Laser-Based Texturing of Graphene to Locally Tune Electrical Potential and Surface Chemistry

Manoj Tripathi,*† Alice King,‡ Giuseppe Fratta,§ Manuela Meloni,† Matthew Large,‡ Jonathan P. Salvage,§ Nicola Maria Pugno,¶∥⊥ and Alan B. Dalton*†

†Department of Physics and Astronomy, University of Sussex, Brighton BN1 9RH, U.K.
‡School of Pharmacy and Biomolecular Science, University of Brighton, Brighton BN2 4GJ, U.K.
§Laboratory of Bio-Inspired & Graphene Nanomechanics, Department of Civil, Environmental and Mechanical Engineering, University of Trento, Via Mesiano 77, I-38123 Trento, Italy
¶School of Engineering and Materials Science, Queen Mary University of London, Mile End Road, E1 4NS London, U.K.
⊥Ket Lab, Edoardo Amaldi Foundation, Italian Space Agency, Via del Politecnico snc, I-00133 Roma, Italy

Supporting Information

ABSTRACT: A simple procedure of producing three-dimensional blisters of graphene through irradiation of the visible range laser by Raman spectrometer has been presented. Fabrication of different volumes of the blisters and their characterization were carried out with Raman spectroscopy by tuning the irradiation dose. The produced blisters showed a consistency in altitude and a remarkable change in functionality, adhesion force map and local contact potential difference as compared to untreated monolayer graphene and naturally occurred graphene nanobubbles. Nevertheless, bilayer graphene is unaffected in the applied laser doses. The laser irradiation led to lattice expansion of carbon atoms and introduced oxygenc functional groups with the structural disorder. The internal pressure of the gaseous molecules was evaluated by monitoring the shape of the graphene blisters and nanobubbles. High-resolution Raman mapping showed the impact of laser-affected area and the defect density ($n_d$) is reported as a function of displacement. Our results reveal ease of applicability of the Raman laser for the imaging and texturing of graphene pointing toward the possibility of the desirable and cost-effective laser writing at the submicron scale by tuning photochemistry of graphene which is pivotal for numerous applications.

INTRODUCTION

The extraordinary properties of graphene have drawn a great deal of interest in the field of mechanics,1 electronics,2,3 photonics,4 and virtually all disciplines of science and technology.5−7 The rapid progress in this emerging field resulted in tuning the extraordinary properties of graphene by altering the shape and dimensions,7,8 substrate interactions,5 and controlled functionalization.5,10 Inducing controlled and optimized strain in the graphene is another arising technology.5 On the other hand, biaxial strain in graphene can enhance the electronic−phonon coupling and potentially turn graphene into a superconductor.18

Patterning and texturing of graphene through a laser (light amplification by stimulated emission of radiation) is one of the efficient ways for straining and altering electrical and mechanical properties.19−21 Controlled and refined laser treatment can induce subtle chemical and structural changes in graphene, which leads to tailoring bandgap, the lattice expansion, functionalization, and conductive channel formation. It can be produced by a variety of laser treatments such as continuous wave,22 nanosecond23 and femtosecond lasers.24−26 Recently, complex and stable 3D structures of chemical vapor deposition graphene are produced by tuning the dose of irradiation to achieve desirable electrical and optical properties.26,27 Nevertheless, it will be more efficient to produce 3D structure by a simple laboratory instrument using visible radiation.21

