future optimization a thoroughly understanding of the interplay between strain variations and external mechanical influences, such as ‘global’ strain are of great importance. Also in view of the promise to tailor electronic properties by strain engineering a detailed knowledge of this particular interplay is crucial.

In this work we discuss a molecular modeling study analyzing this interplay. Under compressive strain nanometer-scale strain variations may be intrinsically mingled with the formation of static ripples. We therefore focus our analysis on applied tensile strain. As for the physical origin of the nanometer-scale strain variations in graphene, thermal fluctuations as well as frozen ripples from the fabrication process and atomic defects in substrates like hBN are conceivable. We consider both the effect of thermal fluctuations and of a static Gaussian potential to model defects in a hBN substrate. For both potential sources of nanometer-scale strain variations we observe an intriguing non-monotonic variation of the average atomic displacements with increasing externally applied tensile strain. Arising from the non-linear elastic properties of graphene, this variation allows us to predict experimentally observable signatures of nanometer-scale strain variations in the measurement of the carrier mobility as well as in the measurement of
the line-widths of the Raman active G- and 2D-mode.

II. METHODS

Aiming for system sizes beyond the reach of first-principles approaches we base our molecular modeling study on the empirical long-range carbon bond order potential (LCBOP). Among a multitude of force fields for carbon material LCBOP exhibits a range of features that are critical to the targeted properties. Specifically, it yields sound velocities, a bending rigidity and elastic constants that match the experimental values within $\sim 10\%$. Of particular relevance for the present study is hereby the faithful representation of the highly nonlinear elastic properties of graphene. This is illustrated in Fig. 1 which shows the calculated stress in a graphene sheet as a function of externally applied strain as obtained with density-functional theory (DFT) using the CASTEP package (library pseudopotentials, 650 eV cutoff energy, and $7 \times 7$ k-point sampling) and treating electronic exchange and correlation at the level of the PBE functional.

Already at rather low strain values the stress-strain curve deviates from the linear behavior expected for a material with constant Young’s modulus of 1.02 TP [1]. Reproducing the findings of Kalosakas et al., the LCBOP data faithfully reproduces this behavior and achieves a match to the first-principles calculations within $\sim 10\%$.

All simulations were carried out with the LAMMPS package. The standard simulation cell contains a rectangular sheet of graphene with a side length $L$ of approximately 10 nm (3680 atoms), corresponding to optimized LCBOP carbon-carbon distances. Periodic boundary conditions are applied and a vacuum layer of 20 Å separates the graphene sheet from its periodic images. Static nanometer-scale strain variations induced e.g. by lattice imperfections in an underlying hBN substrate are modeled by subjecting the graphene sheet to the influence of an isotropic Gaussian potential. This potential is characterized by three parameters: its width, its strength and its perpendicular distance to the graphene sheet. Based on the discussion above, the width of the Gaussian potential must be much smaller than 100 nm. We performed simulations varying the width of the potential between one and four times the interatomic carbon-carbon distance in unstrained graphene (1.42 - 5.68 Å), which results in a strain variation of a few nanometer in size (see below). The conclusions put forward below are identically derived for all widths, which is why we restrict the detailed presentation of the results below to the case of three times the interatomic carbon-carbon distance. Preceding calculations suggest a maximum potential difference induced by point defects in hBN of the order of 100 meV. We find this confirmed by our own DFT calculations for a nitrogen surface vacancy and a boron interstitial. We correspondingly performed simulations varying the strength of the Gaussian potential in 10 meV steps between -100 meV (attractive) and 100 meV (repulsive). We obtain qualitatively the same results for all cases, and discuss below some selected values for the potential strength. The perpendicular distance of the Gaussian potential to the graphene sheet is fixed at 0.25 Å to induce out-of-plane displacements in the graphene sheet.
of similar magnitude as the in-plane displacements over the entire range of applied external strain. By doing so, we obtain both maximum in-plane forces $F_{x,y}$ and out-of-plane forces $F_z$ on the graphene sheet that are typically of the order of several meV/Å (1 meV/Å = 1.6 pN). This is illustrated in Fig. 2.

