Growth and properties of few-layer graphene prepared by chemical vapor deposition

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ABSTRACT

The structure and the electrical, mechanical and optical properties of few-layer graphene (FLG) synthesized by chemical vapor deposition (CVD) on a Ni-coated substrate were studied. Atomic resolution transmission electron microscope (TEM) images show highly crystalline single-layer parts of the sample changing to multi-layer domains where crystal boundaries are connected by chemical bonds. This suggests two different growth mechanisms. CVD and carbon segregation participate in the growth process and are responsible for the different structural formations found. Measurements of the electrical and mechanical properties on the centimeter scale provide evidence of a large scale structural continuity: (1) in the temperature dependence of the electrical conductivity, a non-zero value near 0 K indicates the metallic character of electronic transport; (2) Young’s modulus of a pristine polycarbonate film (1.37 GPa) improves significantly when covered with FLG (1.85 GPa). The latter indicates an extraordinary Young modulus value of the FLG-coating of TPa orders of magnitude. Raman and optical spectroscopy support the previous conclusions. The sample can be used as a flexible and transparent electrode and is suitable for use as special membranes to detect and study individual molecules in high-resolution TEM.

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1. Introduction

The preparation of two-dimensional single-layer atomic crystals [1,2] and the study of their peculiar properties have become a hot issue now that single-layer graphene can be obtained by micro-mechanical cleavage methods [3], showing fascinating physical properties, such as thermodynamic stability, extremely high charge-carrier mobility and mechanical stiffness [4]. Furthermore, graphene has already been showing promise for applications such as transistors [5], transparent electrodes [6–8], liquid crystal devices [9], ultra-capacitors [10], and ultra-tough paper [11]. Therefore, the fabrication of graphene sheets of both large area and high-quality has become an important challenge. There are several ways to produce single or few-layer graphene, such as mechanical cleavage [3], epitaxy growth in ultra-high vacuum [12], atmospheric pressure graphitization of SiC [13], chemical oxidation of graphite [14,15], and chemical vapor deposition (CVD) using transition metals as catalysts [8,16–21]. Among them, the latter (CVD growth) is considered as one of the most promising, thanks to its flexibility; it can produce not only large-area graphene sheets, but also various chemically tuned two-dimensional (2D) materials, such as substitutionally doped graphene [22], 13C-graphene [23], and hexagonal boron nitride films [24], simply by changing gas sources. Growing graphite by using CVD at ambient pressure has been tried for a long time, but the fact that single-layer graphene could be synthesized this way was proven only very recently by Reina et al.
Few-layer graphene (FLG) was synthesized by the CVD method, as previously reported [8,17], with some modifications. The schematic diagram of the CVD set-up is provided in supplementary information. A 300 nm thick Ni film was coated onto a SiO2/Si substrate using an electron-beam evaporator. The Ni-coated substrate was positioned at the center of a quartz tube and heated to 1000 °C at a 40 °C/min heating rate, under a flow of argon and hydrogen (Ar/H2 = 1, 1000 sccm). The substrate was kept at 1000 °C in the Ar/H2 flow for 20 min to anneal the Ni film. The CVD growth of FLG was conducted at 960–970 °C, under a continuous flow of carrier gases, Ar and H2, mixed with less than 5 vol.% methane (CH4:Ar:H2 = 250:1000:4000 sccm). The reaction time was considered as the time from the moment when methane was introduced to the gas mass flow controller to the moment when the methane input-valve was closed. Due to the relatively fast flow, methane, “transported” by carrier gases, reached the middle of the quartz tube with the Ni-coated substrate within a few seconds. In this study, the reaction time was varied from 30 s to 7 min.

After the CVD reaction, the sample in the quartz tube was cooled down to 400 °C at the rate 8.5 °C/min, under a flow of argon and hydrogen. Then the gas valve for hydrogen was cooled down to 400 °C at the rate 8.5 °C/min, under a flow of argon and hydrogen. Then the gas valve for hydrogen was closed and the sample was cooled to room temperature under argon atmosphere.

Poly(bisphenol A carbonate) (PC) was dissolved in chloroform (solid content: ~15 wt.%). The PC/chloroform solution was spin-coated onto the FLG/Ni/SiO2/Si substrate with 500 rpm for 2 min. The coating was homogeneous with ~10 μm thickness. The FLG/PC film was released from the substrate by chemical etching of the Ni-layer with a FeCl3 (1 M) solution, followed, in some cases, by a concentrated HCl solution. The etching time varied depending on the substrate’s size from 30 min to 12 h. The FLG/PC film was rinsed several times with DI-water and dried with nitrogen gas.