Raman spectroscopy is considered as a sensitive and non-destructive laboratory tool for characterizing the chemical, mechanical, and electronic properties of graphene through vibrating carbon atoms.28−30 Raman spectroscopy is based on the vibrational transition of molecules occurring in the ground
electronic state. The irradiation of the Raman beam in the visible range of light changes the lattice constant of the sample leading to variations in phonon frequencies. Nevertheless, the choice of laser is one of the primary concerns that is integrated into the Raman spectrometer system. Prolong exposure to laser excitation energies of 2.33 eV (532 nm) and 1.87 eV (660 nm) at the focusing area up to few microns can result in damage or modification in the sample. Low-molecular-weight polymers are susceptible to initiate photochemistry/photophysics-induced morphology long before burning. The same applies to the adsorbed species in air conditions over catalysis. Nevertheless, tuned laser energy has been used to improve electrical properties in conjugated polymers by establishing planar conformations and higher conjugation length. In this study, Raman lasers (532 and 660 nm) at different power densities (mJ/μm²) have been irradiated over mechanically exfoliated graphene on silica. Atomic force microscopy (AFM) measures the maximum surface area affected under laser irradiation. A lower dose of power density (2.5–3.7 mJ/μm²) has been used for analysis, whereas higher doses (511–753 mJ/μm²) were used for the modifications. Different regions of single and bilayer graphene have been targeted in air condition. The 3D graphene structures produced showed distinguished morphology, adhesion force, and local surface potentials from untreated graphene crystal. The generation of 3D blisters is achieved through lattice expansion of carbon atoms and presence of trapped molecules. In addition, laser-induced partial oxidation of the graphene blisters has been verified from Raman spectroscopy. The areas affected by the laser irradiation were analyzed with Raman mapping as a function of displacement, and the defect density (n_d) was calculated. By tuning the power density of the laser treatment, desirable patterns of the altered electrical and mechanical properties are attainable. Thus, here, Raman spectrometer has been established as a detection tool which is utile for the modification of graphene.

### RESULTS AND DISCUSSION

**Morphology and Local Surface Potential of Blisters and Graphene Nanobubble.** Raman lasers have been used for characterization (thickness, oxidation, strain) and fabrication of graphene 3D structure by changing the power density.
Lower power density (2.55 mJ/μm²) was used for the thickness measurement at the initial stage (see Figure S1 for details), whereas higher power densities (511 and 753 mJ/μm²) were implemented for the fabrication at different volumes. Mechanically exfoliated graphene was used as the test sample, which comprises single layer (1LG) and bilayer (2LG) over 300 nm oxide layer of silica on the silicon substrate, Figure 1a. The conferred region is crucial to investigate the significance of subsurfaces, i.e., a single layer of graphene beneath 2LG and silica substrate under 1LG. Under irradiation of power density (511 mJ/μm²), arrays of (12 × 10) of 3D blisters of graphene are produced on 1LG only, nevertheless, 2LG was unaffected (Figure 1b). It depicts the vital role of interlayer coupling between the graphene layers in 2LG for distributing laser-induced heat energy by interlayer sliding. 1LG on silica substrate is resilient to slide due to higher shear strength and lower thermal conductivity of silica (1 W/(m K))-trapped heat energy between graphene and silica interface cause further heating of 1LG with time (10–20 s), which then leads to a short-range distortion and buckling of the graphene layer.

Controlled exposure of irradiation produces consistent altitude of graphene blisters of average height 2.5 ± 0.5 nm for power density 511 mJ/μm² and up to 6 ± 1.5 nm for 753 mJ/μm², Figures 1c–f and S2. Higher altitude (up to 20 nm) is reported in a controlled environment of argon and nitrogen gas by femtosecond laser. Nevertheless, continuous exposure of Raman laser at the high-power density in air condition at 33% of relative humidity can bring chemical changes in the graphene that will be discussed in the subsequent sections.

The surface chemistry (i.e., functionalization) of the produced blisters is different from the naturally ensued graphene nanobubbles produced during preparation of the sample. The adhesion force map in Figure 2a distinguishes chemical structure of the laser-induced graphene blisters from the 1LG basal plain surface, wrinkles, ripples, and the graphene nanobubbles. The graphene nanobubbles are used as a reference 3D structure, which are produced by strain in the graphene, the presence of trapped molecules, and the
deposition technique. These structural disorders are difficult to control during physical deposition of graphene but are useful to compare chemical and electrical characteristics from the laser-induced blisters. The adhesion force map illustrates the interaction between the silicon tip apex, elastic penetration, and the sample when the tip is moving vertically away from the surface, Figure S3. The adhesion force is linked with the contact area at the tip−graphene interface and the functional group attached on the graphene. Graphene nanobubble and the blisters resemble in adhesion force features, i.e., the adhesion force is higher at the central region of the structures than its surrounding curvature. It is due to the higher contact area between the tip apex and the central region of the blisters and at the graphene bubble. These central regions temporarily acted as deformed elastic membrane under normal force applied by tip apex. One can study the variation in adhesion force as a function of the graphene nanobubble volume. The present work is limited to comparison of properties between laser-induced and self-grown graphene structures.

