Reversible defect engineering in graphene grain boundaries

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Research efforts in large area graphene synthesis have been focused on increasing grain size. Here, it is shown that, beyond 1μm grain size, grain boundary engineering determines the electronic properties of the monolayer. It is established by chemical vapor deposition experiments and first-principle calculations that there is a thermodynamic correlation between the vapor phase chemistry and carbon potential at grain boundaries and triple junctions. As a result, boundary formation can be controlled, and well-formed boundaries can be intentionally made defective, reversibly. In 100μm long channels this aspect is demonstrated by reversibly changing room temperature electronic mobilities from 1000 to 20,000 cm² V⁻¹ s⁻¹. Water permeation experiments show that changes are localized to grain boundaries. Electron microscopy is further used to correlate the global vapor phase conditions and the boundary defect types. Such thermodynamic control is essential to enable consistent growth and control of two-dimensional layer properties over large areas.
he chemical vapor deposition (CVD) of graphene for large area applications is typically carried out by transition metal (Cu, Pt, Co, Cr, Rb etc.) catalyzed hydrocarbon (methane, $\text{C}_2\text{H}_6$, $\text{C}_3\text{H}_8$ etc.) decomposition at temperatures in excess of 950 °C. Continuous monolayers are formed on the metal surface by the nucleation and coalescence of graphene domains. Hence, grain boundaries (GB) and triple junctions that are a part of any polycrystalline material are also present in large area monolayer graphene\(^{2–9}\). These GB have been extensively studied and their defect density has been theoretically shown to be mainly dependent on the extent of misorientation between the grains.\(^{10–12}\) However, in CVD graphene, the defect density at a boundary is not determined by the misorientation angle alone. Tsen et al.\(^{13,14}\) studied the effect of source gas flow conditions on the quality and type of GB structures formed in CVD grown graphene. Using transmission electron microscopic (TEM) analysis of the boundary along with electrical transport measurements, they reported that the samples grown at higher methane flows had smaller grains, uniform coverage and most importantly, lower GB resistance. Samples grown using lower methane flows were found to have boundaries with overlaps and voids, resulting in higher resistance.\(^{14}\) The results are markedly important in two aspects. First, the GB defect density was shown to depend not only on the geometric aspects of the grains such as tilt misorientation and edge type, but also on gas flow conditions. Second, electrical transport characterization was found to be an effective tool to inspect the defect density of GB. Their study, however, did not address the correlation between the physico-chemical state of the growth system and defects at GB. More recently, calculations by Dong et al.\(^{15}\) showed that one can toggle between covalently bonded boundaries and hydrogen terminated edges by changing hydrogen pressure.

In this article, it is shown that there exists a thermodynamic relationship between the gas phase and defects in the deposited monolayer. Such defects, which are shown to be predominantly located at the domain coalescence boundaries, can be healed and then re-created, by exposing the monolayer to different corresponding carbon potentials. Hence, monolayer properties can be tuned. As an illustration, it is shown that the charge mobility ($\mu_{\text{e}}$) and sheet resistance ($R_{\text{s}}$) can be reversibly changed by more than an order of magnitude. This control has allowed us to synthesize graphene with $\mu_{\text{e}}$ of 20,000 cm$^2$ V$^{-1}$ s$^{-1}$ in device with dimensions of 100 μm × 100 μm. Our results are compared against reported literature in Supplementary Note 1. All these improvements on the monolayer happen in a regime in which Raman measurements\(^{16,17}\) are insensitive to the changes occurring in it. Hence, to supplement the electrical transport measurements, water permeation measurement technique is used to identify and spatially locate the incorporated changes in the monolayer. The results highlight the importance of GB closure in obtaining graphene with the best of electronic properties over large areas, in the regime in which the yields of increasing grain size start diminishing (Supplementary Note 2).\(^{18}\) In addition, there are many applications that can benefit from the controlled introduction of defects at graphene boundaries. These include the ability to control molecular permeation and sensitivity to various chemicals.\(^{19,20}\)

### Results

#### Nucleation, growth, and annealing at constant $\Delta G$

Graphene synthesis using Cu catalyzed dehydrogenation of methane can be described by the following equation.

$$\text{CH}_4 \leftrightarrow \text{C}_{\text{Cu}} + 2\text{H}_2$$  \hspace{1cm} (1)