The effect of thermal fluctuations is modeled through a Nose-Hoover thermostat [7] with the temperature set to 300 K. To accommodate the effect of thermal expansion the lateral cell size was equilibrated in 4 ns molecular dynamics (MD) runs at zero applied pressure and using the velocity-Verlet integration scheme with 0.001 ps time steps. As the obtained fluctuations in the atomic positions in the graphene sheet are known to scale with the side length of the sheet [5], simulations were performed for various side lengths/number of carbon atoms in the cell to obtain the scaling relation between the atomic displacements and side length $L$.

For both the static Gaussian potential and the dynamic thermal fluctuation simulations the external strain is applied by rescaling the lateral simulation cell size. Specifically, we consider the effect of uniaxial strain along the zigzag direction ($x$ direction, cf. Fig. 2) and along the armchair direction ($y$ direction, cf. Fig. 2), as well as the effect of equibiaxial strain ($x$ and $y$ direction). In the uniaxial cases, the simulation cell size was kept fixed in the respective other direction. The applied strain is always varied from 0% to 0.5% in steps of 0.1% and from 0.5% to 15% in steps of 0.5%.

The central outcome of the simulations for a given external strain are the average atomic displacements and the relative changes in bond length. Distinguishing the average in-plane displacement $d_{ip}$, the average out-of-plane displacement $d_{oop}$, and the average total displacement $d_{tot}$, these are defined as

$$d_{ip} = \frac{1}{N} \sum_{i=1}^{N} \sqrt{(x_i - x_{i,0})^2 + (y_i - y_{i,0})^2},$$

$$d_{oop} = \frac{1}{N} \sum_{i=1}^{N} |z_i - z_{i,0}|,$$

$$d_{tot} = \frac{1}{N} \sum_{i=1}^{N} \sqrt{(x_i - x_{i,0})^2 + (y_i - y_{i,0})^2 + (z_i - z_{i,0})^2},$$

(1)

where $(x_i, y_i, z_i)$ is the position of atom $i$ at the applied Gaussian potential (MM) or at 300 K (MD), $(x_{i,0}, y_{i,0}, z_{i,0})$ is its equilibrium position at 0 K without applied Gaussian potential, and $N$ is the total number of atoms in the simulation cell.

For the case of the static Gaussian potential, these displacements are evaluated after relaxing the atomic positions under the applied strain until residual forces fell below 0.5 meV/Å. For the thermal fluctuations the displacements are obtained from MD simulations. Starting from the relaxed zero-strain geometry (see above), the system was first equilibrated under the applied strain over 4 ns in the (NVT) ensemble. The displacements are then obtained as time averages over 6 ns (NVE) trajectories.

