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Synthesis of graphene and graphene nanostructures by ion implantation and pulsed laser annealing

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In this paper, we report a systematic study that shows how the numerous processing parameters associated with ion implantation (II) and pulsed laser annealing (PLA) can be manipulated to control the quantity and quality of graphene (G), few-layer graphene (FLG), and other carbon nanostructures selectively synthesized in crystalline SiC (c-SiC). Controlled implantations of Si− plus C− and Au+ ions in c-SiC showed that both the thickness of the amorphous layer formed by ion damage and the doping effect of the implanted Au enhance the formation of G and FLG during PLA. The relative contributions of the amorphous and doping effects were studied separately, and thermal simulation calculations were used to estimate surface temperatures and to help understand the phase changes occurring during PLA. In addition to the amorphous layer thickness and catalytic doping effects, other enhancement effects were found to depend on other ion species, the annealing environment, PLA fluence and number of pulses, and even laser frequency. Optimum II and PLA conditions are identified and possible mechanisms for selective synthesis of G, FLG, and carbon nanostructures are discussed. Published by AIP Publishing. [http://dx.doi.org/10.1063/1.4955137]

INTRODUCTION

In previous experiments, we demonstrated that ion implantation (II) and pulsed laser annealing (PLA) could be applied to selectively synthesize graphene (G) and few-layer graphene (FLG) on crystalline silicon carbide (c-SiC), only where ions were implanted, with nanoscale precision and control, and without altering the surrounding or underlying c-SiC.1,2 The motivation for these initial experiments was to develop a processing approach using multi-ion beam lithography3 combined with thermal or PLA to write nanoscale G features directly onto insulating c-SiC in a manner that avoided using many of the conventional lithographic processing techniques (chemical etching, laydown/lift-off, transfer, etc.). Since II and PLA are processing approaches widely used in current device manufacturing, we considered this a promising approach for some G nanoscale device and sensor applications.

While this processing approach was shown to be quite successful, it left a number of unresolved questions concerning the exact mechanisms controlling the synthesis of the G and FLG. Additional experiments since have shown that the origins of these ambiguities are closely allied with the advantageous strengths of the II and PLA processing techniques. Namely, II and PLA are inherently non-equilibrium processing techniques (ion damage, doping, cascade quenching, etc.; laser rapid heating/melting, resolidification, etc.); each has multiple processing control features (implanted ion species, dose/dose-rate, energy/depth-profiles, laser fluence, pulse duration, number of pulses, etc.); and the processing environment also affects results.

In this paper, we report on controlled experiments that explore how the multiple II and PLA processing parameters can be combined and controlled to selectively synthesize graphene nano-structures; identify how these parameters can be optimized to achieve desired results; and discuss the mechanisms responsible for synthesis of G, FLG, and other graphene nano-structures.

EXPERIMENTAL METHODS

Single crystalline 4H and 6H SiC wafers were purchased from II-VI, Inc. Nanoscale patterning of graphene features utilized a multi-ion beam lithography, nanofabrication, and engineering (MionLiNE) system at the University of Florida (UF).3 For some experiments, it was desirable to do a variety of experiments on a single SiC sample to eliminate potential sample differences, and in these cases, broad area implantations were done at the Australian National University (ANU).

Pulsed laser annealing was performed at UF with a 193 nm wavelength, 25 ns pulse-width JPSA IX-260 ArF laser. The laser has a double fly’s eye beam homogenizer system to produce spatially uniform irradiations. A mask system can control laser exposure areas such as 25 μm × 25 μm, and a precision X-Y positioning table and microscope system can...
be programmed to position exposure areas with a resolution of \( \pm 1 \mu m \) and stage movement repeatability of \( \pm 0.1 \mu m \). Laser fluence, number of pulses, and frequency can be varied over a wide range.

Sample analyses included Raman spectra using a Horiba MicroRaman Spectrometer with a 532 nm green laser, Auger electron spectroscopy (AES), X-ray diffraction, scanning electron microscopy (SEM), thermal simulation calculations, and ion scattering/channeling performed at ANU. Complementary high-resolution cross-sectional transmission electron microscopy (HR-XTEM) was performed at UF to directly image the near-surface regions of the SiC wafers using a JEOL 2010F transmission electron microscope operated at 200 kV. Samples were prepared using an FEI DB235 dual scanning electron microscope/focused ion beam (FIB) system with methods described elsewhere.4–7 Prior to performing FIB cross-section milling, the specimens were coated with \( \sim 100 \) nm of Cr, and occasionally Pt, to protect the surface of the specimens;8 this Cr layer was not present during PLA.