$\text{C}_{\text{Cu}}$ refers to the carbon monolayer (graphene) on the Cu surface. Equation (1) represents only the thermodynamic terminal points, whereas the internal pathways of the system can be many and are still being debated.\(^{21}\) The free-energy change driving the growth or supersaturation ($\Delta G$) is then given by

$$\Delta G = R\text{T}ln(\frac{P_2}{P_1})$$  \hspace{1cm} (2)

where, $K = P_2/\text{P}_1$, $K^\text{eq}$ is the value $K$ at equilibrium, $P_\text{CH}_4$, $P_\text{H}_2$, $T$, and $R$ represent $\text{CH}_4$ partial pressure, $\text{H}_2$ partial pressure, temperature, and the gas constant, respectively. Details of the experimental conditions used and the resulting supersaturation, calculated using standard formation energies\(^{22}\), are given in the Supplementary Note 3 for a range of methane partial pressures. Four groups of samples, $G_1$–$G_4$, were studied as summarized in Table 1. The growth details of four specific samples, $S_1$–$S_4$, one from each of these groups, to be discussed in this paper are presented in Fig. 1a below and also described in Table 1.

In the schematic representation shown in Fig. 1a, a heat-up stage and Cu foil anneal stage are used to set the reactor temperature and anneal the substrate, respectively. Supersaturation, $\Delta G = G_{\text{reactants}} – G_{\text{products}}$, is established in the reactor in the growth stage during which sequential methane dehydrogenation produces a population of adatom species on the Cu surface. Graphene growth then proceeds through the four stages of incubation, nucleation, growth, and coalescence. Following the 6-minute point, certain samples, see Table 1, are treated with two kinds of post-growth anneal procedures PGA-1 and PGA-2, without breaking vacuum. Finally, the samples are cooled down under the growth condition to room temperature.

### Table 1 List of sample types and descriptions

| Sample group type | Description |
|-------------------|-------------|
| $G_1$             | Standard graphene. Samples grown at a constant $\Delta G$ of 36 kJ mol$^{-1}$ from 0 and up to 14 min. After 6 min a complete monolayer is observed in the SEM and is called visually coalesced graphene (VCG). This 6-min graphene is sample $S_1$. Points corresponding to sample $S_1$ are indicated in Fig. 1a, i. Electrical data from sample $S_1$ are presented in Supplementary Table 1, row number 5. |
| $G_2$             | Reverse annealed graphene. Samples obtained by first growing up to point VCG at 36 kJ mol$^{-1}$, sample $S_1$ above, and then exposing to $\Delta G$ values higher than 36 kJ mol$^{-1}$ in stage PGA-1 in Fig. 1a without breaking vacuum. Points corresponding to the specific sample $S_2$ from this group, which was annealed at $\Delta G$ of 82 kJ mol$^{-1}$, are indicated in Fig. 2a, b. Electrical data from sample $S_2$ is presented in Supplementary Table 1, row number number 14. |
| $G_3$             | Annealed graphene. Samples obtained by first growing up to point VCG at 36 kJ mol$^{-1}$, sample $S_1$ above, and then exposing to $\Delta G$ values lower than 82 kJ mol$^{-1}$ and down to 36 kJ mol$^{-1}$ in stage PGA-2 in Fig. 1a, without breaking vacuum. Point corresponding to the sample $S_3$ from this group, re-annealed at $\Delta G$ of 36 kJ mol$^{-1}$ is indicated in Fig. 2b. Electrical data from sample $S_3$ is presented in Supplementary Table 1, row number number 18. |
| $G_4$             | Hydrogen-treated graphene. Samples obtained by first performing growth as in sample $S_2$ above and then exposing the sample to pure hydrogen ambient, $\Delta G < 0$ in PGA-2 without breaking vacuum. Electrical data from sample $S_4$ is presented in Supplementary Table 1, row number 19. |
At $\Delta G \leq 32 \text{ kJ mol}^{-1}$, graphene grains either did not nucleate or the rate was so slow that no growing domains were observed until a reasonable growth period of 20 min. Upon increasing the $\Delta G$ to 36 kJ mol$^{-1}$, group $G_1$, the surface remained unmodified up to 1 min as shown in Fig. 1b. Nuclei of size $\sim 2 \mu$m were observed at 2.5 min as shown in Fig. 1c. The individual grains then grew and started to coalesce. Figure 1d shows a partially covered graphene layer after 4 min of growth. Finally, after about 6 min, the Cu surface was “completely covered” as can be seen in Fig. 1e. The graphene film after 6 min of growth will be referred to as “visually coalesced graphene (VCG)” as the SEM micrographs indicate complete coverage and thereafter no further change is detected by SEM. This is sample $S_1$. Surface images of Cu growth surface at 10 mins and 13 mins are shown in figure 1f, g, respectively. The average film defect density extracted from Raman spectroscopic measurements, presented in figure 1h, (see Supplementary Note 4 for Raman measurement details), also mirrors this temporal evolution. The $I_D/I_G$ ratio decreases sharply up to a growth time of 6 min and then saturates in the period from 6 to 14 min. Thus, it appears from these characterization methods that monolayer deposition is complete by 6 min. Graphene growth by CVD on Cu is typically modeled as a self-limiting catalytic process, which can explain the SEM snapshots and kinetic growth saturation previously observed in $G_1$.

