Graphene Films with Large Domain Size by a Two-Step Chemical Vapor Deposition Process

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ABSTRACT The fundamental properties of graphene are making it an attractive material for a wide variety of applications. Various techniques have been developed to produce graphene and recently we discovered the synthesis of large area graphene by chemical vapor deposition (CVD) of methane on Cu foils. We also showed that graphene growth on Cu is a surface-mediated process and the films were polycrystalline with domains having an area of tens of square micrometers. In this paper, we report on the effect of growth parameters such as temperature, and methane flow rate and partial pressure on the growth rate, domain size, and surface coverage of graphene as determined by Raman spectroscopy, and transmission and scanning electron microscopy. On the basis of the results, we developed a two-step CVD process to synthesize graphene films with domains having an area of hundreds of square micrometers. Scanning electron microscopy and Raman spectroscopy clearly show an increase in domain size by changing the growth parameters. Transmission electron microscopy further shows that the domains are crystallographically rotated with respect to each other with a range of angles from about 15° to nearly 30°. Electrical transport measurements performed on back-gated FETs show that overall films with larger domains tend to have higher carrier mobility up to about 16 000 cm² V⁻¹ s⁻¹ at room temperature.

KEYWORDS Graphene, domains, mobility, chemical vapor deposition, nucleation, Raman spectroscopy

The fundamental properties of graphene are making it a very attractive material for use in electronics, optoelectronics, nanoelectromechanical systems, chemical and biosensing, and many other applications. Compared to other graphene synthesis processes reported to date, such as mechanical exfoliation of graphite, reduction of graphene oxide, or epitaxial growth on SiC substrates, graphene growth on metal substrates has the distinct advantage of being able to provide very large-area graphene films transferrable to other substrates. This advantage is especially true for the case of graphene growth on Cu substrates by chemical vapor deposition (CVD) of methane as reported by our group and reproduced by other groups. CVD of graphene on Cu yields a uniform graphene film whose size is limited only by the size of the Cu substrate and the growth system.

The presence of graphene on metal surfaces was observed as early as 1969 when the nomenclature “monolayer graphite” (MG) was first introduced by John May in his rationalization of low energy electron diffraction (LEED) patterns that were, theretofores, unassigned. Later, Blakely and his research undertook extensive studies of mono- and bilayer graphene on Ni substrates, and several reviews (among others) of studies of growth of graphene on a wide variety of metal substrates are available. It has been shown that graphene can grow across metal steps and grain boundaries, and graphene domains of a few hundred square micrometers have been observed on Ru substrates. We have observed graphene domains of tens of square micrometers on Cu substrates using a C isotope labeling technique. The C isotope labeling technique further demonstrated that graphene growth on Cu is surface-mediated, that is, C species decomposed from methane nucleate on the Cu surface and the nuclei grow to form islands and then domains (we note that some researchers may refer to domains as “grains”) that cover the metal surface in its entirety. In our previous work, we also observed that the domains might be in part defective as indicated by the appearance of D bands in the Raman spectra at certain locations. The interdomain D-band defects are believed to arise from the misalignment of the domains as they come together to fully cover the Cu surface. Because of the presence of such interdomain defects we decided to try to develop processes to increase the domain size as an approach to decrease the density of such defects that are identifiable by the presence of the Raman D band.

We studied the effect of growth parameters such as temperature (T), methane flow rate (JMe), and methane partial pressure (PMe) on the domain size of graphene grown...
on polycrystalline Cu. It should be noted that here we did not consider the effect of Cu grain orientation. Although graphene nuclei/islands on different Cu grains may have different shapes and densities, as reported previously,\(^\text{12}\) at this time we believe that the shape of such islands may be controlled mainly by the growth conditions rather than the crystallographic orientation of the Cu substrate. We thus performed our investigation simply by comparing graphene islands with similar shape without concern about the specific Cu grain crystallographic orientation. In the case of the present parameter space, most of the graphene islands have a starlike shape and uniform density across the Cu foil, as shown by the scanning electron microscopy (SEM) images in Figure 1 of partially grown graphene under different \(T\), \(J_{\text{Me}}\), and \(P_{\text{Me}}\). To control the graphene domain density, we performed experiments where we used the precursor flux and temperature to change the degree of supersaturation of active C-containing species (most probably CH\(_x\)) on the surface of the Cu to promote graphene nucleation. What we observed is that the density of graphene nuclei can be decreased (domain size increases) as \(T\) is increased (Figures 1a,b), or as \(J_{\text{Me}}\) and \(P_{\text{Me}}\) are decreased (Figure 1b,c) and (Figure 1c,d), respectively. That is, high \(T\) and low \(J_{\text{Me}}\) and \(P_{\text{Me}}\) were found to yield a low density of graphene nuclei and thus large domain size.

