The Effect of Humidity on the Interlayer Interaction of Bi-layer Graphene

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The lubricating ability of graphite largely depends on the environmental humidity, essentially the amount of water in between its layers. In general, intercalated molecules in layered materials modify their extraordinary properties by interacting with the layers. To understand the interaction of intercalated water molecules with graphene layers, we performed Raman measurements on bi-layer graphene at various humidity levels and observed an additional peak close to that of the low-frequency layer breathing mode between two graphene layers. The additional peak is attributed to the vibration between an intercalated water layer and the graphene layers. We further propose that the monolayer coverage of water increases between bilayer graphene with increasing environmental humidity while the interaction between the water layer and graphene layers remains approximately unchanged, until too much water is intercalated to keep the monolayer structure, at just over 50% relative humidity. Notably, the results suggest that unexpectedly humidity could be an important factor affecting the properties of layered materials, as it significantly modifies the interlayer interaction.

I. INTRODUCTION

Graphite, the most prominent layered material, has been used as a good lubricant for about 200 years. However, an abnormally high wear rate of graphite brushes in electrical machines aboard aircraft during World War II was reported.1 Later in the 1960s, NASA found that graphite lost its lubrication in space.1 It is not the low pressure, but the low humidity, that is primarily responsible for the reduced lubricating ability at high altitude or in space, as a weakened interlayer interaction (by intercalated water as reasonably believed) is essential for the easy shear parallel to the graphene hexagonal planes.2 The importance of a weakened interlayer interaction for graphite to be lubricant is again confirmed by later work successfully increasing the lubricating ability of graphite under vacuum by introducing functional groups to it.3,4 A complete understanding of this has not been achieved as it is not yet clear how the interlayer water interacts with the graphene layers, while some may still doubt if water intercalates at all.

Layered materials containing different numbers of layers can now be produced, either by exfoliating from the bulk,5 or by stacking one layer onto another.6 Structures made from different numbers of layers may possess unique properties.7 Intercalated molecules further modify these properties. This vast research area focuses in general on three questions — how, and how many, intercalating molecules can be introduced between the layers, and what effect they have on pristine layered material. The properties of layered materials largely depend on their interlayer interaction.8 It is therefore important to quantify the effect of intercalated molecules on this interaction.

In our work we start with a simple system, bi-layer graphene (the simplest layered structure) with intercalated water (the most common polar molecule), while the amount of water is varied by adjusting the humidity (one of the most common atmospheric conditions).

Much work has been done on water in between graphene oxide layers (GO), as it is hydrophilic — water easily comes between the layers and attaches to them via non-covalent bonds.9 Early work reported interlayer spacings of 0.63 nm for dry GO10 and 1.2 nm for hydrated GO.11 The amount of intercalated water can be most accurately obtained by measuring the atomic ratio of carbon to oxygen.12 Nair et al. found that permeation of water in GO is unimpeded, and attributed that to a low-friction flow of monolayer water (held by hydrogen bonds) between GO layers.13 Cerveny et al. also reported a water monolayer between GO layers, and it expanded the GO from the interlayer spacing at 0.57 nm to 0.79 nm, when the water content went from 0 to 25 wt%.14 25 wt% interlayer water corresponds to approximately 1 water molecule in every 2 hexagons of carbon. Further expansion to 1.1 nm was observed at 100% relative humidity (RH).15 Kim et al. measured the diffusivity and dielectric constant of intercalated water in GO layers and found values for both an order of magnitude less than in bulk water,16 which is further evidence that water has a novel structure when contained between GO layers.

Water is also an essential part of the structure of graphene layers.17 Moreover, adsorption of water from the air makes graphene p-type doped.18 Yavari et al., by relating the resistivity of graphene to the humidity, found that this doping was stronger when the humidity was increased, but in general it was weak and can be reversed.19 Note that this doping was on the graphene-air interface and it is not clear what doping intercalated water might induce.

Techniques used to study water between graphene layers include broadband dielectric spectroscopy (BDS, for
and was first measured at 127 cm\(^{-1}\).

**III. RESULTS AND DISCUSSION**

The bilayer graphene samples were prepared on a Si wafer by exfoliation and the fast pick-up technique.\(^\text{30}\) The pick-up was at 100 °C and the release was at 160 °C. The diameter of the overlapping area of the two graphene layers is about 10 \(\mu\)m. The sample was placed in a chamber with a humidity sensor, described previously.\(^\text{31}\) Dry nitrogen flows through the chamber to decrease the humidity while wet nitrogen (bubbled through de-ionised water before entering the chamber) is used to increase it. We performed the room-temperature non-polarised Raman measurements with a Renishaw inVia spectrometer in the backscattering geometry with a confocal microscope. The system has a resolution of 1.3 cm\(^{-1}\). We used a 531 nm laser. The laser power on the humidity chamber was 1.03 mW and there was a further 15% reduction of power by the chamber window. The size of the laser spot was about 2 \(\mu\)m. We should point out that the LBM is only observable under resonance, so the laser wavelength required is determined by the stacking angle of the two graphene layers. The transfer technique we use here gives random angles. With only the fixed 531 nm laser line, we prepared several samples and studied one in which the LBM is in resonance with the laser.