The time evolution of film defect density as measured by Raman spectroscopy. The charge mobility ($\mu_{cm}$) and sheet resistance ($R_s$) evolution with growth time. While $\mu_{cm}$ shows a four-fold change after visual coalescence at 6 min, Raman defect density remains almost unchanged. A Gompertz-Sigmoidal fit (blue continuous line in i) to the charge mobility is also shown.
14 min. These observations belong to the sample set G₁ in which a constant Δ₆ of 36 kJ mol⁻¹ was maintained through all stages of growth. This sharp change in electronic properties, beyond the point at which the other structural characterization methods and Raman defect density measurements do not suggest any significant change in the quality of graphene is the first interesting observation. All the electrical measurements, except for samples grown below 4 min (where the film was not completely coalesced), were performed on 100 µm by 100 µm graphene devices (See Supplementary Notes 5–7, for more details regarding electrical measurements).

Tuning GB and electron mobility by varying Δ₆. Further experiments, sample group G₂ in Table 1, were conducted to determine whether μ_cm and sheet resistance can be further improved with the grain size unchanged. Graphene layers were deposited with Δ₆ conditions kept the same as in sample S₁ in the growth stage (see Fig. 1a). This ensured that the average graphene grain sizes were constant across all samples in this group. Post the VCG stage, these samples were annealed without breaking vacuum for 6 min with higher Δ₆ values up to 112 kJ mol⁻¹. A different growth run was used for each value of Δ₆ used for annealing (see Supplementary Table 1). This is denoted as PGA-1 in Fig. 1a. At higher Δ₆, growth transients were expected to attain the final steady states of interest within 6 min. Hence, a time sequence such as the one in Fig. 1i was not performed for these higher Δ₆ values.

The rather dramatic effects of increasing Δ₆ on the saturation μ_cm are shown in Fig. 2a and sheet resistance is shown in Supplementary Note 6. The charge mobility μ_cm increases by 20-fold from 1000 to 20,000 cm² V⁻¹ s⁻¹. The standard deviation in the observed values was < 10% of the mean values and is shown in the Fig. 2a. Correspondingly, there is a further five-fold reduction in the steady state sheet resistance value to 159 Ω⁻¹ cm⁻¹ on raising Δ₆ from 36 to 82 kJ mol⁻¹. This is the lowest reported sheet resistance thus far (see Supplementary Note 1 and 2 for a compilation of sheet resistances and μ_cm reported in the literature) in large area CVD graphene and over such large device sizes (100 µm × 100 µm). The Raman I_D/I_G ratio again remains flat. Thus, from the VCG stage at which the structural characterization methods seem to become insensitive to changes in the graphene layer, further processing has thus resulted in a 20-fold increase in μ_cm. This value is also five-fold higher than the saturation mobility observed under a constant Δ₆ = 36 kJ mol⁻¹ in G₁ (Fig. 1i). The range of sheet resistances achieved here surpass the values obtained in the literature and its comparison is presented in Supplementary Figure 2a.

