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Charging and discharging of graphene in ambient conditions studied with scanning probe microscopy

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By means of scanning probe microscopy we are able to inject charges in isolated graphene sheets deposited on SiO_2/Si wafers and characterize the discharge induced by water in controlled ambient conditions. Contact potential differences between the graphene surface and the probe tip, measured by Kelvin probe microscopy, show a linear relationship with the tip bias during charge injection. The discharge depends on relative humidity and decays exponentially with time constants of the order of tens of minutes. We propose that graphene discharges through the water film adsorbed on the SiO₂ surface. © 2009 American Institute of Physics. [DOI: 10.1063/1.3149770]

Charge injection combined with scanning probe microscopy (SPM) technique provides a powerful tool to study the electrostatic properties of objects in the nanoscale. It allows to measure electrostatic forces in the vicinity of the charged object to gain insight into the distribution of charges within the object, as demonstrated, e.g., in carbon nanotubes¹ and silicon nanoparticles.² Of equal interest is the use of these techniques to learn about the electrostatic interaction between the object under study and its environment. This is of fundamental importance in surface science. The electronic properties of nanosized objects are strongly affected by the substrate they are deposited on and by molecular adsorbates, which are both difficult to control and require careful characterization. At present, an example attracting huge interest is graphene on silicon oxide. Graphene, a two-dimensional crystal of carbon atoms arranged in a triangular lattice,³ is entirely exposed to its environment, which critically limits its performances as an electronic material.⁴ In particular, water and other molecules adsorb on the surface of graphene and can modify its carrier concentration. This property has been used to detect individual gas molecules adsorbed on graphene.³

In this letter we present an experimental procedure to charge individual few layer graphene (FLG) films deposited on a SiO_2/Si wafer using SPM techniques. Once charged the FLG films are characterized using scanning polarization force microscopy (SPFM) and Kelvin probe microscopy (KPM). FLG films are found to discharge into the Si substrate through the water film adsorbed on the SiO₂ surface.

All the experiments shown here were carried out at room temperature with a commercial atomic force microscope (Agilent 5500) modified to operate in the SPFM mode. The SPFM operational mode has been reported in detail elsewhere^{6,7} and only a brief description is given here. SPFM is an electrostatic SPM mode in which a conductive tip is set at about 10–20 nm from the surface and biased to a few volts.⁸ Together with KPM, it allows one to obtain simulta-

neously information about topography, polarizability, and contact potential difference (CPD) between the tip and the surface.

The microscope head was enclosed in a glove box to control the relative humidity (RH) of the environment. FLG films on silicon wafers were prepared using the conventional micromechanical exfoliation technique,⁹ where a flake of bulk Kish graphite is repeatedly cleaved with an adhesive tape and pressed down onto a silicon wafer coated with 280 nm thermally grown silicon oxide.

Charge injection is performed as follows. The silicon substrate is connected to ground during the experiment. An image in contact mode of the FLG film is taken with the tip grounded [Fig. 1(a), top left]. Then, the tip is placed above the center of the FLG film, brought into contact with it and a bias (V_{ini}) is applied to the tip within the -10 to 10 V range [Fig. 1(a), top right]. After a few minutes the tip is retracted to about 200 nm from the surface, keeping the tip biased [Fig. 1(a), lower left]. With this procedure the FLG film becomes charged acquiring an initial electric potential V_0 , different from V_{inj} , which evolves with time as discussed below. Once the tip is retracted from the surface, V_{inj} on the tip is switched off and a dual SPFM+KPM image is taken [Fig. 1(a), lower right]. Using this method we can guarantee noncontact conditions during all throughout the experiment on the FLG film and avoid the formation of a water neck between the tip and the sample¹⁰ that would perturb the discharge process. An image in contact mode with the tip grounded can be performed to discharge the FLG, as can be seen by comparing KPM images before charging and after discharging.