### III. RESULTS

#### A. Effect of a static Gaussian potential

Figure 3 summarizes the obtained average displacements under applied equibiaxial strain and Gaussian potentials of strength $\pm 10$ meV, $\pm 50$ meV and $\pm 90$ meV. For all three potential strengths and irrespective of whether the potential is attractive or repulsive the average out-of-plane displacement $d_{oop}$ decreases monotonically with increasing strain. As the Gaussian potential and thus the force exerted on the graphene sheet is constant, this indicates an increase in the out-of-plane bending stiffness. The average in-plane displacement $d_{ip}$ exhibits exactly the opposite behavior, i.e. it increases with increasing strain. This reflects the decrease of the Young’s modulus with increasing strain that was discussed in Section II. Intriguingly, the increase of $d_{ip}$ is in all cases more than linear with strain. The reason
for this is that the decrease of Young’s modulus is more than linear with the strain (not shown). If one assumes a simple linear elastic model, in which the strong coupling between the in-plane and out-of-plane deformation is ignored, one finds the trivial result that the in-plane displacements should be independent of the strain. As we see a large increase in in-plane deformations, the difference between the in-plane displacement at finite strain and the one at zero strain quantifies how much the in-plane displacements are beyond linear elasticity. In our case, the in-plane displacements are up to a factor 5 larger than those expected from the simple linear elastic model. The opposite behavior of the average out-of-plane and in-plane displacement has the consequence, that the average total displacement $d_{\text{tot}}$ reaches a minimum for a certain amount of strain (see arrow). The precise value for this amount depends on the potential strength (as apparent from Fig. 3) and on the potential width (not shown). The stronger and narrower the potential, the higher the strain required to reach the minimum. In fact, for a potential width of only once or twice the size of the interatomic carbon-carbon bond, the minimum is always above 15% strain for the explored values of potential strength and for the considered, fixed perpendicular distance between the center of the potential and the graphene sheet. This is approximately equal to the breaking strength of graphene, and will, therefore, not be reachable in experiments.

Both the average in-plane and out-of-plane displacements depend on the width and the strength of the potential. We observe that the average out-of-plane displacements scale linearly with the strength of the potential, which is explained from the fact that the out-of-plane force scales linearly with the strength of the potential. However, the out-of-plane force is largely independent of the width of the potential, as the distance to the graphene sheet is fixed to 0.25 Å, which is much smaller than the explored widths in the simulation. Therefore, there is hardly any effect on the average out-of-plane displacements when the width of the potential is changed from one to four interatomic carbon-carbon bonds. The average in-plane displacements, on the other hand, are highly sensitive to both the width and the strength of the potential, since the in-plane force depends on both of these parameters.

Apart from the average displacements it is also instructive to analyze the spatial distribution of the individual atomic displacements over the graphene sheet. Figure 4(a) compiles corresponding data for two different equibiaxial strain values. The overall increase (decrease) of the in-plane (out-of-plane) displacements for the larger strain value (10%) is again clearly visible. The two types of displacements exhibit qualitatively different spatial distributions, as one would naturally expect from the qualitatively different spatial distribution of the in-plane and out-of-plane forces shown in Fig. 2(b). The out-of-plane displacements are largest exactly at the center of the sheet where the potential is applied and decay radially outwards. In contrast, the in-plane displacement distribution has a ring-like form, i.e. maximum displacements are found at some radial distance away from the source of the disturbance. This distance, or in other words the size of the ring, increases with the width of the potential, but is otherwise largely independent of the strength of the potential and the applied strain.

The fact that the in-plane and out-of-plane displacements differ in their spatial distributions, where out-of-plane displacements are largest at the center of the potential while in-plane displacements exhibit a ring-like form, directly translates to the carbon-carbon bond lengths within the sheet. We quantify this via the nearest neighbour vector $\mathbf{r}_{ij}$ and the relative change in bond length $r_{ij}^{\text{rel}}$ between neighbouring carbon atoms $i$ and $j$:

\[
\mathbf{r}_{ij} = (x_j - x_i, y_j - y_i, z_j - z_i),
\]

\[
r_{ij}^{\text{rel}} = \frac{|\mathbf{r}_{ij} - \mathbf{r}_{ij,0}|}{|\mathbf{r}_{ij,0}|}.
\]

