On the Significance of a Carbon-Rich Background in Plasma-Based Graphene Oxide Reduction

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The reduction of oxygen concentration in graphene oxide is demonstrated using electron beam generated plasmas produced in three different gas backgrounds: Ar, Ar/H₂ and Ar/CH₄. Plasma diagnostics and surface characterizations are combined to determine the influence of each working gas on the chemical composition and structure of graphene oxide before and after reduction. The results suggest that argon treatment alone allows the removal of weakly bound oxygen in graphene oxide by inert ion bombardment, while the addition of hydrogen adds reactive species that enhance the removal of oxygen. However, without a carbon source, removing oxygen from graphene oxide can lead to the formation of defects and vacancies. We find that methane provides not only hydrogen but also the carbon necessary to restore the graphitic plane by healing defects generated during reduction.
1. Introduction
Graphene, the two-dimensional sp² bonded carbon sheet, has attracted much attention for its outstanding properties. One of the routes to large-area synthesis of graphene, or at least graphene-like material, is by reducing the oxygen concentration in graphene oxide (GO). GO is produced through a derivative of the well-known oxidation method of graphite described by Hummers and Offeman [1], which yields a solution of GO flakes that can be readily spin cast over large, arbitrary substrates. Fully oxidized GO is an insulator. However, it has been shown that by controlling the oxygen concentration, one can tune its electrical conductivity [2-4]. Different techniques can reduce graphene oxide including wet approaches using for example, solutions of hydrazine hydrate [5] or dry processes like heating [6,7] and plasma techniques [8-10]. Plasmas are the tool of choice across a broad range of industries given their ability synthesize and modify materials over large areas with nanoscale precision. Surface modification is of particular interest for making physical and chemical changes in the top few layers of materials while maintaining the properties of the bulk. When plasma processing atomically thin films like GO, considerable care must be taken to prevent any etching of the material. For such films, electron beam generated plasmas are ideal due to their high plasma densities and inherently low electron temperatures, which results in a large flux of low energy ions at substrate surfaces. Typical ion kinetic energies do not exceed a few eV [11,12] which is insufficient energy to sputter surface atoms but enough to promote surface diffusion and stimulate desorption [13].

In previous work [10], X-ray photoelectron spectroscopy (XPS) results demonstrated the ability to cleanly and controllably reduce the oxygen concentration in GO from 45 at.% to 5 at.% using electron-beam generated plasmas produced in argon/methane (Ar/CH₄) backgrounds. However, unlike other reduction approaches [14] the process was found to increase the sp² cluster size. The present work clearly delineates the role of a carbon-rich background gas in preserving the graphitic plane from defects during the reduction process by comparing the results using electron beam generated plasmas produced in Ar, Ar/H₂ and Ar/CH₄ to reduce GO. The treated material was characterized to follow the chemical and structural changes using surface energy measurements as well as X-ray photoelectron and Raman spectroscopies. To draw further distinctions between the various background gases, we measure plasma densities and relative ion fluxes to adjacent surfaces in each background. The results provide additional insight and support the general argument that a carbon-rich working gas, in this case methane, delivers carbon to the surface and is thus a better choice for reduction in terms of the physico-chemical characteristics of the reduced graphene oxide.

2. Experimental
The experimental detail of the plasma processing system and the reduction protocol has been described elsewhere [10]. Briefly, the system employs a sheet-like, pulsed, high-energy (≈ 2 keV) electron beam that ionizes and excites the background gas to produce plasma. The electron beam is collimated with a 150 Gauss magnetic field to provide a well-defined and localized ionization volume.

The graphene oxide films were produced by the Hummers method [1] and dispersed in water by sonication. The solution was then spin cast on SiO₂ (300 nm)/Si substrates and

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dried in nitrogen. The samples were then placed on a stage located 2.5 cm from the electron beam axis and the system was evacuated to a base pressure of $<10^{-5}$ Torr. Plasma processing was performed at 90 mTorr pressure, which was achieved by introducing the gases through mass flow controllers and by throttling the turbo-molecular pump. Argon was the main constituent of the process environment with reactive gases ($H_2$ or $CH_4$) set to 5% of the total gas flow. The samples were treated for 5 minutes at a 10% duty factor (2 ms pulse width and a period of 20 ms), which is equivalent to 30 seconds of plasma exposure.

After plasma treatment, the samples were characterized and compared using Raman spectroscopy and X-ray photoelectron spectroscopy (XPS). The contact angles of three liquids—deionized distilled water, ethylene glycol and diiodomethane—were measured to determine the components of the surface energy following the method of Owens and Wendt [15], where the relation between the contact angle and the surface energy is described by:

$$\gamma_L(1 + \cos \theta) = 2 \left( \sqrt{\gamma_S^d \gamma_L^d} + \sqrt{\gamma_S^p \gamma_L^p} \right),$$

with

$$\gamma_S = \gamma_S^d + \gamma_S^p,$$

where $\theta$ is the measured equilibrium contact angle of the liquid on the surface, $\gamma_S$ is the surface energy of the surface to be measured, $\gamma_L$ is the surface tension of the used liquid, and the superscripts $d$ and $p$ indicate the dispersive and polar components of the surface and liquid energy or tension, respectively. Thus, knowing the surface tensions of the three liquids, and their polar and dispersive components, the measured contact angles can be used to derive the total surface energy and its polar and dispersive components. For this work, the average contact angle of three to five drops was used.