The FLG was transferred onto arbitrary substrates such as a silicon wafer, a quartz slide, a KBr pellet, and a TEM grid, by selective dissolution of the PC layer from the FLG/PC film, using chloroform. The substrate was chosen depending on the purpose of the characterization. Raman spectra of samples were obtained with a Raman microscope equipped with a Jobin–Yvon LabRam spectrometer with a spectral resolution of 4 cm⁻¹. The laser excitation wavelength was 633 nm. All measurements were performed using a 100× objective lens and a D0.3 filter. Optical absorption of the FLG was studied using a UV/Vis/NIR Perkin–Elmer Lambda spectrometer. Scanning electron microscopy (SEM) analysis was carried out with a high-resolution FEI XL30 SFEG analytical SEM operated at 0.8 kV or 5 kV. Transmission electron microscopy (TEM) was performed using an imaging-side spherical aberration corrected Titan 80-300 (FEI, Netherlands), operated at 80 kV. Annular dark field scanning transmission electron microscopy (ADF-STEM) was done to determine the number of layers of the FLG on a Quantifoil grid having a 3.5 μm hole. Atomically resolved high-resolution (HR-TEM) images were taken to characterize the atomic structure of the FLG. A value of Cs = 15 μm with an underfocus of ca. 10 nm was used. Atoms appear black at these conditions. Optical microscopy was used to evaluate the layer distribution of an FLG sheet with an area of 4 × 4 mm² on a silicon wafer having a 300 nm thick thermal oxide layer at the top. The FLG/SiO2/Si samples were also used to study the temperature dependence of electrical conductivity by the 4-probe measurement method. For this purpose, four gold contacts were evaporated on top of the graphene layer. The outer electrodes were connected to a Keithley 2400 source-meter providing a constant current of 1 μA, and the voltage drop between the inner electrodes was recorded with a Keithley 2000 voltmeter. The channel length for the voltage drop was 400 μm. The sample was placed into the sample chamber and annealed at 120 °C in vacuum (10⁻⁷ mbar) for 24 h before the measurement. The sheet conductance of the sample was measured as a function of temperature from 300 K to 4 K.

Besides the T-dependence measured on the silicon substrate, the sheet resistance of a large-area (2 × 2 cm²) transparent FLG/PC film was measured also in a four-probe configuration, using the Keithley 225 current source and sensitive Keithley 181 digital nanovoltmeters for voltage reading. Young’s modulus of the FLG was determined from the elastic part of the stress vs. strain curve, measured by a force transducer Hottinger Baldwin Messtechnik, type 52.
450 °C, the molecule dissociates and becomes reactive. Dissociation of the H₂ molecules in the presence of Ni and other metals was clearly demonstrated in an experiment done by Haluška et al. [26]. Atomic hydrogen then acts as an etching agent reducing preferably amorphous carbon that contains unsaturated dangling bounds. For CVD graphene growth we found that a critical amount of hydrogen is also necessary to synthesize FLG because hydrogen keeps a balance between the production of reactive hydrocarbonaceous radicals and the etching of the graphite layer during the CVD process. If the ratio of methane to hydrogen is too low, the etching reaction becomes much faster than the formation of graphene layers. This was also experimentally proved in a recent work of Kong’s group [20]. The reaction time was varied between 30 s and 7 min. The SEM image in Fig. 1a shows that multilayer graphene is formed already at a very short reaction time (30 s). The SEM image contrast is given by the amount of secondary electrons generated in the first few atomic layers of the sample. In the standard calculation (for 20 kV) [27], a higher SEM yield is expected for Ni than for C. Thus, the black parts can be identified as the thickest graphitic layers (Ni is covered by the graphite) and white as the thinnest graphene layers (most secondary electrons from Ni can escape). An empty Ni surface can be ruled out, as even in the thinnest areas, folds and crinkles of the graphene layers can be discerned.