However, when \(J_{\text{Me}}\) and \(P_{\text{Me}}\) are less than a critical value, graphene does not nucleate and above these critical values graphene nuclei can form but, in a given range of values, graphene growth terminates before full surface coverage as shown in Figure 1d. In this case, the growth conditions under which partial coverage occurred were growth temperature \(T = 1035\, ^\circ\text{C}\), methane flow rate \(J_{\text{Me}} = 7\, \text{scm}\), and methane partial pressure \(P_{\text{Me}} = 160\, \text{mTorr}\). Under these conditions even if the surface is continuously exposed to methane, full coverage is still not achieved. The precise growth conditions for partial coverage may vary from growth system to growth.

**FIGURE 1.** SEM images of partially grown graphene under different growth conditions: \(T\) (°C)/\(J_{\text{Me}}\) (scm)/\(P_{\text{Me}}\) (mTorr): (a) 985/35/460, (b) 1035/35/460, (c) 1035/7/460, (d) 1035/7/160. Scale bars are 10 \(\mu\)m.
system. To fully cover the Cu surface the partial pressure of CH₄ must be increased; for example for J_{Me} > 35 sccm and/or P_{Me} > 500 mTorr in our growth chamber, complete surface coverage is obtained within 2 to 3 min.

The effects of J_{Me} and P_{Me} on graphene growth kinetics were further investigated by using the C isotope labeling technique together with ex situ micro-Raman spectroscopy. In these experiments the Cu surface was exposed to ^{13}CH₄ and ^{12}CH₄ (normal methane) sequentially. Since graphene growth on Cu occurs by surface adsorption, the isotope distribution in the local graphene regions will reflect the dosing sequence employed and can be mapped according to their different Raman mode frequencies. For example, the G band of ^{13}C graphene is located at ∼1520 cm⁻¹ while that of ^{12}C graphene is at ∼1580 cm⁻¹ (Figure 5g). By integrating the intensity of the G band of ^{13}C graphene over 1490−1550 cm⁻¹, the ^{13}C graphene regions show as bright regions in the G^{13}-band maps while the ^{12}C graphene regions show up as dark regions (Figure 2b,f), and vice versa for the G^{12}-band maps integrating the intensity of the G band of ^{13}C graphene between 1550−1610 cm⁻¹ (Figure 2c,g). The regions that show up as dark in both maps are gaps uncovered by graphene, which can also be easily distinguished in the optical micrograph (e.g., Figure 2a). As shown in Figure 2, although the Cu surface was exposed to the alternating C isotopes a total of eight times, for the case of P_{Me} = 160 mTorr graphene growth terminated after the sixth dose with a maximum coverage of ∼90%; whereas when P_{Me} was increased to 285 mTorr, graphene growth terminated after the fourth dose and the surface was fully covered. The graphene growth rate was calculated by measuring the area of the isotopically labeled regions. Here we define the coverage rate, v_{coverage}, as the increase of graphene coverage (graphene area, A_{graphene}, divided by the total Cu area, A_{Cu}) within unit time (t)

\[ v_{coverage} = \frac{dA_{graphene}}{A_{Cu} dt} \tag{1} \]

The average area growth rate of graphene domains, v_{domain}, is related to v_{coverage} as

\[ v_{domain} = \frac{v_{coverage}}{n} \tag{2} \]

where n is the domain (i.e., nucleus) density.

Figure 3 shows a summary of the growth process of graphene on Cu at two different pressures, 285 mTorr and 160 mTorr. Figure 3a shows that as the Cu surface is exposed to methane at a P = 285 mTorr, the surface is fully covered with graphene after about 1.5 min whereas at a gas pressure of 160 mTorr the Cu surface reaches only 90% coverage after about 3 min and never reaches full coverage even after continued exposure to methane. Figure 3b further shows that as the surface coverage increases or as the exposed Cu surface area decreases the graphene growth rate
even though C graphene does not nucleate and no graphene is observed supersaturated. In the case of an undersaturated Cu surface, be considered: (1) undersaturated, (2) saturated, and (3) P of C an undersaturated surface, graphene nuclei do not form). decreases dramatically. We attribute the decrease in growth rate to the fact that the C species are supplied by the available Cu-catalyzed decomposition of methane.

In summary, graphene growth on Cu likely proceeds as follows:

1. Exposure of Cu to methane and hydrogen.
2. Catalytic decomposition of methane on Cu to form C_2H_4.
3. Depending upon the temperature, methane pressure, methane flow, and hydrogen partial pressure, the Cu surface is either undersaturated, saturated, or supersaturated with C_2H_4 species.
4. Formation of nuclei as a result of local supersaturation of C_2H_4.
5. Nuclei grow to form graphene islands (in the case of an undersaturated surface, graphene nuclei do not form).
6. Full Cu surface coverage by graphene under certain T, P and J.