Additional experiments were performed to determine whether the rise in mobility in Fig. 2a is reversible. Graphene layers were deposited under conditions that yielded the highest μ_cm and lowest sheet resistance values (after a treatment under Δ₆ = 82 kJ mol⁻¹), sample S₂. Without breaking vacuum these samples were now exposed to decreasing values of Δ₆ down to 36 kJ mol⁻¹ for 4 min and 12 min. A different growth run was used for each value of lower Δ₆ used during the reversal. It was observed that the difference in μ_cm values between the 4-min exposure and the 12-min exposure was within experimental error. Hence, all subsequent exposures to lower Δ₆ were limited to 4 min. The results of these measurements are presented in Fig. 2b. Indeed, the μ_cm values decrease as Δ₆ is lowered in the reactor and trace back the preceding rise almost exactly but for a small hysteresis.

Discussion

An atomistic picture of the nucleation, growth, and coalescence process of graphene is now discussed to help analyze these mobility results and the ones to be discussed in the next section. It is shown that the reversible phenomena is due to grain boundary thermodynamics and not owing to growth kinetics. From a growth perspective, it is important to establish this fact, as thermodynamic control is more robust than a kinetic one. The atomistic picture is shown schematically in Fig. 3a. We use the potential of carbon atoms at various locations in the growth environment to first qualitatively explain the observed phenomena. We later use density functional theory (DFT) calculations to support these conclusions. The graphene growth reaction starts by setting a carbon potential in the vapor phase (C_v) in the form of the methane partial pressure. Methane decomposes on the catalyst and yields a surface carbon potential (C_s) crystalline graphene domains with a potential C_s (carbon potential within the graphene grain) nucleate and grow by consuming the surface carbon. C_s at any point on the growth surface will vary with time and will be determined by the balance between carbon addition by adsorption and decomposition and carbon removal by desorption, nucleation, and attachment to growing edges. With an increase in coverage, as the domains get closer to form boundaries, the C_s values will get lowered due to reduced catalyst site availability and increased consumption. The reduced C_s will lower lateral growth rates and in the extreme case prevent complete coalescence in reasonable time periods. In addition to these three potentials, carbon atoms in the defect structures such as GB,

**Fig. 2** Reversibility in mobility with reversal in thermodynamic conditions. a Effect of annealing sample S₁ under increasing supersaturations (Δ₆) on mobility (μ_cm) and Raman I_D/I_G ratio. μ_cm is seen to increase significantly with Δ₆ and then saturate. The averaged defect density as measured by Raman is insensitive to this change. The sample S₂ (Δ₆ = 82 kJ mol⁻¹) is indicated. b Reversal of μ_cm on reversal of supersaturation, indicating that the phenomenon is thermodynamic in origin and not kinetic. The small hysteresis can be attributed to kinetics. The sheet resistance vs supersaturation and resistance vs gate bias plots are included in the Supplementary Note 6. Samples S₂ and S₃ are indicated.
triple junction and line defects in the graphene monolayer will be at higher carbon potentials \( (C_s) \) than \( C_g \) and are shown in Fig. 3a as a band.

If \( C_s \) became equal to \( C_g \) during growth, a coverage saturation will happen despite a global supersaturation \( (C_s \) being higher than \( C_g) \). If \( C_s \) were raised by increasing \( C_v \) (increasing \( \Delta G \)), the degree of coverage will increase in a given time of growth. Such growth is termed kinetically controlled and the coverage saturation owing to kinetic limitations has been described before by Celibi et al. using a Gompertzian model\(^{24}\). Thus, the increase in mobility with \( \Delta G \) or methane partial pressures in Fig. 2a can be attributed to “more faster and complete coverage” in the time frames involved, in the sense that the grain boundary closure is more complete. If this were true, that is, the improvements in mobility in Fig. 2a were purely owing to kinetic reasons as just discussed, then re-annealing the samples that had attained the highest mobilities in lower methane partial pressures \( (C_g) \) still higher than \( C_d \) should not reverse the mobility gains as observed in Fig. 2b. The reversal is thus proof of the improvements, which happen post grain coalescence, having a thermodynamic origin.

