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Cu oxidation kinetics through graphene and its effect on the electrical properties of graphene†

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The oxidation kinetics of Cu through graphene were evaluated from the surface coverage of Cu oxide (F_{ox}) by varying the oxidation time ($t_{\text{ox}} = 10\text{--}360$ min) and temperature ($T_{\text{ox}} = 180\text{--}240$ °C) under an air environment. F_{ox} , as a function of time, well followed the Johnson–Mehl–Avrami–Kolmogorov equation; thus, the activation energy of Cu oxidation was estimated as 1.5 eV. Transmission electron microscopy studies revealed that Cu_2O formed on the top of the graphene at grain boundaries (G-GBs), indicating that Cu_2O growth was governed by the out-diffusion of Cu through G-GBs. Further, the effect of Cu oxidation on graphene quality was investigated by measuring the electrical properties of graphene after transferring. The variation of the sheet resistance (R_s) as a function of t_{ox} at all T_{ox} was converted into one curve as a function of F_{ox} . R_s of $250 \Omega \text{ sq}^{-1}$ was constant, similar to that of as-grown graphene up to $F_{\text{ox}} = 15\%$, and then increased with F_{ox} . The Hall measurement revealed that the carrier concentration remained constant in the entire range of F_{ox} , and R_s was solely related to the decrease in the Hall mobility. The variation in Hall mobility was examined according to the graphene percolation probability model, simulating electrical conduction on G-GBs during Cu_2O evolution. This model well explains the constant Hall mobility within $F_{\text{ox}} = 15\%$ and drastic F_{ox} degradation of 15–50% by the concept that the electrical conduction of graphene is disconnected by Cu_2O formation along with the G-GBs. Therefore, we systematically developed the oxidation kinetics of Cu through graphene and simultaneously examined the changes in the electrical properties of graphene.

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Introduction

Since the experimental discovery of graphene by the scotch-tape method in 2004,¹ high-quality single layer graphene has been grown from various hydrocarbon precursors by chemical vapor deposition (CVD) on Cu film. As a metal catalyst for the CVD process,^{2,3} Cu foil not only dissociates hydrocarbon precursors but also self-limits the growth of graphene by its extremely low solubility for carbon.^{2,4} Nevertheless, under the random nucleation and growth process of CVD, the graphene layer has a polycrystalline nature on the Cu surface. The graphene grain boundaries (G-GBs) inherently inhibit charge transport, severely degrading the mobility of charge carriers.^{5–8} Moreover, the G-GBs significantly affect other physical properties including, the mechanical strength and gas-molecule impermeability of

graphene.^{9,10} To grow a graphene layer with large grain size, researchers have carefully tuned the graphene-growth conditions.^{11–13} Furthermore, the characterization of the G-GBs, such as grain size linked to the physical properties of graphene, has been a major topic of concern.

Raman spectroscopy is among the most efficient characterization methods of graphene. The D peak (which appears at $\sim 1350 \text{ cm}^{-1}$ on the Raman spectrum) represents the breakage of the sp^2 bond by structural defects in the graphene lattice.¹⁴ Raman spectroscopy provides various quantitative information of graphene, such as the number of graphene layers (from the intensity ratio of the G and D peaks) and doping amount (from the shift of the G or 2D peak).^{15–17} However, Raman spectroscopy cannot provide information over the entire sample area because the laser beam size is limited to a few μm .² Furthermore, the graphene must be transferred to an SiO_2 substrate for accurate analysis. Transmission electron microscopy (TEM) is clearly advantageous because its atomic resolution enables direct observation of the graphene lattice.¹⁸ However, sample preparation is quite difficult, and the observed area is extremely small. The measurement of the electrical properties of graphene requires the transfer of graphene on the insulating substrate for fabricating a device. Because various defects (*e.g.*, wrinkles, cracks, holes, and

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organic residues) are introduced during the transfer process,¹⁹ the measured property cannot be correlated with the quality of the as-grown graphene. Therefore, a method for the analysis of the as-grown graphene on Cu is required for proper tuning of the graphene-growth condition.

In this respect, the oxidation of Cu underlying graphene has been proposed by several researchers as a useful method to simply observe the graphene grain.^{20–26} Because the hexagonal arrangement of carbon atoms in graphene is sufficiently compact to inhibit the penetration of oxygen (and consequently the oxidation of Cu),²⁷ Cu is selectively oxidized by oxidizing species, such as OH[−] or O[−], which penetrate the G-GBs.^{22–24} These radicals are typically dissociated from H₂O at elevated temperatures (approximately 200 °C) or under UV exposure.^{23,24} Energy calculations using density functional theory and TEM

observations have shown that oxygen radicals penetrate the G-GBs without bond breaking. Oxygens penetrating the boundary at room temperature was reported to form Cu oxide at the graphene–Cu interface.²⁵ Moreover, the lateral diffusion of oxygen at the graphene–Cu interface is hindered by van der Waals forces between the Cu and graphene.²⁶ Although the results of oxidation kinetics depend on the states of the graphene (*e.g.*, grain size, shape, and stitching between grains) and the Cu foil (*e.g.*, crystallographic orientation),²⁸ the oxidation of Cu through the grain boundary of the graphene layer has not been systematically investigated. Furthermore, the extent to which Cu oxidation damages the electrical property of graphene is unknown.

Here, Cu oxidation through graphene was investigated as a function of temperature (180–240 °C) and time (10–360 min).