Therefore, there are three Cu surface conditions that can be considered: (1) undersaturated, (2) saturated, and (3) supersaturated. In the case of an undersaturated Cu surface, graphene does not nucleate and graphene is observed even though C_2H_4 may be present in the vapor phase and on the Cu surface. In the case of the saturated surface graphene, nuclei form, graphene grows to a certain island size, and then it stops because the amount of C_2H_4 available from the exposed Cu surface is insufficient to continue driving the C attachment to the island edges and the Cu surface is only partially covered with graphene islands. That is, graphene islands, the Cu surface, and the vapor phase are in equilibrium. In the case where the Cu surface is supersaturated on the other hand, there is always enough methane to form sufficient C_2H_4 to drive the reaction between the C_2H_4 at the surface and the edges of graphene islands. In this last case, graphene islands grow until neighboring islands connect to each other to fully cover the Cu surface.

On the basis of the above observations and the presence of defects at the interdomain regions, as determined by the presence of a higher concentration of Raman D-band, we developed a two-step synthesis technique with the process flow shown in Figure 4. For a given temperature (usually high temperature) nuclei are formed in step I at low J_Me and P_Me followed by step II, where the partial pressure of the methane is increased to promote full surface coverage. The resulting domains of the continuous graphene films grown by this technique were also delineated by C isotope labeling where 13CH_4 was used in step I followed by 12CH_4 (normal methane) in step II, as shown in Figure 5. By comparing the G'-band Raman maps, Figure 5b,e, it is clear that at low J_Me and P_Me, the number of graphene nuclei is much smaller than that for high J_Me and P_Me. As a consequence of the lower density of graphene nuclei, graphene can grow into larger domains. It is important to note that for a given temperature and partial pressure condition, once the nuclei density is set no significant new graphene nuclei are formed and subsequent changes in growth conditions only affect the graphene growth rate (we do not exclude that significant change in growth conditions may promote new nucleation). The Raman maps of D bands (Figure 5c,f), a measure of defects in graphene, which are located at ~1310 and ~1350 cm^{-1} for 13C graphene and 12C graphene, respectively, show that graphene films with larger domains have a lower density of defects arising from nucleation centers and interdomain regions.

To further understand the domain structure and the impact of domains on electrical properties, we performed transmission electron microscopy (TEM) and electrical transport measurements using field effect transistor (FET). Figure 6a,b shows a high-resolution TEM image and a mask-filtered image of a monolayer graphene film at a domain boundary. As can be seen from the fast Fourier transform (FFT) images in Figure 6c,e, the FFTs from the white and black box (Figure 6c,d, respectively) in Figure 6a show a single set of a hexagonal spot pattern rotated relative to one another in each image. The whole FFT of Figure 6a shown in Figure 6e shows the combination of the two FFT patterns rotated relative to one another by approximately 18.5°. The FFTs in Figure 6c–e show clear evidence of the presence of domain boundaries in this image. Magnified images in Figure 6f,g of the graphene in the different domains show how the hexagonal graphene structure is oriented within the imaged graphene domain. Measurements on tens of domains show that the domain misorientation ranges from 13...
to about 30° (since graphene has a hexagonal structure, anything over 30° is simply 60−X). On the basis of the TEM results, together with the fact that the domain size of the films studied is much smaller than the Cu grain size (hundreds of micrometers to millimeters) and that graphene has a hexagonal structure and Cu is cubic, it is likely that the graphene is not growing epitaxially on Cu except perhaps on (111) grains; we plan to further address this issue in the future.

We measured the transport properties of CVD graphene having two different average domain sizes, (linear length) of 6 and 20 µm, and compared them with those of graphene exfoliated from natural graphite. The effective mobility ($\mu$) was extracted using the mobility model introduced by Kim et al.\textsuperscript{24} where the $R$–$V_{bg}$ curves were fit to give a constant value of mobility and intrinsic carrier concentration for each of the measured devices. Figure 7 shows a summary of the transport properties of CVD graphene in comparison to exfoliated graphene. Figure 7a shows the device layout of a back-gated FET where the light regions are Ni contacts and the dark region is the graphene. Figure 7c shows a summary of the mobility of many devices for the three different types of graphene: graphene with 6 and 20 µm domains, and graphene exfoliated from natural graphite. The mobility for many devices from films with small domains, 6 µm, is in the range of ranges from 800 to about 7000 cm$^2$ V$^{-1}$ s$^{-1}$. In contrast, devices with larger graphene domains, 20 µm, have a much higher range of mobility, 800 to 16000 cm$^2$ V$^{-1}$ s$^{-1}$ compared to 2500 to over 40 000 cm$^2$ V$^{-1}$ s$^{-1}$ for exfoliated graphene. Obviously, even films with large domains have devices with low mobility. The lowest mobility for both 6 and 20 µm domain films may be associated not only with the presence of domain boundaries but also with the presence of wrinkles and other defects some of which are induced by the transfer process. The observation of high mobility in CVD graphene that is in the same range as that observed in exfoliated graphene suggests that we are improving the material quality based on our fundamental understanding of the graphene microstructure as well as its development during growth. There is still much work to be done before we have a complete understanding of the effect of structural defects and transport properties but we are beginning to develop the basic process variables that will help create higher quality graphene.