RESULTS AND DISCUSSION

While our initial experiments1,2 successfully demonstrated that II and PLA could be combined to selectively synthesized G and FLG on SiC, and with nanoscale precision, the results raised a number of questions concerning the contributing mechanisms and the optimum processing parameters. For example, laser fluences of 1.0–1.2 J/cm\(^2\) were required to initiate graphitization of unimplanted c-SiC. But when other c-SiC samples were implanted with a variety of different ions [Si (self-dopant), Ge (isoelectronic), Au (noble/catalytic), and Cu (catalytic)], the laser threshold-fluence for the onset of graphitization, \( \Phi_G \), varied widely—\( \Phi_G \) (Cu) \( \sim 0.1 \) J/cm\(^2\), \( \Phi_G \) (Au) \( \sim 0.3 \) J/cm\(^2\), \( \Phi_G \) (Ge) \( \sim 0.5 \) J/cm\(^2\). It is well known that ion implantation doses as low as \( \sim 10^{14}–10^{15} \) ions/cm\(^2\) can damage c-SiC and can render the near-surface material to an amorphous state.9 Previous research has shown that the threshold for melting the surface of a-SiC is strongly dependent on the thickness of the amorphous SiC layer.10–12 So, it was unclear whether the strong dependence we observed for the \( \Phi_G \) threshold on implanted ion species was due to the amorphous surface layer produced by the ion implantation process, or to a catalytic effect due to the implanted ion species (doping), or both. To investigate further, we devised experiments to study these effects separately.

The effects of amorphous SiC surface layers

Amorphous layers of various thicknesses were produced on the (000-1) (C-face) of 6H-SiC single crystalline wafers by ion implantation of Si\(^-\) and C\(^-\) negative ions at projected ranges and fluences to produce stoichiometric distributions. This approach produces a relatively “pure” a-SiC layer. Stopping and Range of Ions in Matter (SRIM)\(^{13}\) calculations were used to estimate the ion energies and fluences needed to produce various thickness a-SiC layers, and the experimental conditions are shown in Table I. Ion scattering/channeling measurements were used to measure the resulting amorphous layer thicknesses, and these were confirmed by HR-XTEM as illustrated in Figure 1. The ion implantations and ion beam analyses were performed at ANU and HR-XTEM at UF. Ion beam analysis was done with 2.0 MeV He\(^+\), scattering angle 110°, and 20° exit angle relative to the surface.

This approach of using stoichiometric implantation produces a uniform a-SiC layer as Figure 1(b) indicates for the 20 nm thick layer (sample 1). Using samples 1–5, PLA parameters could be optimized over the range of thicknesses from 20–217 nm for the “pure” amorphous layers and compared to results when dopant atoms are implanted into these

<table>
<thead>
<tr>
<th>Sample number</th>
<th>Si E (keV) ((10^{15} \text{Si}^-/\text{cm}^2))</th>
<th>C E (keV) ((10^{15} \text{C}^-/\text{cm}^2))</th>
<th>Amorphous layer (d_a) (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>10</td>
<td>1.5</td>
<td>1.75</td>
</tr>
<tr>
<td>2</td>
<td>40</td>
<td>1.5</td>
<td>1.5</td>
</tr>
<tr>
<td>3</td>
<td>60</td>
<td>1.5</td>
<td>27</td>
</tr>
<tr>
<td>4</td>
<td>80</td>
<td>1.5</td>
<td>37</td>
</tr>
<tr>
<td>5</td>
<td>160</td>
<td>1.8</td>
<td>75</td>
</tr>
</tbody>
</table>

FIG. 1. (a) Representative ion scattering channeling measurements for various thickness a-SiC layers on c-SiC and (b) HR-XTEM of 20 nm thick a-SiC on c-SiC.
layers. But, first, it is useful to estimate how amorphous layer thicknesses affect surface temperatures for our laser annealing conditions.

**Thermal simulations**

Early studies by Baeri et al.

and related numerical simulations were conducted by Dutto et al.

and Hedler et al.