Thermodynamically, the ability to form a particular boundary structure that will complete the coalescence process, will require the global carbon potential, \( C_v \), to be greater than \( C_g \) as shown in Fig. 3b. If in spite of this scenario it did not form, then the effect is kinetic. Such a situation will arise if \( C_d \) determined by the relative rates of adsorption–attachment and desorption were to become lesser than \( C_v \). On the other hand, if the situation shown in Fig. 3b prevailed, then, thermodynamic constraints will come into play. Those boundary structures that have \( C_d > C_v \) will never form as it will not be energetically favorable for carbon from the vapor phase to become part of these defect structures. Increasing/decreasing \( C_v \) will then add/remove carbon from these structures, thus healing/forming defects, vacant carbon sites, reversibly at these locations. The spatial and temporal variation of \( C_g \) is difficult to estimate quantitatively. Hence, in the next section we compare the energetics of \( C_s \), with that of boundary energies to determine if such an effect is possible.

Dong et al.\(^{15}\) have shown that by changing the hydrogen potential in the reactor, one can replace covalently bonded boundaries by hydrogen terminated ones and vice-versa. The experimental work presented here involves changing the carbon potential in the reactor resulting in reversible exchange of carbon atoms between the boundary and the vapor phase. To model this process (See Supplementary Note 8 for the thermochemistry considerations) density functional simulations were carried out to compare the energetic differences between different states of the grain and its boundaries. The thermodynamic reversibility at the grain boundary implies that there are two states, a grain boundary and a defective grain boundary, between which the system can toggle depending on the carbon potential in the system. The defective grain boundary will have to be formed by the breaking of carbon–carbon bonds. It is expected that the resulting dangling bonds will get passivated by hydrogen in the hydrocarbon environment of the reactor. This resultant entity can be approximated by a graphene edge terminated by hydrogen. The reaction that represents this process can be expressed as:

\[
C_v - C_s - C + 3H_2 \leftrightarrow 2C - H + CH_4
\]  

(3)

“C′” represents grain boundary carbon and the equation shows a defect removal and its replacement by an edge. In order to determine whether an edge state with lower energy can exist, two graphene grain boundary configurations and their corresponding edges, Fig. 4a, b, were chosen. DFT calculations were performed (see details in Supplementary Note 8) to determine the energies of the grain boundary and the edge. Data from JANAF tables.
were used to calculate the potentials of hydrogen from $\text{H}_2$ and carbon from $\text{CH}_4$. These results were then used to calculate the free energy change for the reaction in equation (3) ($\Delta G = G_{\text{Reactants}} - G_{\text{Products}}$). The $\Delta G$ variation with methane partial pressure is plotted in Fig. 4c.

A positive value means that the reaction can proceed in the forward direction, i.e., an edge can replace a boundary as it has lower energy. A negative value means that the boundary will be stable against such breakup and defect creation. The plot shows that indeed, a pressure range exists, in which the edge has lower energy than the boundary. There is a crossover after which the reverse is true. The crossover pressure is dependent on the grain boundary configuration. Thus, if at the temperature of graphene growth, there are boundary configurations with positive $\Delta G$, then the phenomenon observed will occur at these sites. This will then result in the thermodynamic reversibility observed.

In the previous discussion, we had hypothesized that vacant carbon sites can be annihilated and created reversibly at GB causing the mobility reversal with change in vapor phase carbon potential. First principles calculations supported this idea. In this section we provide experimental proof. A simple experiment was first performed in which graphene samples with decreasing grain sizes, obtained by increasing supersaturation in the nucleation stage itself (see Supplementary Note 9), were exposed to 5 min of pure hydrogen at a pressure of 400 Torr and 1000 °C. The resulting change in the sheet resistance before and after hydrogen treatment is shown in Fig. 5. It is clearly seen that under the same hydrogen treatment, which is expected to create defects in the graphene film, samples grown under higher $\Delta G$ and hence smaller grain sizes (or larger grain boundary length per unit area) not only have a larger sheet resistance but also the highest increase on hydrogen annealing. This indicates that the defects are predominantly created at the GB.

