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Low pressure RF plasma modification of the surface of three different nano-carbon materials

Abstract: Well-ordered nano-carbon materials, like multi-wall carbon nanotubes, graphene oxide, graphene due to their unique physical and chemical properties, are candidates for promising applications.

In this work thin multilayered graphene, single layer graphene oxide layers and highly oriented pyrolytic graphite (HOPG) surface were treated by RF activated N\(_2\) gas plasma at nominally room temperature. Negative bias in the 0–200 V range and treatment time of 10 min was applied. Surface chemical alterations were followed by X-ray photoelectron spectroscopy (XPS). The applied treatments resulted in a significant build-up of nitrogen in the surface of these nano-carbon materials. The amount of nitrogen varied between 4 and 10 atomic %, depending on type of carbon and on biasing conditions. Evaluating the high-resolution N1s XP spectral region, typically three different chemical bonding states of the nitrogen were delineated. Peak component at 398.3 eV is assigned to C=–N–C type, at 399.7 eV to sp\(^2\) N in melamine-type ring structure and at 400.9 eV to N substituting carbon in a graphite-like environment. Identical chemical bonding of the nitrogen was detected on the surface of HOPG treated in the same way for comparison.

Keywords: RF plasma, surface modification, nano-carbon materials, XPS

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1 Introduction

Well-ordered nano-carbon materials, among others, multi-wall carbon nanotubes, graphene oxide, graphene due to their unique physical and chemical properties, are candidates for promising application in various areas ranging from novel structural materials, field emission devices to pharmaceutical drug-delivery vehicles [1-4]. Many of these applications require some kind of surface modification of these materials. Moreover, surface groups may significantly influence the wettability, acid/base properties or even the shape of the carbon nanoparticles [5-6]. Concerning drug carrier application, reactive surface groups are needed for attachment of bioactive molecules. When using them for reinforcing in polymeric matrices, not only the dispersion, but also strong covalent attachment to the matrix is required [7].

Recently, we have investigated application of low-pressure radio frequency (RF) plasma activation of N\(_2\) and to explore this approach to the covalent attachment of the nitrogen to various carbon materials [8]. In this work thin films of graphene oxide, multilayered graphene and highly oriented pyrolytic graphite, were treated by RF activated N\(_2\) gas plasma at nominally room temperature and the amount of built-in nitrogen and its chemical states were studied by X-ray photoelectron spectroscopy (XPS).

2 Experimental

2.1 Samples

Thin layers of powdered graphene (GR) obtained by disintegration [9] of synthetic graphite (Aldrich, USA) and commercial single layer graphene oxide (SLGO) flakes (Cheaptubes, USA), were deposited dropwise from ultrasonicated alcoholic slurry onto a stainless steel holder (Ø 9 mm), suitable also for XPS measurements. Pieces of freshly cleaved highly oriented pyrolytic graphite (HOPG) samples of similar size were used as reference materials.
2.2 Treatments

Plasma treatments were performed in the stainless steel sample preparation chamber of the XPS instrument (base pressure $<1 \times 10^{-4}$ Pa). $N_2$ flow regulated by a bleeding valve to set the pressure to 5 Pa. Constant RF power of 100 W at 13.56 MHz was applied through a matching circuit to a stainless steel coil built into the preparation chamber. The sample bias was varied between 0 and -200 V. The treatment time was always 10 min.

2.3 Characterisation

The plasma treated samples were characterised by X-ray photoelectron spectroscopy using a Kratos XSAM 800 spectrometer (Mg $K_{\alpha}$, 1253.6 eV excitation, fixed analyzer transmission mode). Low resolution survey spectra were recorded in the 150–1300 eV kinetic energy range with 0.5 eV steps. High resolution spectra of the O1s, N1s and C1s were acquired by 0.1 eV steps and 1 s dwell time. For the determination of the chemical shifts of the hetero-atoms (N and O), each set of spectra was carefully referenced to the C1s line of the sp$^2$ type graphitic carbon, set at 284.1 ± 0.2 eV BE (binding energy). Quantitative analysis, based on peak area intensities after removal of the Shirley-type background, was performed by the Kratos Vision 2 and the XPS MultiQuant 7 programs [10,11] (http://aki.ttfk.mta.hu/XMQpages/XMQhome.htm). Experimentally determined photo-ionization cross-section data of Evans et al. [12] and asymmetry parameters of Reilman et al. [13] were used for the calculation. The chemical states of the implanted nitrogen were evaluated by applying the peak decomposition procedure by using the Kratos Vision 2 software.