We performed similar calculations for our different experimental conditions. A finite element analysis was used to estimate the surface temperature during the PLA process. We used an unsteady-state energy balance equation with rectangular coordinates (x-, y-, and z-axes) to approximate the surface temperatures of ion-implanted samples after exposure to one 25 ns laser pulse as shown by

\[
k \left[ \frac{\partial^2 T}{\partial x^2} + \frac{\partial^2 T}{\partial y^2} + \frac{\partial^2 T}{\partial z^2} \right] + \alpha I(z, t) = \rho C_p \frac{\partial T}{\partial t},
\]

where \( T \) is temperature, \( \rho \) is density, \( C_p \) is heat capacity, \( \alpha \) is absorption coefficient, \( k \) is thermal conductivity (in units of \( \text{W/cm K} \)), \( z \) is distance from the sample surface, \( t \) is time, and \( I \) is absorbed power during laser exposure. This can be represented as a function of \( z \) and \( t \) by using

\[
I(z, t) = I_0(t)(1 - R) \exp(-az),
\]

where \( I_0 \) is the incident laser intensity at the sample surface \((z = 0)\) and \( R \) is the reflectivity of the specimen.

Assuming the SiC surface melts within the depth of laser’s penetration depth \( dp \), the heat that is required for SiC to transform from solid phase to liquid phase could be represented as a function of mass and latent heat of fusion

\[
Q = mL,
\]

where \( L \) is the latent heat of fusion and \( m \) is the mass of SiC surface with a depth of laser’s penetration depth, \( dp = 1/\alpha \). \( m \) is presented as

\[
m = \rho A dp = \rho A/\alpha,
\]

where \( \rho \) is density, \( A \) is area, \( dp \) is laser’s penetration depth, and \( \alpha \) is absorption coefficient. Therefore, when \( A \) is 1 cm\(^2\) and the penetration depth \( dp \) is 9.1 nm, this latent heat of fusion \( Q \) is 3.14 \( \times 10^{-3} \) J, which means it requires a laser fluence of 3.14 \( \times 10^{-3} \) J/cm\(^2\) for solid to liquid SiC phase change.

Figure 2 shows the laser fluence required to melt a given thickness of a-SiC on c-SiC, based on our thermal simulation results. We used 2445 K as the a-SiC melting temperature, and the remaining thermal properties of a-SiC and c-SiC are listed in Table II. The trend of decreasing fluence with increasing thickness is due to the poor thermal conductivity of a-SiC compared to c-SiC. When the laser energy is absorbed in the amorphous layer, the heat is conducted relatively slowly until it reaches the c-SiC substrate where is conducted away much more rapidly. Consequently, the thinner the amorphous layer, the more efficiently the heat is conducted away and so higher laser fluences are required to melt thin amorphous layer than thick ones. Based on the latent heat of fusion calculation, the fluence that required for solid to liquid SiC surface phase change is much smaller than the fluence required to heat the entire sample from room temperature to melting temperature. Therefore, the latent heat of fusion could be negligible during the thermal simulation. It should also be noted that the simulation results are only applicable for the initial pulse. As will be shown, depending on fluence, the first pulse can melt or change the thickness of the amorphous layer; transform it to polycrystalline SiC, G, or other carbon nanostructures, etc.; and such changes can present different thermal properties for subsequent pulses. However, these calculations still constitute a useful guide for estimating the temperature range during PLA. Fortunately, HR-XTEM can confirm whether the sample surface melts or changes phase.

**Raman assessment after PLA a-SiC**

The results of PLA on the various thicknesses of a-SiC layers \( (dp) \) with a variety of laser processing parameters were studied using Raman spectroscopy and HR-XTEM. Figure 3 compares Raman spectra for samples 1–3 and pristine c-SiC. The peaks at 1350 cm\(^{-1}\), 1580 cm\(^{-1}\), and 2700 cm\(^{-1}\) are