Figure 1(b) shows SPFM (top left) and KPM (top right) images of a few microns wide FLG sample. In the SPFM images the FLG films show a high contrast (apparent height) due to the huge difference in the dielectric constant (ε) between SiO₂ and the FLG films.¹¹ The KPM image, taken simultaneously with the SPFM image, shows a CPD slightly less positive in the FLG region compared to the SiO₂ substrate, in agreement with recent results.¹² In Fig. 1(b) we show KPM images of the FLG film after charging it at RH

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FIG. 1. (Color online) (a) Scheme of the experimental charging and measurement procedure used. (b) SPFM (top left) and KPM (top right) images of a FLG film before charging at low RH. Bottom: KPM images of the same film after charging to (left) -8 V and (right) +8 V. (c) Δ CPD CPD after charge injection-CPD discharged) vs voltage applied to the tip during the charging process (V_{inj}) measured from KPM taken after charge injection. Δ CPD appears to be proportional to V_{inj} . Different symbols denote different experiments with different tips and different samples. All experiments were performed at RH < 5%.

<5% with -8 V (down left) and +8 V (lower right), respectively. In the KPM images the FLG film charged with +8 V (-8 V) show a bright (dark) CPD contrast.

In the following, we investigate the charge and discharge of FLG films by measuring the CPD between the FLG film and the tip. We first characterize our experimental setup by measuring the dependence of CPD on V_{inj} . We then study the influence of humidity in the environment, which brings us to consider various discharge mechanisms. In Fig. 1(c) we plot the difference between CPD (Δ CPD) measured on three different FLG films before and after charging as a function of $V_{\rm inj}$ at RH < 5%, showing a linear behavior with a slope $s \sim 0.1$ ($\Delta CPD \approx s \cdot V_{ini}$). Note that after each charge injection and measurement of Δ CPD graphene is brought back to its initial state (by discharging) in order to avoid any charge accumulation effect. Different contributions can influence the charging process and the subsequent Δ CPD measurements: (i) a poor tip-FLG contact, (ii) the geometry of the tip-FLG system, (iii) the discharging of the FLG film into the environment and (iv) the loss of some of the injected charge during the retraction of the tip. In order to check the tip-FLG contact, the contact time between the tip and the FLG film has been increased up to 30 min (the usual contact time was $\sim 1-3$ min). Additionally, the increase in the load applied to the tip during the charging process has been verified. Both parameters had no effect on the measured Δ CPD. This indi-



FIG. 2. (Color online) (a) Time evolution of KPM images of a FLG film charged at +8 V and at 30% RH. (b) Cross section profiles of the same images.

cates that the electrical contact between the FLG film and the tip was good enough. In addition, the high reproducibility of the obtained slope over several independent experiments rules out an uncontrolled loss of charge during injection.

The slope *s* strongly depends on the geometry of the system represented by the tip and the FLG film. We measured $s \sim 0.1$ for FLG films with areas ranging from 6 to 30 μ m². Measurements performed on much larger FLG films (in the order of the tip dimensions, 200–400 μ m²) still show a linear behavior but with $s \sim 0.3$. Thus, to control quantitatively the charging process, one should take into account the influence of the full cantilever-tip-graphene geometry.

CPD on a charged FLG film is not constant with time and FLG films discharge with a decay rate that strongly depends on the RH of the environment. In Fig. 2(a) we show the time evolution of KPM images of a FLG film charged at 8 V and at 30% RH. The images show how the positive charge spreads over the silicon substrate while the initially confined FLG film discharges. The evolution of a cross section is shown in Fig. 2(b). The CPD on the FLG film decreases and CPD values on the SiO₂ surrounding the film increase due to positive charges that are moving away from the FLG film and toward the substrate. It is well known that a clean SiO₂ surface rich in OH superficial groups is a very hydrophilic surface and a few monolayers of water molecules are expected to be adsorbed on the surface for a wide RH range.¹³ Thus, the main discharge process would be through water molecules adsorbed on the FLG film itself¹⁴ and the water film on the SiO₂ substrate. At t=28 min in Fig. 2 the CPD on the FLG and the substrate area films surrounding it level at the same value. From this time on the entire CPD profile lowers with time with minimal changes in its shape. This behavior clearly suggests that the slower discharge of the water film surrounding the FLG film is limiting the faster migration of charges from the FLG film to the water film.