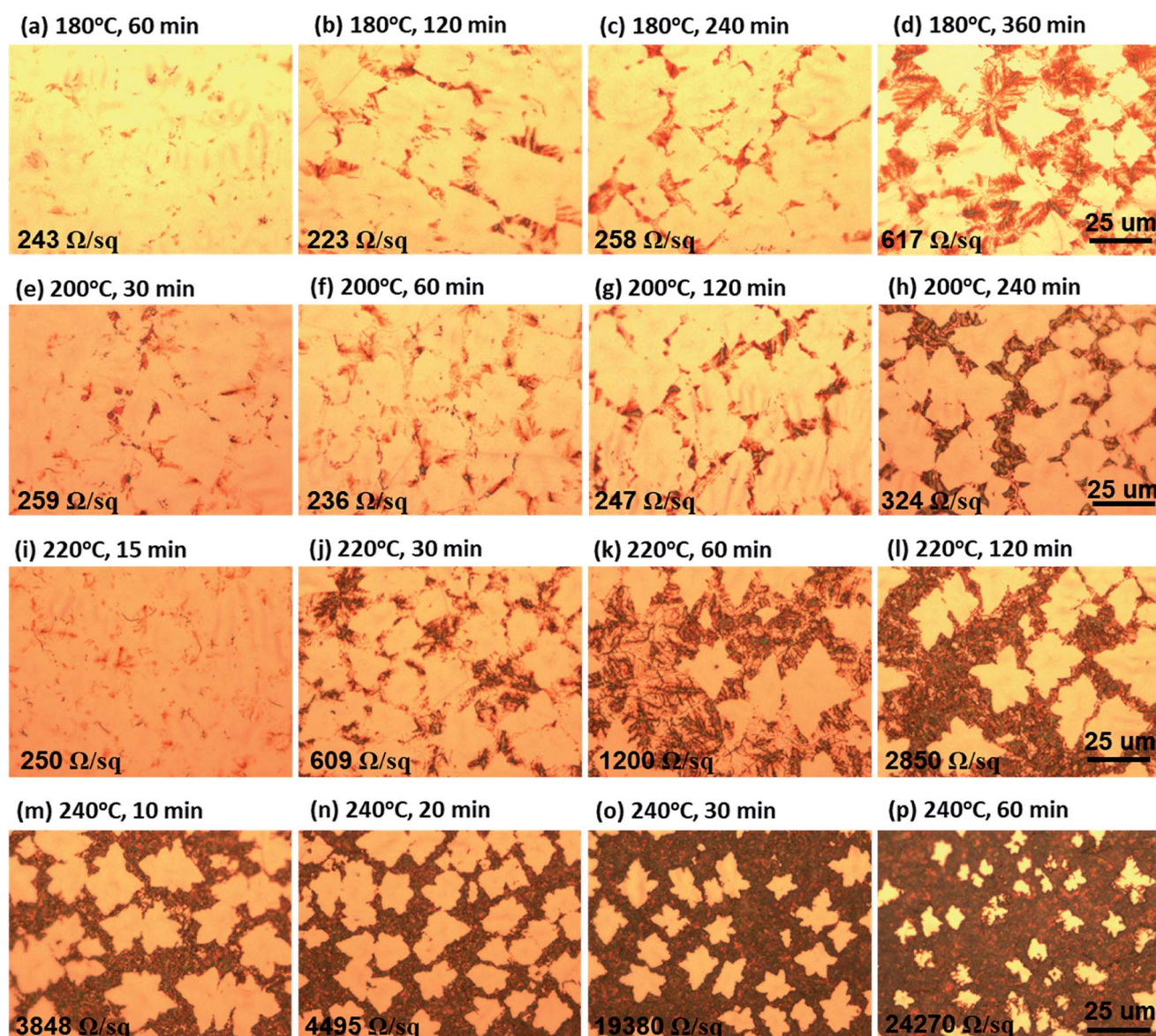


Fig. 1 Optical microscope images of the as-grown graphene after the oxidation at (a)–(d) 180 °C, (e)–(h) 200 °C, (i)–(l) 220 °C, and (m)–(p) 240 °C.



The oxidation kinetics of Cu through the G-GBs was determined based on the Cu oxide coverage that was quantitatively estimated from the optical microscope (OM) images. To investigate the effect of Cu oxidation on graphene quality, all graphene samples were transferred on a polyethylene terephthalate (PET) film and then the electrical properties (*i.e.*, sheet resistance, Hall mobility, and charge carrier density) of these samples were measured as a function of oxidation time and temperature by Hall measurement. We correlated the extent of Cu oxidation and electrical properties of graphene; these results enable the prediction of electrical properties through graphene visualization by Cu oxidation.

Results and discussion

Characterization of as-grown graphene

The single layer graphene (SLG) was synthesized on the Cu foil by rapid thermal chemical vapor deposition (RTCVD) at 1025 °C for 1000 s. As shown in ESI S-I,† no discernible features, *e.g.*, G-GB or other defects, were observed on the as-grown SLG film by the OM image analysis. The sheet resistance (R_s), sheet carrier density (n_s), and Hall mobility (μ) of the SLG transferred on the PET substrate were $254 \Omega \text{ sq}^{-1}$, $1.5 \times 10^{13} \text{ cm}^{-2}$, and $1572 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$, respectively. The n_s of the as-grown SLG was found to be one order of magnitude higher than that of the pristine graphene (without intentional doping) because of the molecular p-

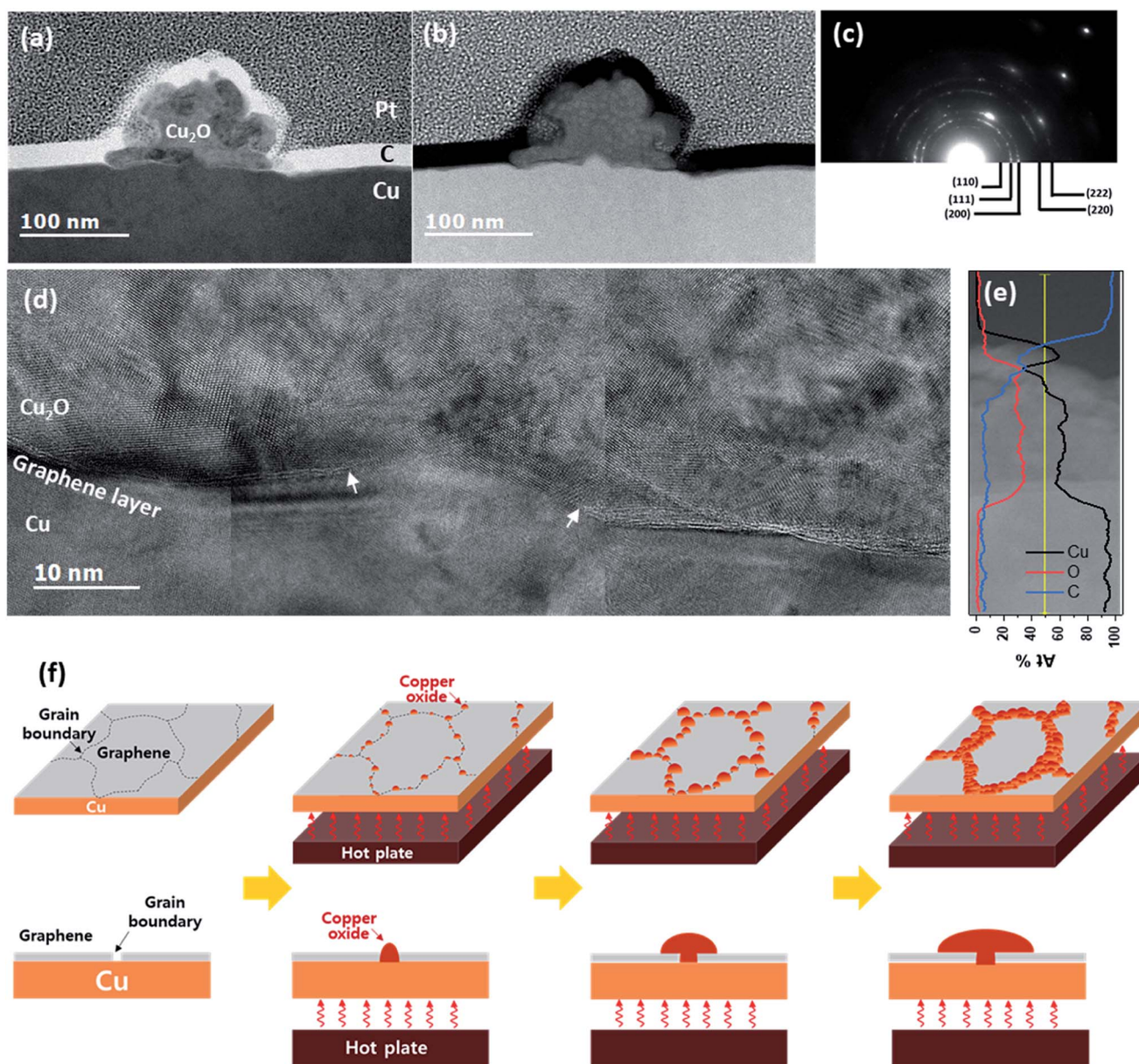


Fig. 2 Cross-sectional TEM images and EDS analysis (a) bright-field STEM image, (b) HAADF STEM image, (c) SAED pattern of Cu_2O , (d) HRTEM image, (e) EDS line spectrum across the interface between Cu and Cu_2O , and (f) schematic diagrams of Cu oxidation through graphene grain boundary.