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Amorphous SiC</th>
<th>Crystalline 6H-SiC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Density, ( \rho ) (g/cm(^3))</td>
<td>2.66</td>
<td>3.21</td>
</tr>
<tr>
<td>Specific heat, ( C_p ) (J/g K)</td>
<td>1.3</td>
<td>1.3</td>
</tr>
<tr>
<td>Thermal conductivity, ( \kappa ) (W/cm K)</td>
<td>0.011</td>
<td>4.9</td>
</tr>
<tr>
<td>Absorption coefficient, ( \alpha ) (10(^6)/cm)</td>
<td>1.1</td>
<td>1.5</td>
</tr>
<tr>
<td>Reflectivity, ( R ) (%)</td>
<td>34.6</td>
<td>40</td>
</tr>
<tr>
<td>Melting point, ( T_m ) (K)</td>
<td>2445</td>
<td>3100</td>
</tr>
<tr>
<td>Latent heat of fusion, ( L ) (J/g)</td>
<td>1300</td>
<td>...</td>
</tr>
</tbody>
</table>
referred to as D, G, and 2D peaks characteristic of graphene, respectively. In general, the G peak indicates sp² bonding (in plane stretching of C-C bonds), 2D represents a greater degree of sp² bonding order (such as G, FLG, and carbon nanostructures), and the D peak represents disorder in the system. As \( d_a \) for samples 1–3 increases, the relative intensities of the D, G, and 2D peaks increase. The trends in the Raman signals due to graphitic carbon can be isolated by normalizing and subtracting the c-SiC signal at 1520 cm\(^{-1}\) and comparing the intensity of the graphitic peaks (\( I_G, I_{2D}, I_D \)). If this is done, the intensity of each spectrum’s D peak (\( I_D \)) is comparable to the intensity of its G peak (\( I_G \)), indicating a significant amount of disorder in all samples. The high degree of disorder also leads to the appearance of the D peak at 1620 cm\(^{-1}\) and the D + G peak at 2930 cm\(^{-1}\).

**HR-XTEM assessment after PLA a-SiC**

To evaluate the structural modifications resulting from PLA, HR-XTEM was performed on both c-SiC and the amorphized samples in Table I. Figure 4 compares PLA results for a 6H c-SiC sample to sample 1 with a 22 nm thick stoichiometric amorphous surface layer. Both were annealed in 1 atmosphere of Ar with 2000, 25 ns, 0.8 J/cm\(^2\) and 50 Hz. At this fluence, both are below the melt threshold for one pulse. After 2000 pulses, Figure 4(a) shows no evidence of change to the c-SiC surface, but Figure 4(b) shows that the amorphous surface layer has decomposed forming 4–5 G layers on the crystalline substrate and an amorphous layer above containing both Si and C atoms. While 2000 pulses at 0.8 J/cm\(^2\) do not alter the 6H crystal, the rapid heating and cooling reach higher temperatures that are sufficient to free and drive Si atoms toward the surface, leaving C atoms behind to bond and form FLG on the c-SiC surface.

Figure 5 shows HR-XTEM results for sample 4 (\( d_a = 124 \) nm) where PLA at 100 pulses at 0.8 J/cm\(^2\) is above the melt threshold. The results in Figure 5(a) are consistent with the interpretation that the amorphous surface layer melts next to the 6H c-SiC substrate, and repeated pulses cause rapid cooling near the crystalline substrate forming polycrystalline 3C-SiC, and an amorphous region above where carbon nanostructures formed. Figure 5(b) shows that FLG formed at the sample surface; 5(c) shows that carbon nano-onions formed in the middle of the amorphous layer; and 5(d) shows the formation of graphitic nanostructures forming in the light regions within the 3C poly that are trapped amorphous regions.

Carbon nano-and-micro-structures like those we have observed, sometimes referred to as onions because of their shapes, were first observed by Iijima, after heating carbon black rods to extremely high temperatures (>4000 K). Other synthesis methods have since been recognized, including high energy electron irradiation of carbon soot, arc discharge in water from a carbon target, and carbon ion implantation in Cu substrates. All of these methods involve non-equilibrium conditions such as rapid heating of carbon-containing precursors to high temperatures or ion implantation where ion induced damage cascades quench rapidly. The carbon-onions observed in sample 4 most likely form due to the rapid melting and re-solidification of a- or c-SiC that occurs during our 25 ns PLA. In our case, repeated pulses can deplete the amorphous areas of Si leaving excess C atoms to bond with each other in 3-dimensional space into spherical and multi-shell arrangements. However, the HR-XTEM images indicate that the formation of FLG mainly stabilizes at interfaces such as the sample surface or amorphous/crystalline interfaces, or at the amorphous surface. Therefore, a thinner amorphous layer enhances G and FLG formation and limits the possibility of carbon-onions growth.