To understand the relative production of species and their transport to surfaces in the different gas backgrounds, measurements of plasma densities and relative ion fluxes at an adjacent surface were measured in a similar system [16]. Although the system is different than the one used for processing, the salient physics is the same. That is to say, the relative production of species via high-energy electron beams will depend largely on three parameters: beam energy, operating pressure, and relative gas concentration [12]. These three parameters were nominally the same in each system. For these measurements a 1 cm diameter, high-energy (2 keV) electron beam was injected into Ar, Ar/$H_2$ and Ar/$CH_4$ gas mixtures. The total flow was 100 sccm and the pumping speed was throttled using a manual gate valve to set the pressure at 90 mTorr. For the mixtures, the $H_2$ and $CH_4$ flow rates were set to 5% of the total flow (5 sccm) with the remainder being argon. Langmuir probes were used to measure the electron density as a function of distance from the beam axis. The ion fluxes were measured using an energy-resolved mass spectrometer (Hiden Analytical) located approximately 2.5 cm from the beam axis. Details of these measurement techniques can be found in [17].

3. Results and discussion
The differences in species production in the gas backgrounds used for processing can be seen in Figures 1 and 2. From Figure 1, the on-axis (position = 0 cm) electron density is
found to decrease when molecular gases are added and the density drops more rapidly with increasing distance from the beam. This is not surprising since the primary loss mechanism is diffusion to the wall in beam generated plasmas produced in noble gas backgrounds; electron-ion recombination (e.g. \( \text{H}_2^+ + e^- \rightarrow 2\text{H} \)) in the bulk plasma adds an additional loss channel when molecular gases are introduced [18].

From Figure 2, the ion mass spectrum in pure argon is dominated by \( \text{Ar}^+ \) ions. The \( \text{ArH}^+ \) and water ions are indicative of residual water present in the reactor. When \( \text{H}_2 \) is added to the background, the \( \text{Ar}^+ \) ion intensity drops significantly, a large \( \text{H}_3\text{O}^+ \) signal emerges, and traces amounts of \( \text{H}_3^+ \) are observed. Molecular gases (in this case \( \text{H}_2 \)) readily undergo charge exchange reactions with noble gas ions (e.g. \( \text{Ar}^+ + \text{H}_2 \rightarrow \text{ArH}^+ + \text{H} \)), with other molecular gases (e.g. \( \text{H}_2^+ + \text{H}_2\text{O} \rightarrow \text{H}_3\text{O}^+ + \text{H} \)), or themselves (\( \text{H}_2^+ + \text{H}_2 \rightarrow \text{H}_3^+ + \text{H} \)). Considering these charge exchange reactions, the aforementioned electron ion recombination reaction, and direct dissociation of \( \text{H}_2 \) from high-energy beam electrons implies that, despite the relatively low \( \text{H}_2 \) neutral density, a significant amount ionic \( \text{H} \)-containing species (as well as reactive neutrals) will be produced and delivered to the surface. When \( \text{H}_2 \) is replaced by \( \text{CH}_4 \), a significant number of \( \text{C}_n\text{H}_{y}^+ \) ion species are observed. While the myriad of reactions leading to the measured spectrum are rich in gas-phase chemistry and certainly worthy of
continued investigations, the relevant result to this work is that there are an abundance of carbon-containing ions and presumably neutral radicals that are created and delivered to the graphene oxide surface.

The relevant characteristics of graphene oxide, GO reduced in Ar-produced plasmas (Ar-rGO), GO reduced in Ar/H₂-produced plasmas (H₂-rGO) and GO reduced in Ar/CH₄-produced plasmas (CH₄-rGO) are compared in Table 1. The surface energy is clearly modified after plasma treatment and is dependent on the working gas. Generally, the addition of oxygen to a surface will increase the surface energy, transforming the surface from hydrophobic to hydrophilic and thus increasing reactivity. The plasma treatment does not change the dispersive component of the surface energy. This is expected since the plasma treatment changes the concentration of the polar functional groups containing oxygen on the graphene surface. Ar plasma treatment has little effect on the polar component. However, the addition of reactive gases leads to substantial reduction in the polar component and thus the total surface energy. The CH₄-containing plasma produces the largest decrease, bringing the total surface energy to 45.5 mJ/m². These results are similar to those found for hydrazine-reduced graphene oxide, where the value was decreased from 62.1 mJ/m² to 46.7 mJ/m² [19]. The XPS survey results in Table 1 verify that the reduction in surface energy is linked to the decrease in the oxygen concentration. A slight decrease of the oxygen concentration was achieved using argon plasmas while the reduction of oxygen using reactive gas-containing plasmas was substantial.