Higher resolution (inset Figure 2a) of the adhesion force map of the blister illustrating a ring structure. It is in a contrast to the polymeric graphene nanobubble suggesting localized chemical perturbation in the blister. Local surface potential produced by KPFM also illustrates a ring structure of the blistered graphene of distinct contact potential difference (CPD), Figure 2b. Nearly 25−30% decrease is observed in the CPD values of the blistered central region compared to the untreated graphene. This resembles the wrinkled structure of graphene, which shows lower work function due to an increase of gap between graphene and silica substrates, which prohibits p-doping. The distinct CPD values between the inner and the outer ring of the blistered region also suggest oxidation of the graphene. Unlike blistered graphene, nanobubble CPD values are evenly distributed. The CPD values decrease with increasing altitude of the graphene nanobubble, Figure 2c, consequently the local work function (eV). The trends of work function (eV) for different regions are given at Figure 2d, which validates the influence of gap distance of graphene from the silica substrate. The work function of the 1LG measured as 5.02 eV as the highest and 4.79 eV for the 2LG at 33% of relative humidity are in coherence with the published findings. The lowering of work function for the 2LG can be explained by the screening of the charges (holes) originated from the SiO2 substrate by underlying graphene. Notably, the CPD values from the inner region of the ring were used for the work function measurement. This is in compliance with the CPD values measured for the nanobubble altitude up to 12
nm. These results decipher the locally tuned surface energy and electrical property for 1LG.

**Impact of Different Laser Power Density.** The fabrication of blistered graphene can be achieved by controlling three major factors: (1) carbon-lattice expansion at 1LG,26,41 (2) presence of trapped molecules between 1LG and hydrophilic silica interface,35 and (3) oxidation of carbon atoms irradiated in air conditions.27 The expansion of carbon lattice was monitored by investigation of G and 2D peaks at different power densities by two different lasers (λ = 532 and 660 nm), Figure 3a,b. There is a broadening of G and 2D peaks with increasing power density for both lasers indicates increasing temperature.42 This shows the introduction of peaks with increasing power density for both lasers indicates (phonon mode (E_{2g} symmetry) at the Brillouin Zone center, μ which can alter the Raman spectra. Therefore, dispersion of G chemical and structural changes in graphene due to heating, −Liu et al.10 during thermal perturbation for the entire sample. The frequencies of G and 2D peaks strongly depend on the change in charge density either by electrical, chemical, or thermal treatments owing to the static effect on bond lengths and nonadiabatic electron–phonon coupling.31,43 The G band (−1589) is associated with doubly generated (iTO and LO) phonon mode (E_{2g} symmetry) at the Brillouin Zone center, which occurs due to first-order Raman scattering process in graphene.28 The G peak frequency is extremely sensitive to eventual changes in the oscillation strength of electron–phonon interaction near the Fermi level.44 Generally, the phonon frequency of G band does not change by the energy of incident photon involved in Raman process.45 Nevertheless, prolonged exposure (up to 20 s) of the lasers might cause chemical and structural changes in graphene due to heating, which can alter the Raman spectra. Therefore, dispersion of G peak position with a variable irradiated power density (mJ/μm²) suggests a disorder in the carbon lattice.

The dispersion rate (∆G/ν; change in excitation laser) increases with disorder26 with the possible generation of local sp² and sp³ domains in graphene.37 The excitation energy (∆E_l) leads to a raised bandgap (E_g) energy in graphene, which enhances the Raman scattering signal due to the resonance condition in the optical absorption.28,47 Several reports demonstrated the impact of laser for decreasing sp² C=C bond and increment of C–C bond.26,27 We observed a decrease of I_G/I_{2D} ratio with higher power density, Figure 3c suggests an alteration in the hybrid state of sp² carbon atoms up to 50% at the highest power density as compared to the unirradiated graphene.