**FIG. 3.** Raman spectrum for pristine SiC, PLA pristine SiC, sample #1, sample #2, and sample #3.

**FIG. 4.** PLA in 1 atm. Ar with 2000, 25 ns, 0.8 J/cm\(^2\), ArF pulses at 50 Hz. (a) 6H c-SiC and (b) c-SiC with a 22 nm thick stoichiometric II amorphous surface layer 22 nm thick. (Below the “melt” threshold for both a-or-c-SiC).
Catalytic or doping effects; Au implantation

We began these experiments to help differentiate the role of the amorphous surface layers from the ion species used to create the amorphous layers during PLA. The previous results and discussion helped clarify how an amorphous surface layer on c-SiC can affect PLA. Now, we want to see if there is a doping or catalytic effect. In order to isolate any “catalytic effect” of implanted species, two samples were prepared from the same crystal by implanting equal amounts of Si$^+$ (1.5 $\times$ 10$^{15}$ ions/cm$^2$, 10 keV) and C$^-$ (1.5 $\times$ 10$^{15}$ ions/cm$^2$, 4 keV) into 6H-SiC to form 20 nm-thick amorphous SiC surface layers. The first sample was used as a control, and the second sample was additionally implanted with Au$^+$ (5 $\times$ 10$^{15}$ ions/cm$^2$, 30 keV) into the amorphous region. The thickness of the amorphous layer was measured by ion scattering/channeling at ANU and verified as 20 nm by HR-XTEM at the University of Florida [see Figure 1(b)].

When combined with the “amorphous effect,” this offers additional control for utilizing II and PLA to selectively form G on SiC.

Optimizing PLA parameters

The prior experiments and discussions have shown that applying PLA to selectively synthesize G and G-like structures in SiC is affected by both the amorphous surface layers produced by II, and by the species of ion used. In this section, the effects of fluence, number of pulses, frequency of pulse delivery, and sample environment during PLA are studied to further identify optimum conditions.

PLA melting effects; high fluences

To supplement the understanding of the amorphous surface layer and doping effects, PLA of c-SiC was also studied.
PLA without II has also been studied by Lee et al.,24 Hwang et al.,25 and Hattori et al.26 using KrF pulsed lasers (~1.2 J/cm², 25 ns). To evaluate how melting affected the structure and graphene growth for our conditions, HR-XTEM and Raman were used to study PLA fluences from 0.1–1.0 J/cm² at 50 Hz, on 4H c-SiC, both above and below Φm. Thermal simulations were performed to estimate the surface peak temperature of c-SiC from a single pulse at different laser fluences ranging from 0.1 to 1.0 J/cm² as shown in Figure 7. The fluence that melts the surface is defined as Φm, the surface reaches the melting temperature, and c-SiC at the top surface changes from solid to liquid, the laser light reflection increases, and absorption decreases in the liquid state. However, the practical reflectivity during melting is not easily available and was estimated as shown by the red dashed line.

The HR-XTEM images of the SiC crystal were taken after 1, 10, and 100-pulse laser pulses and are compared in Figure 8. For clarity, Figures 8(a), 8(c), and 8(e) are intentionally aligned, so the surface is at the same level. In Figure 8(a), 1 laser pulse at 1 J/cm² melted the c-SiC at the near surface, and recrystallization yields a ~20 nm thick polycrystalline 3C-SiC on the 4H c-SiC substrate, and a thin amorphous layer and few layers of graphene on top. Figure 8(b) magnifies the surface area in Figure 8(a) and shows that 2–3 layers of graphene have formed on top of the surface, but the quality is not the best and these FLG samples appear to be discontinuous. The corresponding Raman spectra showed only tiny G and 2D signals corresponding to sp² bonding and a strong SiC signal, indicating that the original crystal structure was still largely intact. Figure 8(c) is the TEM image after 10 laser annealing pulses at 1 J/cm². Apparently, the amorphous layer increases to around 60 nm, and the polycrystalline 3C-SiC seems still around 20 nm but much more non-uniform. The magnified TEM image in Figure 8(d) shows that many more layers of graphene formed at the interface of the polycrystalline SiC and amorphous layer. The Raman spectra showed that the SiC peak was greatly reduced, and the G, D, and 2D peaks rose sharply signifying increased graphene layer formation. Figure 8(e) is the TEM image after 100 pulses. The total thickness affected by the laser is ~75 nm, which increased slightly compared with Figure 8(c). Interestingly, more and more SiC recrystallizes from the amorphous phase to polycrystalline phase, and it seems the amorphization and the recrystallization are in competition. During this process, some of the inclusions show carbon nano-structures like carbon anions in Figure 8(f).