Table 1: Surface energy calculated from the contact angle measurements of three liquids on untreated and plasma-treated graphene oxide and the corresponding oxygen concentration and oxygen-to-carbon ratio.

<table>
<thead>
<tr>
<th>Quantity</th>
<th>GO</th>
<th>rGO Ar plasma</th>
<th>rGO Ar/H₂ plasma</th>
<th>rGO Ar/CH₄ plasma</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface Energy (mJ/m²)</td>
<td>25</td>
<td>24</td>
<td>16</td>
<td>7.5</td>
</tr>
<tr>
<td>Polar</td>
<td>36</td>
<td>35</td>
<td>36</td>
<td>38</td>
</tr>
<tr>
<td>Dispersive</td>
<td>61</td>
<td>59</td>
<td>52</td>
<td>45.5</td>
</tr>
<tr>
<td>Total</td>
<td>43</td>
<td>36</td>
<td>30</td>
<td>10</td>
</tr>
<tr>
<td>O concentration (at.%)</td>
<td>0.75</td>
<td>0.56</td>
<td>0.43</td>
<td>0.1</td>
</tr>
</tbody>
</table>

The results of the oxygen reduction are best detailed by looking at the C1s high resolution XPS spectra shown in Figure 3. Fitting results show the presence of C-C, C-O, C=O and O-C=O bonds at 284.4 eV, 286.5 eV, 287.6 eV and 288.8 eV, respectively. Their concentrations are shown in Figure 4. For all processing conditions, the C-C concentration clearly increases and is accompanied by the decrease of C-O concentrations. These species can be located either on the basal plane or along the edges of the GO flakes [21]. The contributions from the O-C=O and the C=O peaks, attributed to species mainly located at the GO edges [20], changed only slightly when using pure Ar and Ar/H₂ plasmas. However, the use of CH₄ clearly enhances their reduction. It is important to note that the deposition of amorphous carbon using methane has been ruled out for the treatment conditions in this work [10].
Following our earlier observation that sp\(^2\) cluster size increased when reducing GO in Ar/CH\(_4\) plasmas, [10] we measured the D to G ratios using Raman spectroscopy to calculate the sp\(^2\) cluster size (La) using the empirical formula: 
\[
La (\text{nm}) = \frac{10^3}{227. \frac{I_D}{I_G}}
\]

of Tuinstra and Koenig [21]. Those results are presented in Figure 5. In contrast to the results using Ar/CH\(_4\) plasmas, the D to G ratio, and consequently the sp\(^2\) cluster size decreased or remained constant for GO treated in Ar and Ar/H\(_2\) plasmas.

The results indicate that argon plasmas can reduce the oxygen content. That is, a flux of non-reactive, low energy ions are sufficient to remove weakly bound oxygen (C-O). The addition of charged and neutral hydrogen species enhances the reduction. This is not surprising given the addition of H can produce volatile species like H\(_2\)O or OH, thus making the removal of oxygen easier. It is well known that the defect sites of graphene are more reactive than the carbon in the plane [22], and so the interaction of the species takes place mainly on the edges and at defect sites. However, the removal of oxygen in itself can create a defect, either by leaving a dangling bond or through the simultaneous removal of carbon via, for example,
the desorption of CO or CO$_2$; those defects could be passivated by the uptake of oxygen upon exposure to air.

When the plasma is produced in the presence of methane instead of hydrogen, hydrogen species are still produced and delivered to the surface. In addition, C$_x$H$_y$ species are present and provides carbon that can heal vacancies created by the removal of oxygen. This concept is consistent with noticeable reduction in carboxyl and carbonyl groups (Figure 4) and an increase in the sp$^2$ cluster size of Ar/CH$_4$-rGO over both Ar-rGO and Ar/H$_2$-rGO (Figure 5).

4. Summary
The final oxygen content and surface structure of GO when using electron beam produced plasmas to reduce the oxygen content is strongly dependent on the type of background gas. Argon plasmas provide an inert flux of low energy ions capable of removing a small amount of oxygen, while the addition of hydrogen can stimulate a further reduction of oxygen content via the delivery of hydrogen-containing species. The addition of methane will stimulate further reduction. But, unlike hydrogen plasmas, those generated in methane backgrounds produced a wide range of carbon- and hydrogen-containing species, where the additional carbon could fill vacancies created by the removal of oxygen.

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References

Figure 5 D to G intensity ratio and sp$^2$ cluster size of GO and r-GO plasma treated in Ar, Ar/H$_2$ and Ar/CH$_4$ gas mixtures.