In all the four samples previously discussed, S1–S4, it was noted that while $\mu_{\text{cm}}$ showed variations, Raman defect density remained mostly unaltered. As Raman defect density is an average value obtained from a large area scan (10 $\mu$m × 10 $\mu$m), the changes should be happening at length scales that the mapped average does not reflect. To understand the physical picture behind the reversible defect density-mobility trend, a method involving preferential water permeation through defects in graphene followed by etching of an underlying Ge film was employed on the four samples listed in Table 1. This process creates a replica of the defect structure in the graphene monolayer on the underlying Ge film and renders them optically visible. A detailed time sequence of the etching to be discussed in the following section is presented in the Supplementary Note 10. Optical images are presented in Fig. 6a–h and AFM images in Fig. 6i–l. Figure 6a, e show that when the VCG (sample S1) is examined by this method, the etch pattern formed within just 1 min of permeation, involving 50–100 $\mu$m sized grains, corresponds to the microstructure of the Cu substrate on which the monolayer was synthesized (see Fig. 1b). Thus, most of the major defects in VCG lie at the Cu GB. In contrast, in S2, S3, and S4, samples that have been annealed in methane post VCG, the Cu GB are not revealed even after 60 min of permeation (Fig. 6b–d, f–h). Higher magnification images, second row, show that the etch pattern has an average grain size of 6.7 $\mu$m (standard deviation of 0.8 $\mu$m), which matches well with the values measured using SEM (see Supplementary Note 2). Thus, in these samples the defective regions are predominantly at the coalescence boundaries in the graphene layer.

In addition, in the reverse annealed sample S6, the etch pattern lines are thicker (Fig. 6c, g) compared to the annealed sample S2 (Fig. 6b, f). The AFM scans shown in Fig. 6i–l reveal that difference in the surface after permeation experiments in sample S2 and S6. Larger etch pit widths quantitatively correspond to higher rates of permeation enabled by more defective boundaries. After 100 min etch, the wider pit depth of 474 nm in the reverse annealed sample S6, when compared to 56 nm in S2, clearly shows that reverse annealing had indeed made the GB of the monolayer more defective. Further details on the AFM scans are shown in the Supplementary Note 11. Thus, reverse annealing at lower $\Delta G$ has made the graphene GB more defective. One can argue that the reverse annealing might also cause defects within the grain (HRTEM imaging in the next section throws more light on this aspect) that are not detected by the permeability measurements. To determine if this is so, sample S4 was annealed in pure hydrogen at a pressure of 4 Torr. Even in this case as can be seen from Fig. 6d, h only GB are revealed. If the grain were to have become defective, then one will see more uniform etching of the Ge below. Thus, these results are direct evidence for our thesis that the reversible rise and fall in $\mu_{\text{cm}}$ in Fig. 2b, is owing to a decrease/increase in grain boundary defects with a rise/fall in $\Delta G$. The type of defects (structure) and their density along the GB of different samples were inspected using a high resolution TEM. Imaging in a 200 kV TEM was first performed over large areas to establish that the graphene being studied was indeed predominantly a monolayer using tilt diffraction measurements (see Supplementary Note 12). The samples were then observed with an aberration corrected scanning transmission electron microscope (STEM) under a 60 kV electron beam. A large pristine region of the graphene film in S1 is shown in Fig. 6m. The figure shows that the bulk of graphene is mostly defect free with negligible point or line defect density and that the imaging process does not create any noticeable damage to the lattice. Figure 6n shows a representative grain boundary in S3 and Fig. 6o shows a representative boundary in S4. While the boundary in Fig. 6n shows a string of Stone–Wales defect structures distributed in a region less than 1 nm, that in Fig. 6o shows a region that is at least 2 nm wide. The grains in Fig. 6n have a misorientation angle of 13.8° and the defective atomic columns consist of continuous Stone–Wales (5–7) structures (see Supplementary Note 13 and 14 for details on angular misorientation calculations and hexagon to Stone–Wales ratio calculations). The ratio of these
structures to those of the standard hexagon is > 1. Theoretical predictions based on geometric considerations (Supplementary Note 14) predict a ratio of < 0.6 for a misorientation of ~ 13.8°. Thus, the higher defect density at the boundary in S1, the 6 min VCG, is indicative of a graphene that has not been fully formed. The triple junction that is close to the imaged boundary can also influence the defect density. However, what this simple calculation nevertheless shows is that defect densities at boundaries and especially closer to triple junctions will be much larger than those theoretically anticipated. They in turn can have a significant impact on the properties of the monolayer.