Two-dimensional (2D) peak position shows upshift for the laser source 532 nm as compared to 660 nm, Figure 3d. Unlike G frequency, the 2D band originates from a second-order process, involving two iTO phonons at the Dirac point (K). The incident photon energy changes the phonon energy in the Raman process. As the photon energy increases, phonons farther from K point are required for momentum conservation, which is responsible for significant dispersion.35,48 There is a softening of phonon modes due to lattice expansion for both lasers (λ = 532 and 660 nm) with a gradual increase of irradiation dose. Inset-marked frequencies for 2D- and G peak are measured at lowest power density from each laser. The slopes (S) measured by the linear fit for each curve are S = 1.9 for 532 nm and S = 1.1 for 660 nm showing higher biaxial strain39 for the former. Fairly large values of S have been reported experimentally as 2.45,46 2.6315 and for theoretical prediction as 2.2530 for biaxially strained graphene over hollow silica substrate in the pressurized conditions. The expansion of the carbon lattice through Raman laser irradiation is significantly low as compared with pressurized blistered graphene. This is further confirmed by measuring the topographic profile of the blistered graphene at a different dose of the power density, Figure 3e. The maximum height achieved at highest power density (753 mJ/μm²) is up to 6 ± 1.5 nm, which is less than pressurized blistered graphene.30,51

**Role of Trapped Molecules between Graphene and Silica Interface.** Graphene has shown impermeability to most of the gases,52 thus a differential pressure of several MPa can produce 3D structure.51 The role of trapped molecules to generate pressure and inflate graphene blister under laser irradiation was estimated by a geometrical approximation (Figure S4). The topographical informations (height (h) and width (w = 2r), where r is its radius) were used to evaluate internal pressure P of the graphene nanobubbles and the blisters by assuming hemispherical or cuboidal geometry. In particular, the force equilibrium imposes P2πr² = 2πrε (hemispherical) or Pw² = 4πrε (cuboidal), where r is the graphene thickness (t = 0.35 nm) and ε is its normal stress. Noting that ε = Eε, with ε strain of the graphene and E Young’s modulus of graphene (E around 1 TPa) and that geometrically ε = ∆L/L = ∆h/h, we derive

\[ P = 4Eε/\pi \]

The geometry, stress, strain, and pressure inside the blisters or nanobubbles are reported in Table 1; the pressure is significantly lower in graphene blisters with respect to nanobubbles.

| width, w (nm) | height, h (nm) | strain, ε | pressure, P (MPa) |
|-------------|-------------|---------|------------------|
| **Graphene blister** | | | |
| 156 | 1.87 | 0.024 | 215.15 |
| 498 | 2.75 | 0.011 | 31.08 |
| 484 | 3.40 | 0.014 | 40.64 |
| 506 | 6.54 | 0.026 | 71.52 |
| **Graphene nanobubbles** | | | |
| 175 | 15 | 0.171 | 1371.43 |
| 202 | 16 | 0.158 | 1097.93 |
| 431 | 23 | 0.107 | 346.68 |
| 542 | 46 | 0.170 | 438.45 |
| 620 | 50 | 0.161 | 364.20 |
| 646 | 40 | 0.124 | 268.38 |
| 700 | 38 | 0.109 | 217.14 |

The role of the trapped molecules between 1LG and silica interface is also evaluated by normally pressing graphene blistered at different normal forces from 0.250 to 10 nN. A systematic decrease in altitude of the blister was observed, see Figure S5. We successfully reduced the altitude of the blistered region up to 0.9 ± 0.15 nm at 10 nN of the normal force but were unable to regain its original configuration, which was 0.5 nm (measured by root mean square). The discrepancies in the values indicate the irreversibility of blistered graphene to regain its preceding conformation by normal pressing and also indicates the traces of the trapped molecules that did not escape completely. Separately, we did not observe any evident elevation of graphene blistered at the edge region due to
leakage of the trapped molecule\textsuperscript{35} although contrasting feature in the deformation map and adhesion force map has been observed indicating chemical perturbations (see Figure S6).