As the reaction time increases, a more homogenous coverage is obtained, however, a different number of graphitic layers can still be distinguished (Fig. 1b). Raman spectra of the FLG attached to the nickel substrate after synthesis reveal that the number of graphene layers varies from a single-layer to several layers along the graphene sheet (Fig. 1c). In particular, a significant difference can be seen in the full width at half the maximum (FWHM) of the D*-mode (2600–2800 cm⁻¹), which varies between 35 cm⁻¹ and 90 cm⁻¹. Nevertheless, the line-shape of the D*-mode does not change much, unlike in Raman spectra of samples prepared by the micro-mechanical cleavage method [28]. The very narrow D*-peak (FWHM ~35 cm⁻¹) in the lowest spectrum in Fig. 1c is well fitted by a single Lorentzian-line-shape and is related to single-layer graphene. The second spectrum from below in the same figure shows the D*-mode still relatively narrow but asymmetrical that can be fitted by two Lorentzians. This D*-line-shape is a fingerprint of an AB-stacked bi-layer. The D*-line-shapes in the other three spectra plotted higher in Fig. 1c are wide (FWHM up to 90 cm⁻¹) and symmetrical and they reflect turbostratic stacking of several graphitic layers. This suggests that only a single-layer and, in some cases, a bi-layer can be distinguished, when analyzing the line-shape of the D*-mode in our FLG sample.

In order to study their physical properties, the FLG samples were then transferred onto various arbitrary substrates such as silicon wafers, quartz glasses and KBr pellets, depending on the characterization method. A FLG was transferred first onto a PC film (following the procedure described above) and then, either the FLG/PC film itself was characterized, or it served as a medium for the next transferring process. The presence of the FLG on PC film was confirmed by Raman spectra as well (SI-2). The transfer onto the PC film gives the possibility to study the morphology of the back side of the FLG sheet, originally attached to the Ni-layer. The SEM image in Fig. 2a clearly shows a large amount of amorphous carbon covering the bottom side of the FLG after the nickel layer was etched by a 1 M FeCl₃ solution. The presence of amorphous carbon at the bottom side of the FLG strongly supports a growth mechanism of graphite on nickel by CVD as suggested by Obraztsov et al. [16,29]. On the contrary, the next SEM image (Fig. 2b), taken after further treatment using concentrated HCl for 1 h, shows a clean surface of the FLG, demonstrating how efficiently amorphous carbon can be

![Fig. 1 – SEM image of a CVD-grown graphene sheet on a nickel-coated SiO₂/Si substrate (a) after 30 s and (b) after 7 min; (c) Raman spectra of a CVD-grown graphene sheet on a nickel-coated SiO₂/Si obtained from different positions of the sample (scale bar: 1 µm).](image-url)
removed by this procedure. The cleaning step developed and demonstrated in this work might be very important in applications for transparent electronic devices.

In order to study the structure of FLG, we transferred the FLG onto a Quantifoil grid having 3.5 \( \mu \text{m} \) holes, by dissolution of the PC layer from the FLG surface, using chloroform. No residue of PC was detected on the sample. Fig. 3 represents an ADF-STEM image of an FLG sample. Determined from electron diffraction, the single-layer and bi-layer were first identified and their ADF intensities were taken then as a reference for the ADF-STEM thickness analysis using the formula \( I \approx t \times Z^{1.7} \), where \( I \) is intensity, \( t \) is thickness and \( Z \) is the atomic number of scattering atoms. Surprisingly, the distribution of the number of graphene layers did not depend much on the duration of the CVD process, so that no significant difference in distribution of the number of layers between the samples synthesized for 30 s, 1 min, 2 min and 3 min was identified.

Each of the samples contained single-layer graphene covering about 2% of the area and few-layer graphene (less than 10 layers) on most of the area. The TEM image in Fig. 4a shows a single-layer region of FLG surrounded by a multi-layer region. In some places at the multi-layer boundaries, a crystal face of the graphene layer can be distinguished from the angles of the edges. The atomic resolution image of single-layer graphene (Fig. 4b) shows a perfect single crystal structure in the displayed region, where no defect can be seen. Fig. 4c presents an HR-TEM image of a bi-layer region where a clearly pronounced Moiré pattern indicates a turbostratic arrangement of the layers. Overall, we observe a mixture of AB-stacked and turbostratically arranged layers. The turbostratic arrangement might happen due to a non-equilibrium process during the FLG growth where two parallel mechanisms, a pure CVD process as well as carbon segregation during fast cooling (which can be referred to as quenching), take place [30]. Under these highly non-equilibrium conditions of fast FLG growth, the formation of energetically more favourable tightly stacked graphitic layers is suppressed and, instead, a less ordered turbostratic structure is formed.

Interestingly, in few-layer regions, we found polycrystalline structures with different orientations, but the different domains seemed to be connected by chemical bonds (Fig. 4d). This might be responsible for the huge mechanical strength reported below, even though the inhomogeneous structure of the FLG dominates over a large area. Analysis of the number of graphitic layers from electron diffraction patterns was discussed in detail in the literature [31,32].

