Few-layer graphene on SiC, pyrolitic graphite, and graphene: A Raman scattering study

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The results of micro-Raman scattering measurements performed on three different “graphitic” materials: micro-structured disks of highly oriented pyrolytic graphite, graphene multi-layers thermally decomposed from carbon terminated surface of 4H-SiC and an exfoliated graphene monolayer are presented. Despite its multi-layer character, most parts of the surface of the graphitized SiC substrates shows a single-component, Lorentzian shape, double resonance Raman feature in striking similarity to the case of a single graphene monolayer. Our observation suggests a very weak electronic coupling between graphitic layers on the SiC surface, which therefore can be considered to be graphene multi-layers with a simple (Dirac-like) band structure.

The interest in the properties of graphite-like allotropes of carbon has recently been rekindled when its simplest form, graphene, a mono layer of carbon atoms arranged in a honey comb lattice, was experimentally identified and readily produced by a simple exfoliation technique. The linear dispersion at the K and K’ points of the graphene band structure gives rise to a relativistic-like behavior of free carriers that has many implications for the observed quantum mechanical effects, such as the half-integer quantum Hall effect, and has stimulated considerable research over the past few years. Alternatively, thin graphitic layers can be epitaxially grown on 4H-SiC substrate, by the thermal decomposition of either Si- or C-terminated surface. In this case, the material, herein referred to as few-layer graphene (FLG) on SiC, may be composed of one or many graphene layers, depending on the growth conditions. Although much interesting physics has emerged from the studies of exfoliated graphene structures, the epitaxy of graphene seems to be the most viable production route for electronic applications.

In this letter, we present the results of micro-Raman scattering studies of few-layer graphene on C-terminated surface of SiC (FLG-C-SiC) with a large number of layers, and compare them with the data obtained on disks made of highly oriented pyrolytic graphite (HOPG) with different thicknesses and with the spectra of an exfoliated graphene monolayer. Attention is focused on the form of the so called D̃-band, a well established feature in the Raman scattering spectra of graphite and graphene. Its single-component form is the fingerprint of the simple electronic bands in a graphene monolayer, while a multi-component form is the signature of the complex band structure of multi-layers with Bernal-type (graphite-like) stacking.

Apart from some residual inclusions, most of the surface of the FLG-C-SiC probed with micro-Raman spectroscopy was found to show a single component D’ feature, as for an exfoliated graphene monolayer but in striking contrast to the double-component D̃-feature observed in HOPG disks. This observation supports the view that FLG-C-SiC is a system which displays single band (Dirac-like) dispersion relation of electronic states characteristic of decoupled graphene layers.

The structures used for experiments were disks of HOPG with different thickness, an exfoliated graphene monolayer, and, three FLG-C-SiC samples with a different number of layers. HOPG disks (20-μm in diameter) have been prepared by mechanical cleavage with a pulled glass tip of cylindrical ZYA grade HOPG mesas and by deposition via micro-manipulation onto a silicon substrate with a 100-nm or 300-nm SiO2 layer. Graphitic disks could be further thinned by exfoliation down to a thickness ranging between 2 and 20 nm. A graphene monolayer was prepared by exfoliation of freshly cleaved ZYA grade HOPG on a silicon substrate with a 300 nm thermal oxide substrate as described in the literature. Sub-20 nm thick graphitic disks and the graphene layer were subsequently characterised/identified by AFM. FLG-C-SiC structures were thermally decomposed from 4H-SiC substrates following the procedure detailed in Ref. 5 and 6. Two of them (sample 1 and 2) are 5-10 layers thick while sample 3 consists of 70-90 layers, as determined from the intensity ratio of the Si 92-eV and C 271-eV Auger peaks. Raman scattering spectra have been measured using a confocal microscope with 2μm spatial and 1 nm spectral resolution. Experiments were performed at room temperature using the 632.8 nm line of a HeNe laser.

We start the discussion of the experimental results by presenting the characteristic Raman-scattering spectrum of bulk graphite, obtained here for a thick (~100 nm-high) disk of HOPG (see Fig. 1). The G band at 1582 cm⁻¹ is due to first order (one phonon) Raman scattering process. It is characteristic of sp² hybridization and involves the in-plane optical phonon E₂g near the Γ point of the phonon band structure. As a rule, the overtone (two-phonon process) of the G-band, labeled as the G’-band in Fig. 1 is much less intense. The first order Raman scattering process due to characteristic zone boundary phonons (at 1325 cm⁻¹) is forbidden in defect-free structures and is barely visible in our spectra. However, these
The phonon wave vector implied in both scattering processes is indicated in red.

The Raman scattering process, schematically shown in the inset of Fig. 1, is inherently sensitive to the multi or single-band character of the dispersion relations of the electronic states. In consequence, the D'-feature is known to be composed of at least two-components in graphite (as seen in Fig. 1) and also in other graphene multi-layers with Bernal stacking due to the multi-band character of the electronic states in these systems. In contrast, the unique single Lorentzian form of the D'-feature is the signature of a system with a single-band electronic dispersion. The D' band in the form of a single component peak. We note, however, the existence of some inclusions or spots on the sample surface, which are visible under an optical microscope, and which give rise to a different spectrum characterized by a distinct, more complex shape of the D'-band (see trace (b') in Fig. 3). When scanning the laser spot over the FLG-C-SiC surface we find that most of the sample gives rise to the characteristic spectrum of the graphite monolayer (trace (e) in Fig. 3) and also in other graphene multi-layers with Bernal stacking due to the multi-band character of the electronic states in these systems. In contrast, the unique single Lorentzian form of the D'-feature is the signature of a system with a single-band electronic dispersion, such as the representative graphene monolayer.