![FIG. 4. (Color online) (a) Top view of the graphene sheet depicting the spatial distribution of the atomic displacements under the influence of a Gaussian potential and for equibiaxial tensile strain of 1% (left panels) and 10% (right panels). Compared are the in-plane displacements (upper panels) and the out-of-plane displacements (lower panels). The Gaussian potential of strength 90 meV is centered in the middle of the sheet and the discrete displacement data of the individual C atoms in the sheet has been interpolated to a continuous color map for easier visualization. (b) Relative change in carbon-carbon bond length as a function of the distance to the center where the Gaussian potential is applied. Shown are data for three tensile strain values of 1%, 5% and 10%. The cyan line is a guide to the eye.](image-url)
Here $|r_{ij}|$ is determined at 0 K without applied Gaussian potential. Figure 4(b) shows the relative change of this bond length $r_{ij}^{\text{rel}}$ as a function of the distance to the center where the potential is applied. At the smallest strain shown (1%) the variations are dominated by the out-of-plane displacements and largest changes in the bond lengths are correspondingly obtained at the position of the potential. For larger strains, the in-plane displacements gradually take over. Correspondingly, maximum bond length changes are also observed at larger distances from the applied potential. At a strain of 10% these maximum values are found about 8 Å away from the potential. The resulting diameter of the nanometer-scale strain variation of about 16 Å is therewith almost an order of magnitude larger than the width of the potential (4.26 Å) causing it. This is a consequence of the direct relation between the applied external potential and the resulting deformation potential that arises due to changes in bond lengths. 65,66

Even though the maximum bond changes are quite small (less than 0.2%), it is important to realize that this still results in a significant pseudomagnetic field $B$. The reason for this is that the previous work investigated a graphene structure with a much larger ratio between the out-of-plane displacement ($\sim 2.5$ nm) and the width ($\sim 5$ nm) of the nanometer-scale strain variation and furthermore did not allow the atoms to relax in-plane. As a consequence, the gradient of the relative bond length change $r_{ij}^{\text{rel}}$, which is the important quantity for the strength of the pseudomagnetic field, is in our case much smaller, which thus results in a much smaller pseudomagnetic field. Nevertheless, such small pseudomagnetic fields do affect the uniformity of the pseudomagnetic field in a graphene flake.

As Gaussian potentials or bubbles with different geometries induce inhomogeneous strain distributions in the graphene sheet with regions of compression, elongation and unaffected bond length, it is imperative to consider the change in displacements in the nanometer scale strain variation itself and thus take the atomic positions of the graphene sheet with the Gaussian potential at zero applied external strain as reference position. Figure 5 compiled the result for a Gaussian potential with a strength of 90 meV. The displacement in $x$- and $y$-direction of the atoms, that together form the in-plane displacement, are enhanced when going from 1% to 10% of applied tensile strain. This is fully consistent with the results in Figs. 3 and 4. In Figs. 4(a) and 4(b), the outer red and blue lobes indicate that the atoms are pushed radially outwards with respect to the position where the Gaussian potential is applied. In contrast, the inner lobes represent a region of compression and is a result of the strong coupling between the in-plane and out-of-plane displacement. The out-of-plane displacement $z$ keeps decreasing with applied tensile strain and reaches almost the out-of-plane displacement of the reference configuration at 10% applied strain.

Finally, we consider the change in local geometry of the two-dimensional surface, which is characterized by the discrete Gaussian curvature $K_D$ and the discrete mean curvature $H_D$. The former quantifies the local bending of the surface, whereas the latter measures the relative orientation of edges and normal vectors along a closed path. Both are zero in the case of a flat surface. Figure 6 shows both quantities for a Gaussian potential of 90 meV at an applied strain of 1% and 10%. Both, the discrete Gaussian curvature $K_D$ and discrete mean curvature $H_D$ show that the graphene becomes flatter with increasing
strain, as both quantities decrease at least one order of magnitude. This is consistent with the observed decrease in out-of-plane displacements.

In the simulations with uniaxial strain, both along the zigzag and the armchair direction, we qualitatively obtain equivalent findings as those just summarized for the equibiaxial strain case. This is visible in the average displacements compiled in Fig. 7(a). Also here, we find the decrease (increase) of the average out-of-plane (in-plane) displacements with strain. However, since the ratio of the out-of-plane to in-plane displacements is much larger for uniaxial strain, a minimum in the total average displacement is not found (within the range of strain values below the breaking point of graphene) for the investigated parameters of the Gaussian potential. The breaking of the graphene sheet can be observed as a sudden rise in the in-plane displacements at a strain around 13% along the armchair direction in Fig. 7(a).