We started the measurements at a low RH of 7%. At each RH value throughout the experiment we collected two spectra, one in the low wavenumber range for the LBM and CM, the other in the high range for the G and 2D modes. During the collection of a spectrum (20 – 400 s), slight fluctuations of the RH occurred but did not exceed 1%. We increased the RH from 7% to 22% and then to 37%. Changes in the spectral profile were observed at 37%, so we decreased the RH back to 22%, then increased it again to 30% and 39%, and further increased it to 53%. Losing the signals of the interlayer modes at 53%, we reduced the RH all the way down to 5%, and finally brought it back to 16%.

The spectra are presented in Fig.1. They are vertically shifted for clarity, in the order they were collected, from bottom to top. The spectra in the low and high frequency ranges, collected at the same RH point, are shown at about the same horizontal level (there is a slight difference in the RH values for the spectra on the left and the right of Fig. 1 because of small fluctuations in RH during the collection of spectra, as noted). The key facts are very clear from the spectra, that the intensity ratio of G to 2D significantly decreases at 53 % RH, and an additional peak close to the LBM clearly appears from 22 %.

Spectra were fitted using maximum likelihood estimation. We first subtract the background. For the G and 2D modes, the background of the spectra is relatively flat below the strong Raman peaks. From 21.5 % RH (blue),
we notice a background which then gets stronger. To fit the background, we select a few points on the spectra away from the peak positions and fit them with a 6-term polynomial. For the CM and LBM, although these peaks are intense at low RH, it is challenging to obtain an accurate fit for them as they are close to the cut-off edge of the laser band-reject filter. The background in the range of the CM may consist of contributions from the laser, the edge of the filter, and luminescence. It is difficult and unnecessary to sort every component out. What we do here is to collect two spectra of monolayer graphene in the humidity chamber at 10 % and 40 % RH. The line shapes of the two spectra almost overlap, suggesting that the humidity has no effect on the monolayer graphene. The difference between the spectra of monolayer and bilayer graphene is what we are interested in, as it arises from the presence of the second layer. We use the spectra of monolayer graphene at 40 % RH as background. After subtracting the background, we compare the fittings of various numbers and shapes of peaks by their residuals, and by their values of the Bayesian information criteria (BIC). We show an example of the best fitting for the spectrum collected at 6.2 % RH, in Fig. 2. Here we fit the CM and LBM peaks by two pseudo-Voigt peaks of 57.6 % Gaussian at 59 and 95 cm\(^{-1}\), respectively.

We first focus on the high frequency range. After subtracting the background, we integrate the spectra (data, not fits) over the G and 2D peaks range, and plot the integrated area ratio of G to 2D in Fig. 3.

We then fit the spectra and find that the GM profile at most RH points consists of two peaks, one at around 1580 cm\(^{-1}\), the other at 1590 cm\(^{-1}\). There is an additional peak in the range of 1610–1620 cm\(^{-1}\) at 38.5% and 53.4% RH. These peaks are not resolved in Fig. 1, but are obtained objectively by the fitting routine described above. No consistent shift of G or 2D peaks with RH is
The results are consistent with the amount of water in between the bi-layer graphene changing its interlayer interaction, and being sensitive to the environmental RH, though there is no definite relation. We would like to point out that bi-layer graphene behaves as two monolayers at just 53.4% RH, which can be naturally achieved in the atmosphere of many labs. Additionally, there is an interesting pattern that the ratio slightly goes up and then abruptly drops and it repeats three times — data points 1→2→3, 4→5→7, and 8→9, as shown in Fig. 3.

We now consider the CM and the LBM. We obtain the frequencies and widths of the peaks from the best fits. Fitted parameters for the CM are plotted against RH in Fig. 5. The uncertainties in peak positions from fitting are within the system resolution of 1.3 cm$^{-1}$, while the uncertainties in the width are $\sim 2$ cm$^{-1}$, at various RH values. The sequence of the measurement is numbered in accordance with the G and 2D modes in Fig. 3. No CM is observed at 38.5 % as well as 52.7 % RH. The positions of the CM at various RH values are close, except the very first point at 6.2 %. No consistent change of width is found.

The additional LBM peak is the main finding of this work. In Fig. 6 we plot the peak positions and widths of the LBM, and the integrated-area ratio of the higher LBM to the lower, against RH.