The Raman scattering spectra of these samples show the characteristic G and D' bands on the background of a more or less pronounced signal from the SiC substrate. We find that most of the sample gives rise to the characteristic Raman scattering spectra which are shown in Fig. 3, for three different samples. The important feature of these spectra is the appearance of the D' band in the form of a single component peak. We note, however, the existence of some inclusions or spots on the sample surface, which are visible under an optical microscope, and which give rise to a different spectrum characterized by a distinct, more complex shape of the D'-band (see trace (b') in Fig. 3).

The Raman scattering spectra of our FLG-C-SiC samples can be compared to the characteristic spectrum of the graphene monolayer (trace (e) in Fig. 3) and/or to the spectrum of HOPG disks shown in Fig. 2. Within experimental uncertainty, the energetic position of the G-peak is common for all samples. Not surprisingly, this means that $E_{2g}$-in-plane optical phonons are the same in all these materials. It is worth noticing that the spectra presented here show no sign of the D-band. This demonstrates the weak disorder in our samples. The D-band can also appear in the spectra measured at the edges of the graphitic flakes, but in our experiments the laser spot was always focused on the interior of the measured structures.

The resemblance of the shape of the D'-band in the Raman scattering signal arising from the majority of the surface of our FLG-C-SiC samples to the D'-band measured on the exfoliated graphene flake is remarkable. A single Lorentzian peak is seen for both type of structures.
This is particularly clear for the graphene and the FLG-C-SiC samples 1 and 2, each showing the same half-width of 29 cm$^{-1}$. The fit of the Lorentzian profile to the D'-line is not as good for the case of sample 3 which has as many $\sim$90 layers, and results in a larger half-width ($D = 40$ cm$^{-1}$). The FLG-C-SiC samples differ however from graphene as far as the position of the D'-band is concerned. In our spectra for exfoliated graphene, this band is centered at 2641 cm$^{-1}$, whereas the D'-peak of FLG-C-SiC appears at slightly higher energies. Its actual position fluctuates from one location to another of the laser spot on the sample and from sample to sample, but remains in the range 2655 - 2665 cm$^{-1}$. The Raman scattering signal which arises from the visible inclusions on FLG-C-SiC surfaces shows a D'-band of a more complex form. We note the resemblance to the D'-band characteristic of graphite in general, and more precisely to the one observed for our thin HOPG disks. We speculate that this minority Raman scattering signal (linked to the inclusions) originates from some graphitic residues (Bernal-stacked graphene multilayers) on the surface of our FLG-C-SiC. Decomposing the D'-band characteristic of the graphitic residue in samples 1 into two Lorentzian peaks we find that the amplitude ratio of these peaks is the same as in the case of a HOPG disk with a height of $\sim$3 nm ($\sim$10 monolayers).

The striking correspondence in the shape of the D'-band observed in the majority Raman scattering signal from the FLG-C-SiC samples and from graphene is a clear indication of the similar electronic structures in both these material systems. This strongly supports previous work pointing towards the quasi-two dimensional Dirac-like character of electronic states in FLG-C-SiC samples. Our results are also consistent with recent reports which illustrate that two dimensional carbon layers on C-face SiC substrates are not Bernal stacked. Instead, these layers are generally rotated at specific angles, and in consequence are electronically well decoupled one with respect to another. To some extent, the FLG-C-SiC resembles the so-called “turbostratic” graphite, also composed of non Bernal-stacked, but randomly rotated, layers. A single component D'-band is also characteristic of the “turbostratic” graphite. The differences in the Raman scattering spectra of FLG-C-SiC and of “turbostratic” graphite are however also clear. “Turbostratic” graphite show much wider D'-peaks (typically 50 cm$^{-1}$ in Ref. 18) and a visible D-band, the latter due to the appreciable disorder and/or small size of such graphitic granulates. On the other hand, the D'-peaks in FLG-C-SiC and “turbostratic” graphite are centered at very similar energies, but characteristically higher, by about 20 cm$^{-1}$ as compared to the case of graphene. While the shift of the D'-peak for the FLG-C-SiC (and “turbostratic” graphite) as compared to the case of graphene might be due to a difference in the phonon energies characteristic of each of these systems, we speculate it is more likely due to the possible difference in the Fermi-velocity of Dirac cones in these systems. Recent reports clearly show that the energy position of the D'-band, arising from the double resonant Raman scattering process, can depend on the Fermi-velocity and this parameter seems to be $\sim 10\%$ larger for graphene compared to FLG-C-SiC. While no shift is observed for the G-band, a third possible but unlikely explanation for the observed energy shift of the D' band could also be a natural doping of the FLG-C-SiC.

In conclusion, we have found the majority signal of micro-Raman scattering observed from graphitized carbon-terminated surfaces of SiC shows a double resonant D'-band in the form of a single component peak which indicates the single-band electronic structure of this material. Together with previously reported results of transport spectroscopic and structural investigations, the present data confirms the appearance of Dirac-like electronic states in FLG-C-SiC. Remarkably, this is in spite of its multilayer character. Almost certainly, the layers of FLG-C-SiC are electronically well decoupled, as is the case to some extent in “turbostratic” graphite. This suggests that functional graphene-based devices can be developed using the methods of thermal decomposition (epitaxy) of silicon carbide, opening the way for many potential applications.

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