type doping by the adsorption of imidazole during the wet etching of the Cu foil.²⁹

Cu oxidation kinetics on graphene/Cu foil structure

The as-grown SLG on Cu foil was heat treated under air atmosphere to investigate the Cu oxidation behavior through the graphene. Fig. 1 shows the OM images at each oxidation temperature (T_{ox}) and oxidation time (t_{ox}); Fig. 1(a)–(d), (e)–(h), (i)–(l), and (m)–(p) show the OM images at 180 °C, 200 °C, 220 °C, and 240 °C, respectively. Early heat-treated samples at 180 °C/60 min, 200 °C/30 min, and 220 °C/15 min (Fig. 1(a), (e), and (i), respectively) show dark contrasts at the line expected with G-GBs. Upon increasing the t_{ox} , the dark contrasts not only delineate the grain boundaries of graphene (Fig. 1(c), (g), and (j)) but also increase their line width (Fig. 1(d), (h), and (k), respectively). Eventually, the bright area, presumed to be a graphene grain, is isolated, as shown in Fig. 1(l). At 240 °C, the graphene grain seems to isolate after exposure to air for 10 min and the size of the bright area decreases with increasing t_{ox} . Notably, some samples with similar oxidation morphologies, such as those treated at 180 °C/240 min, 200 °C/120 min, and 220 °C/30 min that are illustrated in Fig. 1(c), (g), and (j), respectively, show that the t_{ox} tends to decrease to half its value with T_{ox} by only 20 °C. As shown in ESI S-II,[†] the energy-dispersive X-ray spectroscopy (EDS) mapping image of sample oxidized at 220 °C for 1 hour clearly shows that a large amount of oxygen is detected in a dark area in OM image which indicates the dark area is Cu oxide. Previous reports^{23,24} on the composition analysis of Cu oxidation under graphene using TEM also suggested that the dark contrast in OM images corresponds to Cu oxide. Therefore, the Cu oxidation, dark contrasts on OM image, begins at the G-GBs and progresses to the periphery of the G-GBs.

The dark area in the sample oxidized at 200 °C for 120 min was analyzed using cross-sectional TEM (Fig. 2). The cross-sectional TEM samples were prepared perpendicular to the

dark lines. Bright-field scanning TEM (STEM) revealed a radially grown morphology on the flat Cu surface (Fig. 2(a)). From the indexing of the selected area diffraction pattern (Fig. 2(c)) and composition analysis of energy dispersive spectroscopy (EDS, Fig. 2(e)), the species was determined as Cu_2O (space group: $Pn\bar{3}m$, $a = 4.288 \text{ \AA}$). Judging from the low contrast in the high-angle annular dark field (HAADF) STEM images (Fig. 2(b)), the Cu_2O had a lower mass density than the Cu substrate. Therefore, the interface between Cu_2O and Cu was very clearly defined. The high resolution TEM (HRTEM) images in this area clarified the graphene layer at the Cu– Cu_2O interface with an opening at the center (marked by arrows in Fig. 2(d)). The graphene layer was also found under other Cu_2O particles (see ESI S-III[†]). The semi-circular shape of the Cu_2O and graphene opening at the center of the interface indicate that Cu_2O is formed from a point source of Cu. Furthermore, graphene existence between Cu_2O and Cu evidenced that the Cu_2O grew by out-diffusion of Cu.³⁰ Indeed, Cu oxidation predominantly occur by the diffusion of Cu cations through Cu oxide.^{31,32} However, these results contradict a few previous reports, which claim that Cu_2O nucleates and grows by oxygen in-diffusion through graphene at the graphene–Cu interface.^{23,24} According to these previous reports, oxygen atoms dissociate from H_2O at the graphene vacancies and GBs then penetrate the G-GBs and form Cu_2O at the graphene–Cu interface.²⁴ The randomly oriented and shaped graphene grains are stitched together, forming abundant pentagon, heptagon, and other non-hexagonal carbon rings at the G-GBs.³³ G-GBs with imperfect stitching and voids would crack during the CVD process. We believe that Cu_2O nucleation was promoted at the non-hexagonal carbon rings or in the imperfect-stitching region of G-GBs but was suppressed at the relatively stable grain boundary. For instance, at 180 °C after 240 min of oxidation, the line width of the grown Cu_2O reached $10 \mu\text{m}$ (Fig. 1(c)), and these locations are considered as the non-hexagonal carbon rings or imperfect-stitching; however, at certain G-GBs where

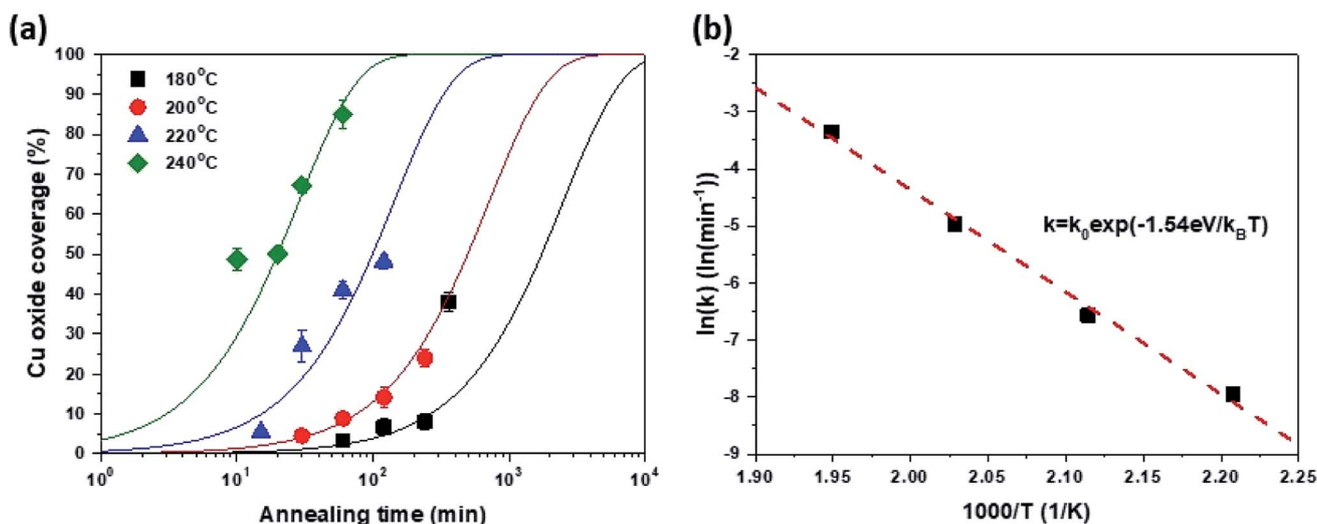


Fig. 3 (a) Cu oxide coverage as a function of oxidation time and (b) oxidation reaction constant as a function of inverse oxidation temperature.