Another important difference between equibiaxial and uniaxial strain is that the lower symmetry of the uniaxial strain leads to an anisotropy in the underlying spatial distributions, which are summarized in Fig. 7(b). For both straining directions it is seen that the spatial distributions of the in-plane and out-of-plane displacements are elongated along the direction of the applied strain. Considering the range of the observed displacements, it is obvious that the displacement of the atoms close to the center of the potential affects a number of neighbor atoms extending far beyond nearest neighbors and even outside of the area of the graphene sheet where the potential has any appreciable strength. To explain the observed spatial distributions of the displacements, we will assume that the range of the displacements, for a given magnitude of displacement of the central atoms, corresponds to a fixed number of carbon atoms along any direction outwards from the center of the potential. Thus, the elongated carbon-carbon bonds along the straining direction should result in a larger range of the displacements. For the in-plane displacements, where the central atoms experience a similar magnitude of displacement in the uniaxial and equibiaxial case, it is indeed seen that the range of the in-plane displacements along the straining direction in both the armchair and zigzag case closely matches the range observed along the corresponding straining direction in the equibiaxial case. For the out-of-plane displacements it is seen that the magnitude of the displacements are much larger in the uniaxial case, which indicates that the out-of-plane bending stiffness is increased less for uniaxial strain than for biaxial strain. As expected, the increased magnitude of the displacements leads to an increased range of the displacements both perpendicular to and along the straining direction. Thus, in this case
the range of the displacements perpendicular to the
while the range of the displacements along the straining
direction are much larger than the equibiaxial case. Note
that as the out-of-plane bending is always perpendicular
to the straining direction, the out-of-plane bending
stiffness is identical for all atoms independently of the
straining direction. The in-plane stretching mode, how-
ever, is expected to soften along the straining direction as
a consequence of the decrease of Young’s modulus with
increasing strain. Thus, a part of the observed increase in
the in-plane displacements along the straining direction
could also be explained by such strain-induced softening.

**B. Effect of thermal fluctuations**

We now proceed to the effect of thermal fluctuations
as determined by the MD simulations at 300 K. The left
and center panel in Fig. 8 show the time-averaged values
of the displacements obtained with Eq. 1 as a function of
applied strain for a side length \( L \) of 2.5 nm and 32 nm respec-
tively. In both panels, we observe that the out-of-plane
displacement \( d_{oo} \) (green dots) decreases with increasing
strain, whereas the in-plane displacement \( d_{ip} \) (red dots)
increases. As a consequence, the total displacement \( d_{ot} \)
(black dots) reaches a minimum for a certain strain value.
These results are thus in perfect agreement with the re-
sults of the MM simulations we discussed before. Overall,
the results of our MM and MD simulations suggest that
the out-of-plane displacements \( d_{oo} \) always decrease with
increasing strain and that in-plane variations always in-
crease with increasing strain. For this, it is not important
what the origin of the strain variation is.

2D crystals, such as graphene, were thought to be ther-
mostatically unstable due to their extremely small
thickness. In particular, the ratio between its effec-
tive thickness, which is defined as the average out-of-
plane fluctuations, and the side length of a rectangular
2D crystal \( L \), diverges if \( L \to \infty \). Therefore, it was a
big surprise that suspended graphene devices could be
made and were stable at room temperature. The reason
why graphene is stable is the strong coupling between
its bending and stretching modes. This coupling is
well described in membrane theory by a so-called crit-
tical exponent \( \xi \). We determine the dependence of
this critical exponent on the applied strain, by determin-
ing the average out-of-plane displacement \( d_{oo} \) as a func-
tion of the side length \( L \) for different values of strain,
cf. Fig. 8. The resulting values are compiled in Ta-
ble I. The exponent \( \xi \) of 0.64 at zero strain is in good
agreement with analytical work done on the statistical
mechanics of flexible membranes, self-consistent screen-
ing approximations, and MD simulations reported in
literature, which give an exponent of 0.58. For higher
strain values, the critical exponent decreases and already
reaches \( \xi = 0.25 \) at 0.5% of strain. This result indicates
that even a small amount of strain stabilizes graphene
(\( \xi < 0.5 \)), e.g. it remains flat in the sense that the aver-
age out-of-plane displacement remains much smaller than
the side length \( L \) even if \( L \to \infty \).