Theoretical studies revealed that absence of the trapped molecule could also lead to the formation of blister up to 1 nm in altitude at a laser wavelength of 93.1 nm, which might be based only on lattice extension.\textsuperscript{41} Extreme irradiation of the laser can lead to ablation of graphene carbon atoms and substrate underneath.\textsuperscript{53} Xing et al. observed graphene threshold damage limit >300 GW/cm\textsuperscript{2} of power density,\textsuperscript{54} nevertheless disruption in sp\textsuperscript{2} carbon bond can be initiated at 14–66 mJ/cm\textsuperscript{2}.\textsuperscript{55} We did not observe any ablation of graphene and silica substrate in the given range of power density. Nevertheless, a wide area (diameter up to 1.8 μm) of distorted graphene is observed by force–distance (\(F–D\)) spectroscopy through adhesion force map Figure 4a. The distorted area of graphene appeared in the annular fashion of radii \(R_h\) and \(R_i\), for \(R_h < R_c\). The affected area associated with \(R_h\) is highly distorted regime as compared to \(R_i–R_h\) region, where \(R_i = 3.3(R_h)\). The produced deformation shapes resemble to the proposed model for the activated D-band scattering in the Raman spectrum of graphene sheet by the point-like defect.\textsuperscript{56} The model defines the generation of the circular area over a graphene of different radii during ions implantation. The inner area has a higher structural disordered region than the outer area where the Raman D-band is activated and lattice structure is preserved. We monitored the introduction of the point defects toward blistered center by analyzing the Raman spectra over the blistered region at low power density. A trend of \(I_D/I_G\) ratio is observed, which has the higher values at the central region of the blister, Figure 4c. The total area contributing to the D peak scattering is proportional to the number of point defects, giving rise to \(I_D/I_G = A/(L_d)^2\) where \(A \approx 100 \text{nm}^2\) (for the low defect density) and \(L_d\) is the size of the crystalline sp\textsuperscript{2} clusters used to quantify the density of defects \((n_D)\) from the following eq \textsuperscript{246,56}.

![Figure 4](image)

**Figure 4.** Affected area of the blister. Area affected by laser power density (0.511 mJ/μm\textsuperscript{2}) shown by (a) adhesion force map. The bright regions represent the higher values of the adhesion force between the tip apex and the surface. (b) Schematic image of the disordered region at two different radii \(R_h\) and \(R_i\) represent inner and outer circles, respectively. (c) The density of defects \((n_D, 1/\text{cm}^2)\) and \(I_D/I_G\) ratio as a function of displacement.

![Figure 5](image)

**Figure 5.** Correlation between topography and functionalization of graphene blister. (a) The optical microscopy of produced blisters (i.e., post-treated) at 100× optical lens. The separation between each blister is 2 μm (b–d). Raman map (resolution 100 nm, sampling data 1581) for \(I_D/I_G\) ratio, 2D peak intensity, and G peak width in contrast to untreated graphene showing distinct signature from untreated single-layer graphene, scale bar is 200 nm. (e) AFM 3D morphology of individual blister along with untreated surface. (f) Raman spectra of D- and G peak as a function of displacement showing a broadening of D and G peak at the blistered region. (g, h) There is broadening and red shift in G peak as a function of displacement. (i) The equivalent topography of the same blister shows maximum altitude of 6 ± 1.5 nm.
Using finite values of $I_D/I_G$ at a fixed wavelength ($\lambda$), size of defect density ($n_d$) as a function of displacement is reported in Figure 4c. It clearly shows that the defect density decreases farther from the blistered region. The sizes of the defect density are in close proximity with simulation results showed for a height $\approx$ 1 nm produced by defect density $\sim$ (10$^6$ cm$^{-2}$).\textsuperscript{20}