relatively stable G-GBs, still no Cu_2O was formed. We surmise that the initial nucleation process of Cu_2O was generated by oxygen in-diffusion through the G-GBs as previously reported but that Cu_2O grew by the dominant out-diffusion of Cu cations at the G-GBs. Fig. 2(f) shows the schematic sequence of Cu oxidation through graphene based on the results of the current study. During the oxidation, Cu oxide was initially formed at G-GB and further growth was proceeded by out-diffusion of Cu through the Cu oxide. Otherwise, if Cu oxide growth was proceeded by in-diffusion of oxygen to G-GBs, not by out-diffusion of Cu cation, Cu oxide would be encapsulated by the graphene layer. Consequently, the graphene layer undergoes tensile stress generated by 1.7 times of volume expansion during Cu oxide formation, and the estimated strain of graphene is about 80% based on Fig. 3(a). Considering fracture occurs on PMMA and PDMS supported graphene only about 8% of strain during the tensile test,^{34,35} it seems impossible to survive graphene without fracture during Cu oxide formation. However, no significant R_s change is observed even if almost G-GBs have been decorated by Cu_2O (Fig. 1(c), (g) and (i)). Moreover, few samples of graphene seem to be isolated by grown Cu_2O , but the measured R_s shows that graphene samples are still electrically connected (Fig. 1(d), (h), (k), and (m)). Therefore, Cu oxide growth proceeds without further graphene damage caused by the volume expansion during Cu_2O formation because of oxidation proceeds by out-diffusion of Cu cation through formed Cu_2O .

The surface coverage of Cu_2O (F_{ox}) was assessed based on the OM images in Fig. 1. Here, the intensity of the OM image was normalized and the fraction of dark regions was estimated from the intensity histogram (for more details refer to ESI S-VI†). Fig. 3(a) shows the F_{ox} as a function of t_{ox} and T_{ox} . It is worth noting that the change in the F_{ox} as a function of t_{ox} agrees well with the typical Johnson–Mehl–Avrami–Kolmogorov (JMAK) equation [eqn (1)] for predicting phase transformation kinetics based on nucleation and growth.³⁶

$$F = 1 - \exp(-kt^n), \quad (1)$$

where F is the fraction of the transformed phase, t is the transformation time, n is the shape factor, and k is the kinetic parameter related to the transformation rate. n is related to nucleation and growth dimension. If the phase change occurs by nucleation and three-dimensional growth, n has a value of approximately 4. If there is no further nucleation during the growth, *i.e.*, nucleation is not a function of time, n has the same value as the shape dimension. And n has a value of about 3 for three-dimensional growth. In our case, n value was well fitted to 1 at all T_{ox} (for more details refer to ESI S-V†), which is shown as solid lines in Fig. 3(a). It means that the nuclei of Cu_2O were formed at the fixed region of G-GBs, and Cu_2O grows one-dimensionally from the G-GBs without additional nucleation. The one-dimensional growth of Cu_2O is well matched experimental result in Fig. 1 and previous discussions that nucleation of Cu_2O is limited at the non-hexagonal carbon rings or imperfect-stitching region and further nucleation of Cu_2O is suppressed. Here, only 2-dimensional growth was considered with no information on the thickness change because the F_{ox} is the surface coverage. Therefore, because the length of G-GB is fixed, one-dimensional growth implies an increase in the width of Cu oxide from G-GBs.

Furthermore, the k value related to the transformation rate was also extracted based on the JMAK equation in Fig. 3(a) and is plotted as a function of $1/T_{\text{ox}}$ in Fig. 3(b). The slope of the curve fitted by the Arrhenius equation was 1.54 eV, which reflects the activation energy of Cu_2O growth through graphene, representing the lateral growth rate of Cu_2O . Therefore, the growth of Cu_2O through graphene is controlled by Cu out-diffusion through Cu_2O because this activation energy is similar to that of Cu diffusion in Cu oxide estimated in the classical Cu oxidation experiment.³⁷ Consequently, it is possible to predict the kinetics of Cu oxidation through graphene at a certain targeting temperature through the JMAK equation using a k value obtained by extrapolation.

Electrical properties of graphene based on Cu oxidation

Previous results showed that Cu oxide grows on the graphene surface by out-diffusion of Cu ions through Cu oxide. To

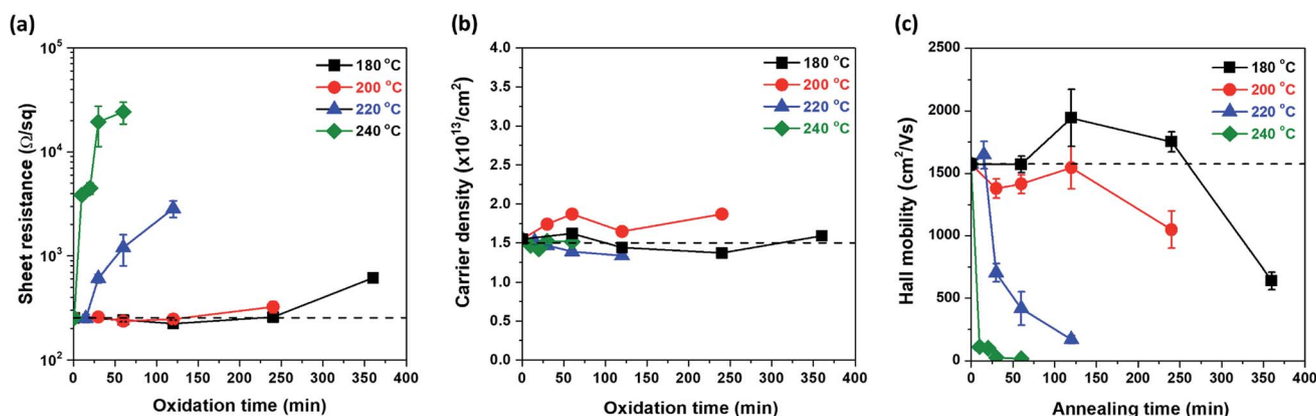


Fig. 4 Electrical properties of graphene as a function of the oxidation time at different oxidation temperatures (180 °C, 200 °C, 220 °C, and 240 °C); (a) sheet resistance (b) sheet carrier density, and (c) Hall mobility with dashed lines which represent the values of the as-grown graphene.



investigate the quality change in the graphene during Cu oxidation, the electrical properties of all samples were measured. All oxidized samples, as shown in Fig. 1, were transferred on a PET film; then, the R_s of graphene was examined by Hall measurement using the van der Pauw structure. The change in R_s as a function of t_{ox} at each T_{ox} is illustrated in Fig. 4(a). At T_{ox} of 180 °C, 200 °C, and 220 °C, the R_s values did not significantly increase compared to those of the as-grown graphene ($R_{s0} = 254 \Omega \text{ sq}^{-1}$) until 240, 120, and 15 min of oxidation, respectively. However, at T_{ox} of 240 °C, R_s sharply increased after only 10 min of oxidation. To determine the cause of the change in R_s , n_s and μ were measured as a function of t_{ox} by Hall measurement (Fig. 4(b) and (c)). Interestingly, n_s maintains a value of $1.5 \times 10^{13} \text{ cm}^{-2}$, regardless of T_{ox} and t_{ox} . However, there was a drastic change μ (Fig. 4(c)).