**TABLE I. Critical exponent \( \xi \) and its error as a function of
the applied strain.** The exponent \( \xi \) at 0% strain is 0.64, which
is in perfect agreement with literature.

| strain [%] | \( \xi \) | error [%] |
|-----------|--------|----------|
| 0         | 0.64   | 10       |
| 0.5       | 0.25   | 8        |
| 1         | 0.23   | 14       |
| 3         | 0.16   | 18       |
| 5         | 0.18   | 18       |
| 10        | 0.14   | 19       |

FIG. 8. Calculated time-average displacements as a function of strain from MD simulations at 300 K. Results are shown for
a side length \( L \) of 2.5 nm (left panel) and 32 nm (middle panel). In agreement with the MM simulations, \( d_{oo} \) decreases with
increasing strain, whereas \( d_{ip} \) increases. The right panel shows how the average out-of-plane displacement scales with the side
length \( L \) for different values of strain: 0% (black), 0.5% (red), 1% (green), 3% (blue), 5% (cyan), 10% (magenta). The slope of
the fitted lines yields the critical exponent of membrane theory, see text.
IV. SUMMARY AND CONCLUSIONS

We presented a molecular modeling study analyzing the effect of externally applied tensile strain on nanometer-scale strain variations of graphene. As source for such nanometer-scale strain variations we considered thermal fluctuations and a static Gaussian potential to model lattice imperfections in an underlying substrate, as the exact physical origins of these nanometer-scale strain variations are hard to capture. For both sources and for both uniaxial and equibiaxial strain the central outcome of our simulations is a decrease of the out-of-plane atomic displacements and a surprising increase of the in-plane atomic displacements with increasing strain. Details of both variations are beyond the expectations of linear elasticity theory, and reflect a strong increase of the bending rigidity and the known decrease of graphene’s Young’s modulus with strain, respectively. In conjunction both trends lead generally to a non-monotonic change in the total average displacement with strain. This displacement first drops with applied strain and then starts to rise again.

This finding suggests an intriguing opportunity to further investigate the influence of nanometer-scale strain variations as the major mobility limitation to graphene. If so, then the mobility of suspended graphene should initially increase with applied tensile strain, only to decrease again after a critical amount of strain. In contrast, we expect the confining potential of sandwich structures of the type hBN/graphene/hBN to largely suppress out-of-plane displacements. Leaving only in-plain strain variations, the mobility should in this case consistently be reduced upon application of tensile strain. The critical role and variation of the in-plane displacements could thereby be further validated through (tip-enhanced) Raman spectroscopy. Both the Raman active G-mode and 2D-mode are in-plane modes and are therefore expected to be sensitive to in-plane strain variations. Assuming the width of these modes to reflect the amount or strength of nanometer-scale strain variations, our simulations suggest an increase of this width with increasing tensile strain. The known splitting of the G-mode in the case of uniaxial strain could thereby represent a complication. However, such a splitting has not been observed for the 2D-mode and does also not occur for the G-mode in the case of equibiaxial strain. As such, externally applied strain appears as a highly suitable experimental knob to further understand the role of nanometer-scale strain variations in determining the material’s properties of high-quality graphene.

The presented insights may be also highly relevant for other two-dimensional crystals with non-linear Young’s modulus. As these materials have an unique electromechanical coupling analogous to graphene, a detailed understanding of this interplay is crucial for future applications of these two-dimensional crystals as well.

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