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**PLA environment effects**

Experiments exploring the effects of the annealing environment showed that PLA in 1 atm of Ar, H₂, or N₂ ambient can significantly lower the graphene yield compared with PLA in vacuums as low as 1–20 mTorr. Figure 9 compares Raman spectra for PLA of c-SiC in Ar ambient and vacuum. Figure 9(a) shows results for PLA with 100 pulses at 0.8 J/cm². The red (Ar) and green (vacuum) spectra have similar G peaks, but slightly different D and 2D peaks. The green spectrum (vacuum) has a narrower D peak and higher 2D peak, implying slightly better graphene quality. Annealing with more laser pulses, such as 500 pulses, helps differentiate the two ambient conditions as shown in Figure 9(b) where the G, D, and 2D peaks rise sharply signifying increased graphene quality. The existence of Ar atoms in the environment creates a pressure that could slow or suppress the free silicon atoms from leaving the surface, and therefore lowers the graphene yield. The situation is similar to the growth of graphene by thermal annealing c-SiC at high temperatures.21

**PLA frequency effects**

The results from Section “PLA Melting Effects” indicated that PLA at high fluences that melt the surface result in relatively poor quality graphene. In experiments to create higher quality G, both lower fluences and pulse delivery times (frequency effects) were explored. It was found that PLA of c-SiC at fluences below 0.8 J/cm² produced no change to the SiC crystals and did not produce G regardless of frequency. But, in PLA experiments at 0.8 J/cm² in vacuum, an unusual effect between the laser repetition rate and graphene yield was discovered. Figure 10 contrasts Raman spectra of SiC after pulsed laser annealing at 50 Hz and 1 Hz. At 0.8 J/cm², the peak

![Graph showing thermal simulations of peak temperatures at the end of 25 ns laser pulses at various fluences](image)

**FIG. 7.** Thermal simulations of peak temperatures at the end of 25 ns laser pulses at various fluences. The solid black line region is from the thermal simulation, and red dashed line indicates the change to liquid has occurred.
surface temperature is \(~2200\) K and is much lower than the c-SiC melting temperature of \(~3100\) K according to the thermal simulation. Figure 10(a) is the Raman spectrum of PLA at 0.8 J/cm\(^2\), with multiple pulses at 50 Hz in vacuum. With increasing number of pulses, the G and 2D peaks do not change significantly and only the D peak appears slightly, but it is still considerably smaller compared with pristine SiC. Figure 10(b) shows the Raman spectrum of PLA SiC at 0.8 J/cm\(^2\), with multiple pulses at 1 Hz in vacuum. The areas that were annealed with 125 pulses or more demonstrate prominent graphitic signals. So, this “frequency effect” indicates that lower repetition rate results in improved graphene growth.

To confirm the graphene growth and quality, HR-XTEM images were taken for these samples. Figure 11 compares the TEM images of PLA at 50 Hz (a) and 1 Hz (b) with 375 pulses at 0.8 J/cm\(^2\). Figure 11(a) shows little and discontinuous graphene growth, which is consistent with Raman spectra results for the same conditions. To the contrary, Figure 11(b) shows 4 layers of graphene epitaxial growth on the c-SiC surface, and there is also a thin amorphous layer on top containing unconnected loose graphene features. When Raman spectra signal was acquired, the signal from both the epitaxial graphene and the loose graphene on top was collected. And, we believe that the defect peak, D, in Figure 10(b) is primarily contributed from the loose graphene features in the amorphous region, and the epitaxial graphene on the c-SiC substrate has very good quality based on these TEM images.