In summary, we investigated the effects of growth parameters such as temperature, methane flow rate, and partial pressure on CVD synthesis of graphene on Cu substrates. High-temperature and low methane flow rate and partial pressure are preferred to generate a low density of graphene nuclei, while high methane flow rate or partial pressure are preferred for continuous large-area graphene films. On the basis of these observations, we developed an isothermal two-step growth process in which a low graphene nuclei density is set followed by achieving full graphene surface coverage by increasing the methane flow rate and partial pressure. Electrical transport data showed that graphene films with large domains have higher mobility than those with small domains predominantly due to a decrease in interdomain defects. Further TEM results indicate that the domains are rotated with respect to each other by as much as 30°. The basic understanding presented in this paper can lead to significant improvements in graphene synthesis and graphene-based electronic devices.

**Experimental Section. Graphene Synthesis.** A split tube furnace with a 6 in. heating zone and a 1 in. outer diameter

| Table 1: Two-Step Graphene Growth Parameters |
|---|---|---|---|---|
| experiments | $T$ (°C) | $J_{Me}$ (sccm) | $P_{Me}$ (Torr) | $t$ (min) | average domain area ($\mu$m$^2$) |
| A Step I | 1035 | 7 | 0.160 | 2.5 | 142 |
| Step II | 2 | 1 |
| B Step I | 1035 | 35 | 0.460 | 0.5 | 33 |
| Step II | 2 | 1 |

FIGURE 5. Optical images and micro-Raman imaging spectroscopy maps of graphene films transferred onto 285 nm SiO$_2$/Si substrates. Optical images (a,d), Raman G$_{12}$-band maps (b,e) and D$_{12}$-band maps (c,f) of sample A (a–c) and B (d–f), respectively. (g) Raman spectra of $^{13}$C graphene, $^{12}$C graphene, $^{13}$C/$^{12}$C junction, wrinkle, nucleation center, and domain boundary, respectively, as marked with corresponding colored circles in (a–c). A summary of the two-step process for conditions A and B is shown in Table 1. Scale bars are 5 µm.
A quartz tube with gas panel having methane and hydrogen was used in a CVD mode to grow graphene films on Cu. The graphene synthesis process presented here is similar to that previously reported with small changes: the 25 µm thick Cu foil was first reduced and annealed at 1035 °C under 2 sccm of H2 at a pressure of 40 mTorr for 20 min to increase the Cu grain size and clean the Cu surface; the graphene growth temperature was changed to the desired value after the initial Cu cleaning and grain growth and a desired amount of methane was introduced into the growth tube; the methane partial pressure was controlled by either methane flow rate or a ball valve between the quartz tube outlet and the pump; the growth time was varied accordingly; the methane and hydrogen gas flow and pressure were kept constant, same values as the growth process, during the furnace cool-down.

Micro-Raman Characterization. Graphene films were transferred onto 285 nm SiO2/Si substrates using poly-(methyl methacrylate) (PMMA) for optical microscopy and micro-Raman imaging spectroscopy (WiTec Alpha 300, 532 nm laser wavelength, 100× objective).

Mobility Measurements. Graphene films were transferred onto p-type Si wafers with a 300 nm thick SiO2 layer using the same transfer method as previously and FET devices were fabricated using standard photolithography and electron beam lithography processes; Ni was used for metal contacts. The channel lengths and widths ranged from 2 to 100 µm and 1 to 15 µm, respectively. The device
structures were designed to permit multiple electrical measurements. All of the electrical measurements were performed using back-gated devices at room temperature in air using a HP 4155 Semiconductor Parameter Analyzer.

Transmission Electron Microscopy. A Cs corrected FEI Titan S/TEM operating at 80 keV was used for TEM imaging. HRTEM image was averaged with HoloWorks 5.0 using 20 individual images using an acquisition time of 0.5 s per each frame for an improved signal/noise ratio. HRTEM was mask filtered using an array mask in Digital Micrograph. Magnifier function in Holoworks 5.0 was used to magnify cropped TEM images.

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FIGURE 7. (a) Optical micrograph of a FET device (top) and (b) a typical plot of the normalized channel resistance ($R_{ch}$) as a function of applied back gate voltage ($V_{bg}$). (c) Carrier mobility as a function of graphene domain size in comparison to exfoliated graphene.