The additional peak appears from the second RH point at 22.4%. We attribute this peak to the vibration between the intercalated water layer and graphene layers, from the following considerations. First, the decreasing G to 2D ratio with increasing RH indicates the weakened interaction between two graphene layers. We therefore expected a redshift of the LBM. Second, the frequency of the LBM of bi-layer graphene does not change much after water intercalation from 22.4 % RH. This suggests that the bi-layer graphene was only partially filled by water. Third, the additional peak is at higher frequency than the original LBM of bi-layer graphene, indicating that it is not from vibrations of two graphene layers but perhaps involves something of smaller mass. Also, the nearly unchanged position of this additional peak at various RH values indicates that it corresponds to an interaction between layers that does not change much with increasing amounts of intercalated water. On both counts, it can be attributed to the water-filled regions. This also provides a reasonable interpretation for the recent XRD measurements on graphite showing unchanged interlayer spacing under various humidity conditions$^{32,33}$ that small amounts of water intercalate between graphene layers, increasing the coverage of water-filled regions, while not expanding the interlayer spacing with increasing humidity. Fourth, adsorption of water on the graphene makes it p-type doped. No evidence shows that doping shifts the LBM. With an increasing amount of water on both sides of the top graphene layer, one would expect an increasing level of doping. Again the approximately unchanged position of the additional peak at various RH values suggests that it is not just a doping-modified LBM.
FIG. 5. (a) Frequency and (b) width of the fitted peak for the CM of bi-layer graphene are plotted against RH. Black solid squares denote the spectra collected at a higher RH point than all the previously collected spectra. Blue open circles denote the spectra collected after reducing the RH from 37.4%, and the blue open triangles are for those collected after reducing the RH from 53.4%. In both cases, the humidity was decreased and then brought up again. The sequence of the measurements are labelled in accordance with Fig. 1. The missing number 6 refers to those of no observable CM signal. The uncertainties in the frequency from fitting are within the system resolution of 1.3 cm$^{-1}$, while the uncertainties in the width are $\sim 2$ cm$^{-1}$.

of graphene layers. In fact, the level of doping is tiny here. It is known that the frequency and width of GM are sensitive to doping. There is no observable shift of the G and 2D modes with humidity in Fig.1 and Fig.4 shows that the frequency and width of the GM are randomly scattered with humidity almost within the very small uncertainty of data. Fifth, the reported monolayer of water between GO layers$^{13}$ seems relevant here. The additional peak in the frequency from fitting are within the system resolution of 1.3 cm$^{-1}$, while the uncertainties in the width are $\sim 2$ cm$^{-1}$.

FIG. 6. (a) Frequency and (b) width of the LBM of bi-layer graphene are plotted against RH. At an RH where there are two LBMs, (c) the integrated area ratio of the LBM from higher to lower frequency is plotted against RH. Black solid squares and diamonds (for the LBM of higher frequency) denote the spectra collected at a higher RH point than all the previously collected spectra. Blue open circles denote the spectra collected after reducing the RH from 37.4%, and the blue open triangles are for those collected after reducing the RH from 53.4%. In both cases, the humidity was decreased and then brought up again. The symbols with a cross are for the LBM of higher frequency. The uncertainties in the frequency and width from fitting are within the system resolution of 1.3 cm$^{-1}$. The sequence of the measurements are numbered in accordance with Fig. 1. The missing numbers refer to those of no clear LBM signal.

amount of water is too much to maintain the layer structure at 52.7 %, where all the interlayer modes disappear. Finally, we apply a linear chain model to obtain a force constant for this additional layer breathing mode. The chain consists of graphene, water and graphene. We consider the nearest-neighbour interaction only. We assume that there is one water molecule in the area of a carbon hexagon, in terms of the density of the water layer, as it has to be condensed enough to form a layer, and held by hydrogen bonds following Nair et al.$^{13}$ (if we consider the
bonding of water in-between two graphene layers to be the same as between GO). We insert $i = 1$, $N = 3$, the reduced mass (of water and carbon) $m = 3.26 \times 10^{-27}$ kg Å$^{-2}$, and the measured LBM frequency to Eq. 1 and obtain the force constant $\alpha_{Gr-water} = 37.9 \times 10^{18}$ N m$^{-3}$, about 1/3 of the force constant between graphene layers. This is a reasonable value for the force constant. All of the LBM peaks are very narrow (width below $\approx 10$ cm$^{-1}$), and there is no abrupt change in the width of fitted peaks, further validating the reliability of the presented results, on which the above discussion is based. The integrated area ratio of the two LBM changes with RH but no clear relation is observed.

IV. CONCLUSION

In summary, we measured the high frequency G and 2D modes, and the low frequency interlayer CM and LBM of bi-layer graphene at various humidity levels. With increasing RH, we observe a decreasing intensity ratio of the G to 2D mode, a downshift of the CM and notably an additional LBM. We conclude that intercalated water molecules form a layer, the interaction of which with the graphene layers is about 1/3 of that between pristine graphene layers. With increasing amounts of water, the interlayer interaction of the bilayer graphene is weakened, while the interaction between graphene and the water layer remains nearly unchanged, until too much water is intercalated to maintain the layer structure at over 50 % RH. Water molecules can be introduced in between graphene layers at relatively low levels of the humidity, and by increasing the humidity the bi-layer graphene behaves as two monolayers, at just over 50 % RH. This suggests that atmospheric humidity could be a crucial parameter in many laboratories affecting the results of graphene-related research and applications.

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