Because the electrical properties depend on both T_{ox} and t_{ox} , it is difficult to quantitatively determine the effect of Cu oxidation on the electrical properties of graphene. Therefore, by expressing T_{ox} and t_{ox} in terms of F_{ox} , the electrical properties of graphene were plotted as a function of F_{ox} (Fig. 5), in which the dashed lines represent the values of the as-grown graphene. As mentioned, n_s was independent of T_{ox} and t_{ox} and remained constant until $F_{\text{ox}} = 85\%$ (Fig. 5(b)). The R_s and μ had different curves for each T_{ox} , but these curves were merged into one curve by plotting it as a function of F_{ox} . It clearly indicates that electrical properties of graphene are strongly related to F_{ox} . From $R_s = 1/n_s q \mu$, the change in R_s as a function of F_{ox} is fully described by the change of μ , inversely corresponding to the change of μ , because of the constant n_s . The change in μ as a F_{ox} is divided into three regions. In the first region (under $F_{\text{ox}} = 15\%$), the value of μ stayed at the as-grown value, and then the value of μ

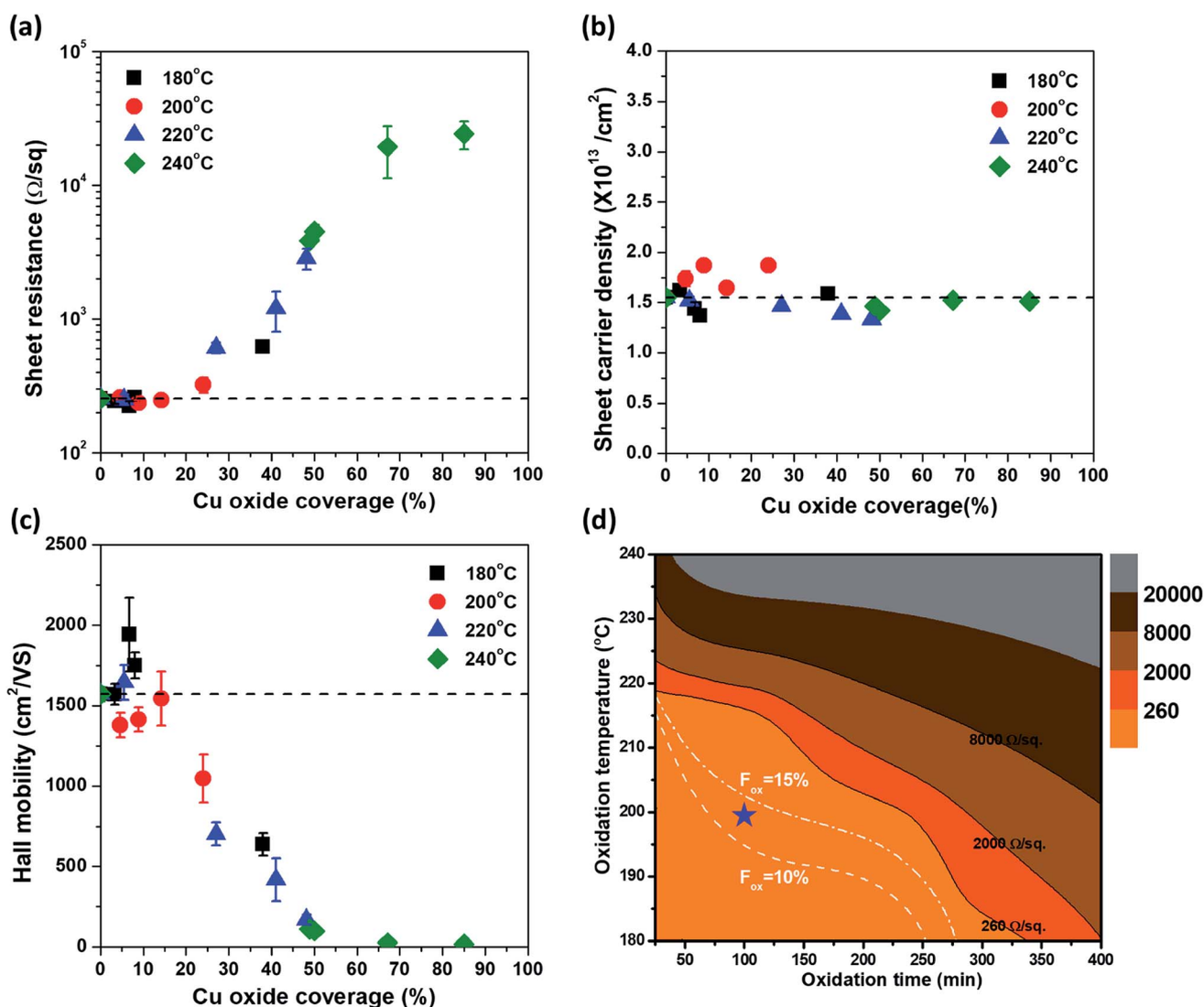


Fig. 5 Electrical properties of graphene as a function of Cu oxide coverage at different oxidation temperature (180 °C, 200 °C, 220 °C, 240 °C); (a) sheet resistance, (b) sheet carrier density, and (c) Hall mobility. The black dashed lines represent the values of the as-grown graphene. (d) Contour map of graphene sheet resistance with white dashed lines which indicate constant F_{ox} lines of 10% and 15%. The blue star represents the optimal condition of G-GB visualization.