The time evolution of the mean CPD measured on the FLG film at different RH conditions exhibits an exponentiallike decay [Fig. 3(a)]. Only the data taken at 50% RH decays to the CPD value of the FLG film before charge injection



FIG. 3. (Color online) (a) Equivalent *RC* circuit modeling the discharge of the FLG film. Some of the components of the circuit can be neglected at low humidity (R_2) and at high humidity ($C_{\rm H_2O/SIO_2}$). (b) Evolution with time of the FLG film CPD value after charging it at +8 V for 10%, 30%, and 50% RH. Exponential decays used to fit the data [Eqs. (1) and (3)] are shown and the CPD value of the discharged FLG film is marked (~1.4 V).

(~1.4 V for that particular sample) for the times measured. Very similar decays were obtained for both positive and negative voltages.

In order to explain our results we model our system as an equivalent circuit formed by capacitors and resistances [Fig. 3(b)]. One capacitor (C_{FLG}) is formed by the FLG film and the SiO₂/Si substrate while the other (C_{H_2O/SiO_2}) is formed by the water film adsorbed on the SiO₂ and the SiO₂/Si substrates (the Si substrate is grounded). R_1 represents the resistance that limits the flow of charges on the FLG film to the water film. R_2 models the discharge of the water film mainly through the diffusion of the water molecules on the SiO₂ surface. The solution of the differential equations of the circuit is in the form

$$V_{\rm FLG}(t) = A_1 e^{m_1 t} + A_2 e^{m_2 t},\tag{1}$$

where

$$m_{1,2} = -\frac{1}{2} \left[\frac{1}{C_1 R_1} + \frac{1}{C_2 R_1} + \frac{1}{C_2 R_2} \right] \pm \frac{1}{2} \sqrt{\left[\frac{1}{C_1 R_1} + \frac{1}{C_2 R_1} + \frac{1}{C_2 R_2} \right]^2 - 4 \frac{1}{C_1 C_2 R_1 R_2}},$$
(2)

and $V_{\text{FLG}}(0) = A_1 + A_2 \equiv V_0$.

At 10% RH we can simplify the model by considering that $R_2 \rightarrow \infty$ and can thus be ignored. We then obtain that

$$V_{\rm FLG}(t) = \frac{C_T V_0}{C_{\rm H_2O/SiO_2}} + \frac{C_T V_0}{C_{\rm FLG}} e^{-t/R_1 C_T},$$
(3)

where C_T stands for the total capacitance $(1/C_T=1/C_{FLG} + 1/C_{H_2O/SiO_2})$ and V_0 is the initial voltage at the C_{FLG} capacitor. Equation (1) shows the voltage decay for the FLG film going from V_0 to $V_F = C_T V_0 / C_{H_2O/SiO_2}$ with a time constant R_1C_T . Both R_1 and C_T depend on RH. The increase in water molecules on the SiO₂ surface in contact with the FLG film will increase the ratio of charge transfer between the FLG and the water film, thereby lowering the resistance in our model. The amount of water adsorbed on SiO₂ would clearly affect C_{H_2O/SiO_2} and water adsorbed on or below the

FLG film could also modify C_{FLG} . The experimental data fitted with this model is shown in Fig. 3(a) with a time constant of \sim 50 min. At high humidity there is a high diffusion of charges on the surface and we can consider that R_2 is so small that no charge is accumulated in $C_{\text{H}_2\text{O/SiO}_2}$. The equivalent RC circuit is thus simplified by a circuit composed of a capacitor C_{FLG} that discharges through a resistance R_1 . The CPD value decays exponentially to its precharging value [1.4 V for RH=50% with a time constant of ~ 10 min, in Fig. 3(a)]. At intermediate humidity the discharge of the FLG film depends strongly on the charge accumulated in the water surrounding it and how fast these charges diffuse away from the FLG film. In our model that means that none of the processes are dominating the discharging and both R_1 and R_2 must be taken in account. Data taken at 30% RH [Fig. 3(a)] can only be fitted then using the two exponential decays shown in Eqs. (1) and (2).

In conclusion, we have presented an experimental procedure to charge individual FLG films deposited on a SiO_2/Si wafer using SPM. We have found that CPD measured on the FLG film using KPM shows a linear relationship with the voltage applied to the tip during the charge injection for both negative and positive voltages ranging from -10 to +10 V. Discharge of FLG films has been studied at different RH conditions. We have found that CPD follows RH dependent exponential decays with time. We have finally shown that FLG films discharge mainly through the water layer adsorbed on the SiO₂ surface.

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