**Functionalization of Graphene.** The impact of the laser to alter the local functionalization of 1LG is investigated as a function of displacement. Graphene blisters at Figure 5a are reinvestigated at a high-resolution physical AFM imaging (of 512 $\times$ 512 pixel $\approx$ 2 $\mu$m $\times$ 1 $\mu$m) and Raman mapping (pixel square $\approx$100 nm $\times$ 100 nm) at a lower dose laser energy for a fixed wavelength. The higher values of $I_D/I_G$ are localized at the blistered region as compared to its surrounding. Figure 5b shows the introduction of Raman active disordered region. The decrease in the intensity of 2D peak (Figure 5c) and broadening of G peak width (Figure 5d) appeared at the central region of the blister. The correlation between blister’s topography and Raman peaks was analyzed through line profile over the Raman mapped region, Figure 5d, (e.g., G peak width) and morphology at Figure 5e. The line profile over Raman map is associated to 20 Raman spectra of D, G, D', and 2D peaks taken at a separation of 100 nm, Figure 5f. It illustrates the orderly increment of the D peak intensity at $\sim$1335 cm$^{-1}$, D’ peak at $\sim$1609 cm$^{-1}$ (high-resolution D’ peak is given at Figure S7). The 3D-topography of the given individual blister at Figure 5e is strongly linked to the red shift in the G peak up to 20 cm$^{-1}$ (Figure 5g) and its broadening up to 50.4 cm$^{-1}$ (Figure 5h). All these factors suggest a feature of partial oxidation with structural disorder,\textsuperscript{46,57} which is confirmed by the presence of the carboxylic group in Fourier transform infrared (FT-IR) spectra at Figure S8. These findings were also observed by Johansson et al.\textsuperscript{26} to induce epoxy and hydroxyl groups after irradiation in air conditions.

Raman laser writing can produce an array of 3D graphene in different designs, periodicity, or a complex script. Here, “US” depicts an acronym for University of Sussex with tuned local CPD potential and adhesion force map at Figure S9. The method illustrated here opens new avenues and possibilities in the graphene-based devices to generate localized pseudomagnetic field,\textsuperscript{16} sustainable and efficient nanoscale motor\textsuperscript{1} with different intercalated molecules, localized functionalization surface of different adhesion forces, and polarities for biosensing applications,\textsuperscript{58} bandgap tailoring,\textsuperscript{59} nanofluid, and actuator.\textsuperscript{60} Here, we proposed an effective and simple method to produce 3D structure of graphene with tuned surface properties.

**CONCLUSIONS**

We demonstrated a procedure of producing the localized 3D structure of single-layer graphene by tuning laser dose from Raman spectrometer. Increase in the dose of power density from $51$ to $753$ mJ/$\mu$m$^2$ surges the volume of the blister. The lattice expansion of carbon atoms increases the gap between 1LG and silica interface and deposition of the oxygenic functional group, which alters the photoinduced chemistry of graphene and is outwardly shown through scanning probe techniques (AFM and KPFM). The introduction of the functional group and higher contact area of deformation causes the higher force of adhesion at the blistered region. The CPD contrast showed the lower work function at the blistered region due to the prohibition in p-doping from the silica substrate, which resembles the wrinkle and graphene nanobubble characteristics. The work function decreases with increasing gap between graphene and the silica substrate. The internal pressures in the blisters are significantly lower than the graphene nanobubble. We did not observe any laser-induced ablation of graphene or any vacancy in the irradiated region, though the structural disorder was received. The decrease in intensity ($I_D/I_{2D}$) with increasing power density shows a reduction in the domain size of the sp$^2$ breathing mode. Phonon softening observed in high-resolution Raman mapping toward the central part of the blistered region shows lattice expansion of the carbon atoms. The defect density measured from $I_D/I_G$ decreases as it goes away from the laser-affected region ($R_0$ to $R_0$), affirmed an annular shape for the distribution of defects. Our results reveal the effectiveness of the Raman spectrometer for the modifications and functional analysis of graphene layer.

**MATERIALS AND METHOD**

Single and bilayer graphene was produced by mechanical exfoliation of graphite (HOPG) and deposited on a silica substrate (300 nm oxide thickness). Deposited graphene on silica substrate was cleaned in an ultrasonic bath in the sequence of isopropanol and deionized water each for 30 min, then heat-treated in vacuum for 3 h at 200 °C.

Raman spectroscopy (spectral resolution 0.3 cm$^{-1}$) has been carried out at 100X objective lens using two lasers. Laser (1): 532 nm (type: solid state, model: RLS3250) and Laser (2): 660 nm (type solid state, model: RL660C100) at different powers (mW). The maximum power used from the laser source for 532 nm is 50 mW and for 660 nm is 100 mW. Variations in the laser powers were carried through ND filters. The higher power densities of the laser were achieved by increasing the time duration (10, 15, and 20 s) of the irradiation while keeping the output power consistent. It should be noted that there is a loss of 5–10% of the power density during reflection from in-built mirrors in the Raman spectrometer. The laser irradiation was repeated over four different single-layer graphene flakes of reproducible textured amplitudes, which are reported at Figures 1, S5, S6, and S8.