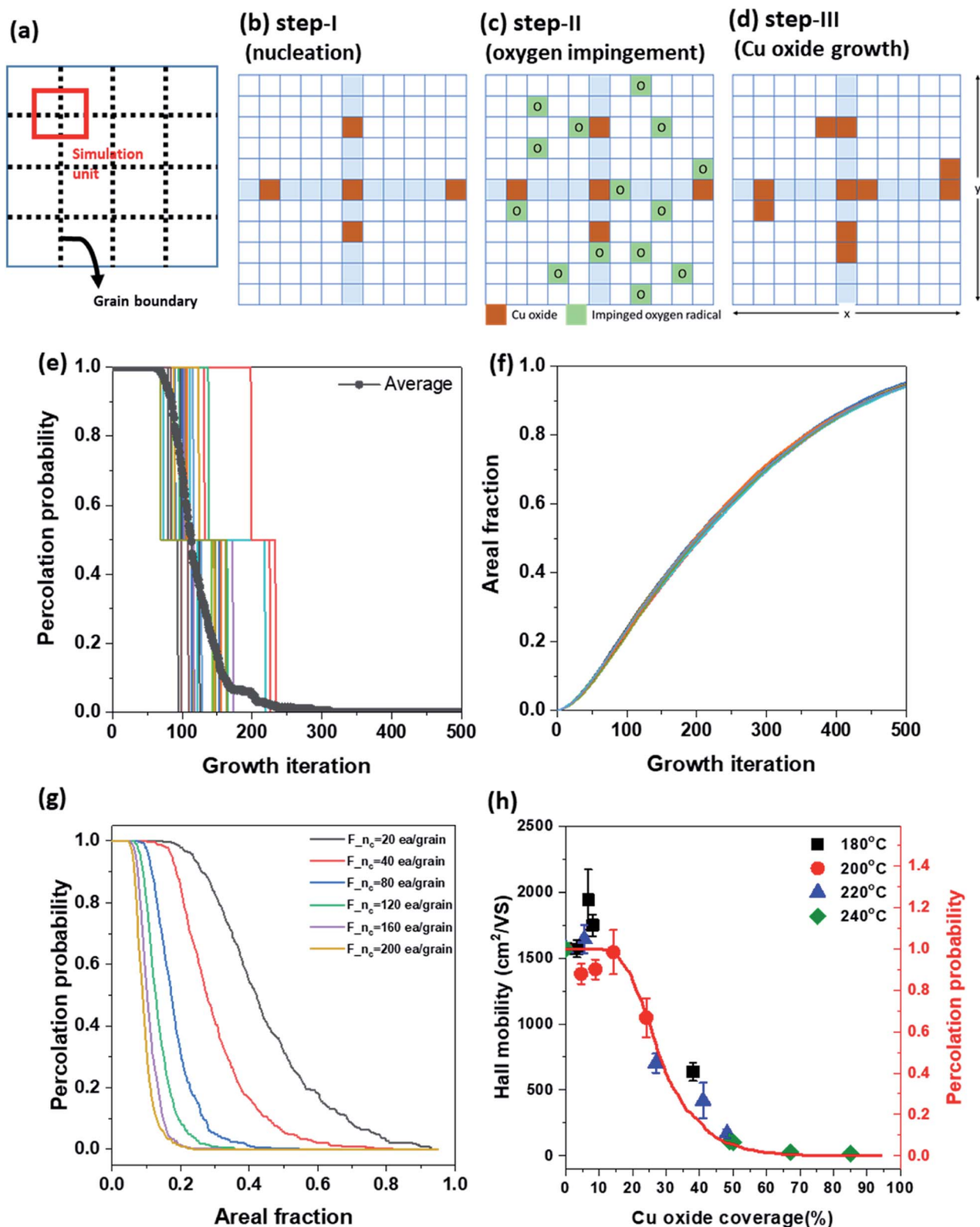


Fig. 6 The relationship between Hall mobility and graphene percolation probability as a function of Cu oxide coverage: (a) schematics of the simulation unit and simulation step ((b) nucleation, (c) oxygen impingement, and (d) Cu oxide growth), (e) percolation probability and (f) areal fraction as a function of growth iteration, (g) percolation probability as a function of oxidation fraction, and (h) Hall mobility and percolation probability at $F_{n_c} = 40$ ea per grain as a function of the oxidation fraction.



Table 1 Oxidation conditions of Cu coated by graphene

| Temperature (°C) | Oxidation time (min) | | | |
|------------------|----------------------|-----|-----|-----|
| 180 | 60 | 120 | 240 | 360 |
| 200 | 30 | 60 | 120 | 240 |
| 220 | 15 | 30 | 60 | 120 |
| 240 | 10 | 20 | 30 | 60 |

decreases linearly with the increase in F_{ox} in the second region ($F_{\text{ox}} = 15\text{--}50\%$). Finally, the value of μ was only a few tens of $\text{cm}^2 \text{V}^{-1} \text{s}^{-1}$ over $F_{\text{ox}} = 50\%$. Herein, Cu_2O is formed on the graphene surface and hence is undamaged by volume expansion (1.7 times) from Cu to Cu_2O . However, as a defect site of the G-GB, the initial Cu_2O nucleation site is the pathway of Cu cation out-diffusion (see the HRTEM image in Fig. 2(d)). The initial nucleation site is considered as a degraded G-GB point even before the Cu_2O growth; thus, it may negligibly affect the electrical properties of graphene up to $F_{\text{ox}} = 15\%$ of Cu_2O growth. However, the continuous out-diffusion of Cu cations through the nucleation sites can expand and propagate the Cu cation diffusion path along with the G-GBs. When F_{ox} exceeds 15%, the propagation of the Cu cation diffusion path becomes severe and μ of graphene is reduced. At F_{ox} values above 50%, most of the graphene grains seem to be electrically disconnected. As electrical measurement inevitably involves charge carrier transfer, the mechanically weak points of the damaged grain boundaries can be further damaged during the transfer process, thereby exaggerating the decreasing tendency of the μ (see ESI S-VI†). From these results, we tried to provide optimal conditions for G-GB visualization through the contour map of R_s as a function of T_{ox} and t_{ox} as shown in Fig. 5(d). The optimal condition requires a F_{ox} enough to measure the graphene grain size at the same time without changing the electrical properties of graphene. Therefore, as mentioned above, the maximum value of F_{ox} can be defined as 15%. On the other hand, G-GB is sufficiently revealed for measuring the graphene grain size when the amount of oxidation is at least about Fig. 1(b) and (f), and the F_{ox} at this time is 10%. The boundary for optimal F_{ox} values is shown as dashed lines in Fig. 5(d). The difference in oxidation time indicated by the distance between dashed lines at 200 °C is 50 min, so the process margin is very wide and oxidation time is relatively short which is compared to 180 °C. Therefore, oxidation at 200 °C for 100 min, can be suggested as an optimal condition for G-GB visualization.

To explain the propagation of damage on G-GBs and electrical disconnection in the view point of Hall mobility variation as a function of F_{ox} , the percolation probability was introduced in a simple schematic model of Cu oxidation at G-GBs. The percolation probability is used for conductivity prediction as a fraction of metallic component in composite materials.³⁸ Because there is no mathematical formula for calculating the percolation probability, the percolation probability of graphene is simulated through a simple model, *i.e.*, the graphene is disconnected as Cu oxide randomly nucleates and grows in G-GBs. The graphene grains in the 2D matrix were assumed to have square shapes to simplify the calculation (dashed line in