CONCLUDING REMARKS

We have shown that ion implantation and pulsed laser annealing can be controlled to synthesize graphene and graphene nanostructures in SiC. While we were aware that these combined techniques offered a wide range of processing parameters, some of the resulting effects were surprising. The observation that the a-SiC phase anneals/melts at lower fluence than c-SiC is expected and occurs for other materials like Si. The enhanced graphene formation associated with the doping effect can also be rationalized. But, the exact mechanism whereby SiC is depleted of Si to allow graphene formation during rapid heating is not certain. Lee et al.\(^{24}\) calculated that the amount of Si that would be lost to sublimation—even assuming melting with 500, 25 ns, KrF laser pulses—was only \(~10^{-2}\) atoms/A\(^2\), and hypothesized a photophysical bond breaking effect. But, some of the other effects we observed such as the frequency, environment, nanostructure formation, and structural changes are not totally consistent with this interpretation.

The mechanism of the Au\(^{+}\) “catalytic effect” for graphitization is still under investigation. Au and Cu metals have
been used in the CVD method for graphene growth,\textsuperscript{20–22} and both exhibit a catalytic effect. The low solubility of carbon in these transition metals leads to precipitation of carbon atoms at the surface that form G when they cool down; this is a self-limiting process. Silicon and gold can form Au-Si eutectic alloys at a much lower melting temperature\textsuperscript{23} and may help remove Si atoms in a Si-C system. However, our concentration of implanted Au is small, and our estimates are that it is unlikely this low concentration of Au is enough to change the absorption characteristics, and thus surface temperature, of the Au\textsuperscript{+} implanted sample compared to the control sample, or to “change” the structure of the amorphous layer. The explanation we propose for graphene growth during PLA of Au\textsuperscript{+} implanted SiC is that the laser heats the amorphous surface to a high temperature, silicon atoms start to de-bond from carbon atoms, and diffuse toward the surface where they escape. The presence of the Au either enhances de-bonding or loss of Si, leaving the surface enriched with C atoms that bond to form graphene structures. However, it is still unverified how the Au could increase the loss rate, or how interactions between Au/Si and Au/C encourage graphene growth. In our study, as a result of Au\textsuperscript{+} implantation, more layers of graphene formed after laser annealing, and we confirmed that Au has a “catalytic effect.” Combined with the “amorphous effect,” this means that we can better control the laser annealing process and selectively graphitize the SiC surface by II and PLA.

It is also not clear how the “frequency effect” phenomenon enhances graphene growth in PLA SiC. The temperature change during PLA increases rapidly during the 25 ns laser pulse, and then quickly drops back to room temperature in a few $\mu$s, based on the thermal simulation.\textsuperscript{27} The only difference between these two conditions is the time interval between two subsequent laser pulses, which is 20 ms for

![FIG. 9. Raman spectra of PLA c-SiC in Ar ambient and vacuum condition.](image)

![FIG. 10. Raman Spectrum for frequency effect. (a) Pulsed laser annealing c-SiC in vacuum, at 0.8 J/cm\textsuperscript{2}, at 50 Hz with 1, 5, 25, 125, 375, 625, and 1000 pulses. (b) Pulsed laser annealing c-SiC in vacuum, at 0.8 J/cm\textsuperscript{2}, at 1 Hz with 1, 5, 25, 125, 375, 625, and 1000 pulses.](image)

![FIG. 11. HR-XTEM images (a) PLA 375 pulses at 0.8 J/cm\textsuperscript{2}, 50 Hz. (b) PLA 375 pulses at 0.8 J/cm\textsuperscript{2}, 1 Hz.](image)
50 Hz and 1000 ms (1 s) for 1 Hz. Even 20 ms is long enough for the surface to be cooled down to room temperature compared with a few μs, but it seems that a longer wait time at room temperature is helping the graphene growth process. In the physical process of PLA c-SiC, silicon atoms are given enough kinetic energy due to the temperature to de-bond with carbon atoms and then can desorb from the SiC surface. After silicon atoms leave the surface, the carbon atoms could bond with each other and form graphene. If the free silicon atoms do not leave or do not leave quickly enough, silicon atoms could inhibit the graphene growth by re-bonding with carbon atoms. In terms of time scales, the de-bonding and bonding process in the chemical reaction is in the range of a few nanoseconds to microseconds, while the surface is still hot, but the mass transportation is in the millisecond or second range when the surface is cool. It seems that the graphene growth during PLA c-SiC may not be reaction controlled but a diffusion controlled process.

We have shown that the numerous processing parameters of II and PLA can be controlled and manipulated to produce the amount and quality of G, FLG, and carbon nanostructures on SiC. And, both are compatible with current device manufacturing methods, so they can have significant advantages for growing and patterning G on SiC.

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