On the other hand, Fig. 6o shows a boundary, which is both wider and higher in defect density than the one in Fig. 6n. Voids are seen at the boundary in addition to standard Stone–Wales defect sites clearly showing the effect of annealing at lower ΔG. This explains the lower μcm (see Supplementary Table 1) and high water permeability in S2. Only covalently bonded GB were observed at the supersaturations used in the work done as part of this research as can be noted from the SEM and STEM images. Overlapped boundaries as reported by other groups were not observed under the conditions used for synthesizing the samples used in this paper (Supplementary Note 3). However, at much lower methane partial pressures and lower growth rates, overlapped GB were also observed. A discussion of the characteristics of those samples is beyond the scope of this paper.

In summary, we show that the configuration of grain boundary defects in graphene monolayers have a thermodynamic relationship with carbon potential in the vapor phase. These defects can be created and healed reversibly. This relationship allows one to control grain boundary structure. The resulting large variation in grain boundary resistance allows the field effect mobility in graphene monolayers to be changed by more than an order of magnitude, from 1000 cm² V⁻¹ s⁻¹ to 20,000 cm² V⁻¹ s⁻¹ and
sheet resistance from 1200 to 159 Ω□−1, the best reported yet for such large device sizes. Most of these microstructural changes that result in such large variations in electrical properties cannot be detected by standard Raman and SEM characterization. The study also shows that GB might be equally, if not more important, than grain size in graphene.

Methods
CVD graphene growth. Graphene was grown in a custom-made CVD reactor22,27. Ultra-pure (99.9999%) hydrogen, and nitrogen flowing through a purifier, 99.98% pure methane and 99.98% pure Cu from Sigma Aldrich were used for synthesis. Although growth was done at 1000 °C, reactor chemistry was changed to vary carbon potential by varying methane and hydrogen flows. Gas flows for the supersaturation used in this article are shown in Supplementary Fig. 3. Total reactor pressure was maintained at 4 Torr for all our experiments. Such a condition result in such large variations in electrical properties cannot be explained by variations in the growth conditions24,25, were conducted using an Olympus made immersion lens etch measurements and AFM

Raman measurements. Raman spectroscopic measurements on these transferred layers were carried out with a LabRAM HR spectrometer fitted with a 532 nm laser source. The instrument was fitted with precision micrometer stage for mapped measurements. Further details on Raman spectroscopic measurements are provided in Supplementary Note 4.

Electrical measurements. Van der Pauw devices for sheet resistance measurements and MOS FET structures for mobility measurements were fabricated using standard optical lithographic techniques and the graphene channel was defined using a mild oxygen plasma etch using an Oxford reactive ion etcher. Except for electrical measurements below 4 min in Fig. 11 all the electrical measurements were made on 100 μm × 100 μm area as shown in Supplementary Note 5. The measurement statistics and fitting examples are also provided in Supplementary Note 6. For the samples grown for less than 4 min, the graphene film was not continuous and hence, electron beam lithographically made devices were used. Details are provided in Supplementary Note 7. All electrical measurements were made after annealing the sample under a current density of 3 × 10−6 Acm−2 for 3 min under a pressure of 4 × 10−5 mbar of air. Given that the average grain size is ~7.08 μm (standard deviation of 0.88 μm), (See Supplementary Fig. 2) the 100 μm channel widths are expected to yield average values that represent electronic properties over large areas.

Etch measurements and AFM. Ge etch experiments to characterize defects, developed in our group24, were conducted using an Olympus made immersion lens setup fitted with an Olympus optical acquisition system at room temperature and in open air. The etch profile measurements25 were performed using a Bruker made atomic force microscope.

TEM. Diffraction measurements were performed using a FEI T20 transmission electron microscope at 200 kV and the STEM measurements were made using an aberration corrected Nion Ultra STEM 100 microscope operated at 60 kV.

Data availability
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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Author contributions
TB and MJ conducted the first-principle studies, RS, RM, and PG performed the high-resolution TEM studies. KB fabricated the device for water permeation studies, and SS and MV conducted the permeation measurements. AM fabricated the devices for electrical studies, and KB performed the electrical transport studies. KB, RP, and SR designed the experiments and wrote the manuscript. All the authors contributed for discussions and analysis leading to the article.

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