AFM characterization was performed with a Bruker Dimension Icon, positioned in an insulated box over an antivibrant stage to minimize environmental noise and building vibrations. Contact potential difference (CPD, volts) and mechanical data were measured from advanced operation mode of PF-KPFM (PeakForce-Kelvin Probe Force Microscopy) and PF-QNM (PeakForce-Quantitative nanomechanical), respectively. PeakForce is a Bruker’s proprietary mode, which allows the collection of both types of information on a single acquisition. Amid the PF-KPFM operation, the standard PeakForce procedure during the first pass of each line scanning, with the tip softly (<1 nN) tapping on top of the sample. In this condition, it gathers topography and mechanical properties of the sample. In the second pass over the same scanning line, the cantilever lifted from the surface up to 10 nm distance to collect CPD data. The proprietary Scansanyst algorithms simplify engaging and parameters for setting procedures, optimizing in real time Scan Rate, PeakForce set point and Feedback Gain settings but was in general turned off during capture once the optimal parameters were achieved to avoid inconsistency forth the final picture.
This type of characterization has been performed using every time the same mounted tip to guarantee as much consistency as possible in the Kelvin Probe data, highly variable, being dependent not only on sample properties but also on the cantilever used in the measurement. A PFQNE-AL tip was chosen for these reasons, being the gold standard of Bruker’s tip for KPFM characterization. It is a soft silicon–nitride tip with 5 nm nominal tip diameter, 300 ± 100 kHz resonant frequency, and 0.8 ± 0.2 N/m spring constant, optimized for electrical modes and with a proprietary reflective coating on the backside. Thermal Tune calibration was performed before each imaging session to verify consistency of resonance frequency and stiffness of the cantilever. The stiffer cantilever has been avoided to minimize the physical damage to the blister.

High-resolution mechanical information acquired by Peak-Force-QNM (Quantitative Nanomechanical Mapping) for Figure 2a, S3, S5, and S6. This mode works with the same procedure described in the first pass of the PF-KPFM. A silicon nitride (model: Scanasyt-air tip) was chosen in these cases, having a nominal ultrasharp tip of 2–3 nm that allows maximum resolution while minimizing contact area, ensuring better consistency and resolution of mechanical information, in particular adhesion force map. The resonance frequency and the stiffness of the cantilever are measured as 70 ± 25 kHz and 0.2–0.8 N/m (nominal 0.4 N/m), respectively.

The adhesion force measurement carried out in PF-QNM approach was operated in a true contact mode at relative humidity (30–35%) at room temperature, the blisters were bearing a certain contact load and may have a larger contact deformation. Thus, the elastic deformation is performed by modest pressing (set point <1 nN) over the blistered region. It is a distance between “jump to contact” and the elastic penetration. Prominent deformation at the blistered membrane leads to the higher contact area between the AFM tip-apex and the deformed region, which is responsible for higher adhesion forces “pull-out”.

The FT-IR measurements were performed using PerkinElmer Spotlight 400 FT-IR Microscope System. The system uses dual-mode single point and mercury cadmium telluride array detector standard with InGaAs array option for optimized NIR imaging. All the measurements were done using the mid-IR (4000–500 cm\(^{-1}\)).

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.8b02815.

Thickness verification of graphene layers and absence of D peak confirmed by Raman spectroscopy; optical image of blister produced at different locations in 1LG; topography of individual blister with adhesion and deformation map produced at a laser dose of 511 mj/\(\mu m^2\); inducing normal force on blisters at different set points with peakforce schematic mechanisms; impact of laser at the edge region of graphene; red shift and broadening of G and 2D peaks with increasing laser dose (PDF)

AUTHOR INFORMATION

Corresponding Authors

*E-mail: m.tripathi@sussex.ac.uk (M.T.).

*E-mail: a.b.dalton@sussex.ac.uk (A.B.D.).

ORCID

Nicola Maria Pugno: 0000-0003-2136-2396

Alan B. Dalton: 0000-0001-8043-1377

Notes

The authors declare no competing financial interest.

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