Fig. 6(a)). Based on the morphological change illustrated in Fig. 1, we assumed the following steps. Step-I-nucleation, at the beginning, Cu oxide randomly nucleates at the G-GBs and there is no additional nucleation during the growth step (Fig. 6(b)). Step-II-oxygen impingement, the oxygen impingement is random (Fig. 6(c)). Step-III-Cu oxide growth, Cu oxide grows when the oxygen impinged near the already existing Cu oxide (Fig. 6(d)). After step-I-nucleation, step-II and step-III were repeated to grow Cu oxide. The random nucleation and continuous growth of Cu_2O according to the iteration was provided in ESI S-VII† in the form of moving images. The connected graphene and Cu oxide coverage were estimated in each growth step using MATLAB software. The percolation probability, $P(F_{\text{ox}})$, was 1 when the conducting path of graphene exists in both the x and y directions, 0.5 when the conducting path of graphene exists only in one direction (x or y), and 0 when the graphene is isolated by Cu oxidation. Fig. 6(e) and (f) show the graphene percolation probability and Cu oxide coverage, respectively, as a function of growth iteration. Each step function in Fig. 6(e) indicates 20 times repeated percolation probability with different initial positions of Cu oxide nucleus, and its average is shown as a gray solid curve. As shown in Fig. 6(f), the Cu oxide coverage increases with growth iteration, regardless of the initial position of the Cu oxide nucleus, whereas the graphene percolation probability is strongly affected by the initial position of the Cu oxide nucleus. Moreover, to investigate the effect of the amount of initial Cu oxide nucleus that might be related to G-GBs quality, the percolation probability was plotted as a function of the Cu oxide coverage [$P(F_{\text{ox}})$] by averaging 200 simulation units and varying the initial nucleus density (F_{n_c}) from 20 to 200 ea/grain (as the solid lines in Fig. 6(g)). As oxidation progressed, there was a region where $P(F_{\text{ox}}) = 1$ was maintained, after which $P(F_{\text{ox}})$ decreased rapidly and eventually the conducting path of graphene disappeared [$P(F_{\text{ox}}) = 0$]. $P(F_{\text{ox}}) = 1$ reflects that the presence of the 2-directional conducting path in graphene and no disturbance to the charge transfer in the graphene. As $P(F_{\text{ox}}) < 1$, the conducting path between graphene grains start to disappear, indicating the disturbance of the charge transfer in the graphene. The simulation results revealed that the F_{ox} range at $P(F_{\text{ox}}) = 1$ was determined by the initial nuclei density of Cu oxide at G-GB, which is related to the defect concentration in G-GB. This change in $P(F_{\text{ox}})$ at $F_{n_c} = 40$ ea per grain is very consistent with the change in the measured Hall mobility as shown in Fig. 6(h). Therefore, the relationship between the oxidized graphene morphology on Cu and its electrical properties was determined and the extent of morphological variation was assessed without affecting the graphene electrical properties. This simple modeling of Cu oxidation through graphene enables the calculation of the percolation probability of spanning graphene as a function of surface coverage of the Cu oxide, which is closely related to the carrier mobility within graphene.

Conclusion

Cu oxidation through graphene was investigated in detail as a function of temperature and time. The kinetics of Cu



oxidation through graphene were quantitatively determined based on the Cu oxide coverage that was determined from the OM image. There are three unique features in discussing Cu₂O growth behavior on graphene. First, Cu₂O growth kinetics undergoes one-dimensional growth. The OM image and JMAK equation analysis shows the Cu₂O nucleation is limited in G-GBs and the Cu₂O one-dimensionally grows with suppressed further nucleation. Second, the Cu₂O growth is governed by dominant Cu cation out-diffusion in according to that; (i) HR-TEM analysis reveals graphene layer is at the Cu–Cu₂O interface, (ii) the Cu oxidation predominantly occurs by dominant out-diffusion of Cu cation, (iii) the graphene layer is not critically damaged by the volume expansion of Cu oxide. Third, F_{ox} of 10–15% confirmed that the graphene grain size could be measured without degradation of the electrical property of graphene, and suggested that the optimal condition considering practical points of experiments such as process margin is oxidation at 200 °C for 100 min. Also, the percolation probability could be predicted by simple modeling of Cu oxidation through graphene, the carrier mobility in graphene can also be estimated.

Methods

Synthesis of graphene

The graphene was synthesized by RTCVD (NPS Corporation) on a 370 × 470 mm² Cu foil (thickness: 0.35 mm, Nippon Mining & Metals Corporation). The Cu foil was loaded into a chamber and heated to 800 °C under CH₄ atmosphere (30 sccm, 550 mTorr). The surface of Cu foil was then cleaned using pre-annealing, and the crystalline orientation of Cu was rearranged. Next, the graphene synthesis was performed by reheating the foil to 1025 °C for 1000 s. The chamber was cooled down to 600 °C and then to the room temperature by injecting CH₄ and N₂ gases, respectively.

Cu oxidation on graphene/Cu foil structure

After the deposition, the graphene grown on Cu foil was cut into specimens of 3 × 3 cm², and the samples were subjected to heat treatment on a hot plate for different durations at various temperatures under air atmosphere. Four different temperatures were chosen: 180 °C, 200 °C, 220 °C, and 240 °C. The oxidation time was varied from 10 to 360 min (Table 1).

Characterization of graphene and Cu oxide

After the oxidation of graphene on the Cu foil, surface images were acquired with OM (Olympus BX50). To identify the dark contrast of OM image, scanning electron microscope and Energy-dispersive X-ray spectroscopy (Merlin Compact, ZEISS) was conducted. The cross-sectional microstructure of graphene on Cu after oxidation was observed using Cs-corrected TEM (ARM 200F, JEOL Ltd.). The as-grown and heat-treated graphenes on Cu were transferred on a polymethyl methacrylate (PMMA) supporting layer in order to investigate the electrical properties of graphene. The PMMA was spin coated onto the graphene on Cu and then the underlying Cu was wet etched in

imidazole-based Cu etchant (ammonium persulfate (0.1 M) + H₂SO₄ + H₂O₂). Thus, the graphene was simultaneously doped during the etching of Cu.²⁹ The floated PMMA/graphene layer was scooped after 8 hours of complete Cu etching onto PET. Finally, the top PMMA layer was removed by acetone. The graphene electrical properties, *i.e.*, sheet resistance, Hall mobility, and sheet carrier density, were measured by using a van der Pauw structure of 8 × 8 mm². Hall measurements were performed under a 0.5 T magnetic field (HL 5500PC, BIO-RAD) at room temperature.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 K. S. Novoselov, A. K. Geim, S. V. Morozov, D. Jiang, Y. Zhang, S. V. Dubonos, I. V. Grigorieva and A. A. Firsov, *Science*, 2004, **306**, 666–669.
- 2 X. S. Li, W. W. Cai, J. H. An, S. Kim, J. Nah, D. X. Yang, R. Piner, A. Velamakanni, I. Jung, E. Tutuc, S. K. Banerjee, L. Colombo and R. S. Ruoff, *Science*, 2009, **324**, 1312–1314.
- 3 H. C. Lee, W. W. Liu, S. P. Chai, A. R. Mohamed, A. Aziz, C. S. Khe, N. M. S. Hidayah and U. Hashim, *RSC Adv.*, 2017, **7**, 28427.
- 4 X. S. Li, C. W. Magnuson, A. Venugopal, R. M. Tromp, J. B. Hannon, E. M. Vogel, L. Colombo and R. S. Ruoff, *J. Am. Chem. Soc.*, 2011, **133**, 2816–2819.
- 5 Q. K. Yu, L. A. Jauregui, W. Wu, R. Colby, J. F. Tian, Z. H. Su, H. L. Cao, Z. H. Liu, D. Pandey, D. G. Wei, T. F. Chung, P. Peng, N. P. Guisinger, E. A. Stach, J. M. Bao, S. S. Pei and Y. P. Chen, *Nat. Mater.*, 2011, **10**, 443–449.
- 6 H. L. Zhou, W. J. Yu, L. X. Liu, R. Cheng, Y. Chen, X. Q. Huang, Y. Liu, Y. Wang, Y. Huang and X. F. Duan, *Nat. Commun.*, 2013, **4**, 2096.
- 7 S. Y. Cho, M. S. Kim, M. Kim, K. J. Kim, H. M. Kim, D. J. Lee, S. H. Lee and K. B. Kim, *Nanoscale*, 2015, **7**, 12820–12827.
- 8 L. A. Jauregui, H. L. Cao, W. Wu, Q. K. Yu and Y. P. Chen, *Solid State Commun.*, 2011, **151**, 1100–1104.
- 9 G. Lopez-Polin, C. Gomez-Navarro, V. Parente, F. Guinea, M. I. Katsnelson, F. Perez-Murano and J. Gomez-Herrero, *Nat. Phys.*, 2015, **11**, 26–31.
- 10 F. Hao, D. N. Fang and Z. P. Xu, *Appl. Phys. Lett.*, 2011, **99**, 041901.