The FLG was also transferred on a quartz slide and on a KBr pellet in order to characterize its optical absorbance behaviour in the IR–NIR–VIS range. In the case of graphene prepared by micro-mechanical cleavage, photon absorbance does not change when the wavelength varies from the visible to the infrared region. The value of the absorbance intensity is given by the fine structure constant or its multiples as the number of graphene layers increases [33,34]. However, our FLG shows a monotonic increase of absorbance as the photon energy increases, with a maximum at 4.6 eV (Fig. 5b). A recent study [35] pointed out that the peak should be attributed to resonance excitons that are formed in two-dimensional semimetals.

We measured the sheet resistance of FLG sheets as a function of the temperature \( R(T) \). All samples prepared at 3 min of
reaction time show almost the same “semiconducting” behaviour: $R(T)$ decreases as the temperature increases (Fig. 5c). However, the total change in resistance from 4 K up to room temperature is only a factor of two [36]. For a semiconductor, the energy gap in the electron energy spectrum would cause electrical conductivity to drop to zero as thermal activation is becoming inefficient. In our case the trend of conductance at low temperatures contradicts the semiconductor behaviour: Sheet conductance plotted in the log–log scale (Fig. 5c: inset) clearly shows convergence of conductance to a non-zero value as $T \to 0\,\text{K}$.

The mechanical properties of the FLG were investigated comparing Young's modulus of a simple PC film (1.37 GPa) with Young's modulus of a double-layer composite FLG/PC film (1.85 GPa). The values were determined from the elastic part of the stress vs. strain curve (SI-4a). We calculated Young's modulus of the FLG by using a formula for Young's modulus of composed layers [37], considering that our FLG sample consists of five graphene layers (ADF-STEM estimate). Then the value of Young's modulus of the FLG is 4.6 TPa. Based on the HR-TEM investigation, even though the layers of FLG contain many polycrystalline domains, the high value of Young's modulus can be explained by the fact that the different crystal domains merge, thus forming mainly chemical bonds between each other (Fig. 4d). Fig. 5d shows the electrical sheet resistance, measured also as a function of strain. In the elastic part below the 2% prolongation, there is no observable change in sheet resistance. At higher strain, sheet resistance increases so that it triples its original value when strain reaches 17%.

Finally, even though there is still a way to go to optimize CVD growth in order to obtain a more homogeneous structure and to eliminate the multi-layer “scales” decorating the surfaces of FLG sheets, we can already use the FLG in two applications. Polycarbonate, which is well-known for its high optical transparency and its good mechanical performance, was chosen as a substrate to produce flexible transparent electrodes based on a FLG/PC film. The sheet resistance vs. the transmittance of the films was shown in SI-5. A sample with 84% transmittance measured at a wavelength of 550 nm has $800\,\Omega/\text{sq}$ of sheet resistance. This value is not as high as that of a doped single-walled carbon nanotubes transparent network produced by the arc-discharge method [38]. However, we believe there is still space for improving the quality of CVD-grown FLG films. The main obstacle limiting the quality of our flexible transparent electrode is the appearance of thick multi-layer “scales” which absorb the visible light (Fig. 5b), but do not contribute to the electrical conductivity, since they form small, non-connected islands. The second application we have realized is a special membrane for TEM study. The unique potential of using graphene as an ideal support film for electron microscopy has been demonstrated
recently [31]. However, the commonly used method for preparing a graphene membrane, namely through micro-mechanical cleavage, requires the use of a complex process that yields one micron-sized graphene at a time. On the contrary, CVD-grown FLG can be transferred by a single operation on a large number of TEM grids. As we have shown in Fig. 4b, there are large enough areas of single-layer graphene that can be used for observation, potentially from nano-scale small particles down to single molecules. We show an example in the supporting material (SI-6).

To conclude, we have synthesized an FLG sheet with an area of several square centimeters by the CVD process, and demonstrated its novel properties. It consists of single- and double-layers with randomly distributed isolated multi-layer “scales” which usually are formed near the grain boundaries of polycrystalline nickel during the growth process. Based on HR-TEM we did show that different crystal domains of the FLG are connected by chemical bonds. This, apparently, leads to extraordinary mechanical properties and reasonably small sheet resistance measured at large area (2 cm$^2$) of our sample despite the macroscopically inhomogeneous structure. The FLG can be used in applications such as a flexible transparent electrode and a special membrane for TEM study.

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Appendix A. Supplementary data


REFERENCES