- 11 Z. Yan, J. Lin, Z. W. Peng, Z. Z. Sun, Y. Zhu, L. Li, C. S. Xiang, E. L. Samuel, C. Kittrell and J. M. Tour, *ACS Nano*, 2012, **6**, 9110–9117.
- 12 S. S. Chen, H. X. Ji, H. Chou, Q. Y. Li, H. Y. Li, J. W. Suk, R. Piner, L. Liao, W. W. Cai and R. S. Ruoff, *Adv. Mater.*, 2013, **25**, 2062–2065.
- 13 Y. F. Hao, M. S. Bharathi, L. Wang, Y. Y. Liu, H. Chen, S. Nie, X. H. Wang, H. Chou, C. Tan, B. Fallahazad, H. Ramanarayanan, C. W. Magnuson, E. Tutuc, B. I. Yakobson, K. F. McCarty, Y. W. Zhang, P. Kim, J. Hone, L. Colombo and R. S. Ruoff, *Science*, 2013, **342**, 720–723.
- 14 L. M. Malard, M. A. Pimenta, G. Dresselhaus and M. S. Dresselhaus, *Phys. Rep.*, 2009, **473**, 51–87.
- 15 A. C. Ferrari and D. M. Basko, *Nat. Nanotechnol.*, 2013, **8**, 235–246.
- 16 F. Tuinstra and J. L. Koenig, *J. Chem. Phys.*, 1970, **53**, 1126–1130.
- 17 A. Das, S. Pisana, B. Chakraborty, S. Piscanec, S. K. Saha, U. V. Waghmare, K. S. Novoselov, H. R. Krishnamurthy, A. K. Geim, A. C. Ferrari and A. K. Sood, *Nat. Nanotechnol.*, 2008, **3**, 210–215.
- 18 K. Kim, Z. Lee, W. Regan, C. Kisielowski, M. F. Crommie and A. Zettl, *ACS Nano*, 2011, **5**, 2142–2146.
- 19 J. W. Suk, A. Kitt, C. W. Magnuson, Y. F. Hao, S. Ahmed, J. H. An, A. K. Swan, B. B. Goldberg and R. S. Ruoff, *ACS Nano*, 2011, **5**, 6916–6924.
- 20 C. C. Jia, J. L. Jiang, L. Gan and X. F. Guo, *Sci. Rep.*, 2012, **2**, 707.
- 21 J. Y. Lee, J. H. Lee, M. J. Kim, J. K. Dash, C. H. Lee, R. Joshi, S. Lee, J. Hone, A. Soon and G. H. Lee, *Carbon*, 2017, **115**, 147–153.
- 22 K. P. Hong, D. Lee, J. B. Choi, Y. Kim and H. Kim, *ACS Appl. Nano Mater.*, 2018, **1**, 2515–2520.
- 23 D. L. Duong, G. H. Han, S. M. Lee, F. Gunes, E. S. Kim, S. T. Kim, H. Kim, Q. H. Ta, K. P. So, S. J. Yoon, S. J. Chae, Y. W. Jo, M. H. Park, S. H. Chae, S. C. Lim, J. Y. Choi and Y. H. Lee, *Nature*, 2012, **490**, 235–239.
- 24 J. Kwak, Y. Jo, S. D. Park, N. Y. Kim, S. Y. Kim, H. J. Shin, Z. Lee, S. Y. Kim and S. Y. Kwon, *Nat. Commun.*, 2017, **8**, 1549.
- 25 C. Kunka, S. Bavdekar, N. G. Rudawski, A. Fournier and G. Subhash, *J. Phys. Mater.*, 2019, **2**, 025005.
- 26 T. Yoon, J. H. Mun, B. J. Cho and T. S. Kim, *Nanoscale*, 2014, **6**, 151–156.
- 27 S. S. Chen, L. Brown, M. Levendorf, W. W. Cai, S. Y. Ju, J. Edgeworth, X. S. Li, C. W. Magnuson, A. Velamakanni, R. D. Piner, J. Y. Kang, J. Park and R. S. Ruoff, *ACS Nano*, 2011, **5**, 1321–1327.
- 28 J. M. Wofford, S. Nie, K. F. McCarty, N. C. Bartelt and O. D. Dubon, *Nano Lett.*, 2010, **10**, 4890–4896.
- 29 K. Jo, S. M. Kim, S. M. Lee, J. H. Kim, H. J. Lee, K. S. Kim, Y. D. Kwon and K. S. Kim, *Carbon*, 2015, **82**, 168–175.
- 30 S. K. Lee, H. C. Hsu and W. H. Tuan, *Mater. Res.-Ibero-Am. J.*, 2016, **19**, 51–56.
- 31 D. W. Bridges, J. P. Baur, G. S. Baur and W. M. Fassell, *J. Electrochem. Soc.*, 1956, **103**, 475–478.
- 32 S. Mrowec and A. Stoklosa, *Oxid. Met.*, 1971, **3**, 291–311.
- 33 L. P. Biro and P. Lambin, *New J. Phys.*, 2013, **15**, 035024.
- 34 H. Jang, Z. H. Dai, K. H. Ha, S. K. Ameri and N. S. Lu, *2D Mater.*, 2020, **7**, 014003.
- 35 S. H. Bae, Y. Lee, B. K. Sharma, H. J. Lee, J. H. Kim and J. H. Ahn, *Carbon*, 2013, **51**, 236–242.
- 36 M. Avrami, *J. Chem. Phys.*, 1939, **7**, 1103–1112.
- 37 Y. Ebisuzaki and W. B. Sanborn, *J. Chem. Educ.*, 1985, **62**, 341–343.
- 38 A. L. Efros and B. I. Shklovskii, *Phys. Status Solidi B*, 1976, **76**, 475–485.