3 Results and discussion

3.1 Chemical composition

The overall surface compositions (calculated by the generally applied “infinitely thick homogeneous sample” model) of the treated samples are summarized in Figs. 1 and 2. The data show that 4-10 atomic % nitrogen can be incorporated into the sample surface at the plasma treatment even at the applied relatively short 10 min reaction time. Somewhat higher N content (~14 atomic %) was measured for the GR sample at 20 min exposure with 200 V bias. The nitrogen attachment to the samples shows significant bias dependence. At low bias (50 V), only a small amount of N could be incorporated into HOPG while similar amount surface N content was found for all samples at 200 V bias. As concerning the bias dependence,
it is interesting to note that the 50 V negative bias does not affect increase of N content above the values obtained at zero bias for GR and SLGO. A very low (1-2 atomic %) oxygen content was measured both on untreated HOPG and on graphene. After all plasma treatments it rises up to 4-6 atomic % and it does not show significant bias dependence. The possible source of the oxygen contamination may be the $\text{O}_2$ and $\text{H}_2\text{O}$ impurity of the reagent gas or desorption of these species from the chamber walls.

### 3.2 Chemical Structure

The incorporation of nitrogen into these nano-carbons will inevitably modify the chemical structure of the ordered graphene type carbon network.

The C1s, N1s and O1s spectra for the HOPG, GR and SLGO samples recorded at different biases (0-200 V) are depicted in Figs. 3-5. The chemical states of the measured elements in the HOPG and GR samples are similar. During the plasma treatment, only their intensities are decreasing, demonstrating that the implanted nitrogen (and oxygen) incorporated into the first 10 nm layer. The C1s spectrum of the SLGO sample is significantly different, because of the extra peaks of the carbon bonded to the large amount of oxygen. Both the elemental and oxygen bonded carbon content decreases with the increasing bias.

The chemical states of oxygen do not vary significantly for the treatments with different biases, while their content changes slightly. The O1s line of oxygen contamination, related to the presence of $\text{O}_2$ and $\text{H}_2\text{O}$, for the HOPG and GR is centred at 532.0 eV. This shift can be most probably...
assigned to C–OH chemical states [14]. The high BE sub-
peak at about 533.5 eV indicates the presence of ketone
and carboxyl groups [14,15].

XP spectra of the N1s energy region for the GR and
SLGO and also for HOPG samples, treated at 100 V bias,
are depicted in Fig. 6. The broad peaks clearly indicate
that the bonded nitrogen atoms have different chemical
environments. Performing peak synthesis fitting with
1.75 eV half-widths components, corresponding to
individual chemical states, three N1s components
denoted further as N1, N2 and N3) were found in all
treated samples, positioned at 398.3 ± 0.3 eV , 399.7 ± 0.3 eV
and 400.9 ± 0.3 eV , corresponding to minimum three
different chemical environments. According to Fig. 5,
the same chemical states of nitrogen were detected also
on the surface of SLGO, although their relative amount
was somewhat different. The assignment of this peak
components to certain bonding states were discussed in a
large number of publications based on experimental works
devoted to the synthesis of CNx type materials [16-23],
N doped graphene [24,25] and nitrogen-containing DLC
coatings [8,26,27]. The controversy existing in rendering
N-C structures to different chemical shifts was discussed
in [16-19,28]. The determined chemical shifts were
assigned in accordance with published data as follows: N1
peak component at 398.3 eV was assigned to C=N–C type
chemical bonds, i.e. to the sp2 nitrogen bonded to two sp2
type carbon, like in pyridine [21,22,27]. The N2 component
at 399.7 eV is attributed to sp2N in diazine or triazine-
type structure [28]. The smallest N3 type N at 400.9 eV
we assigned to N bonded to three graphitic rings, i.e.,
substituting a carbon atom in graphite-like environment
[27]. These assignments were summarised in Table 1.

The ratio of the different chemical states of nitrogen
does not seem to alter with the applied bias. As an
exception, the treatment of SLGO sample where at 200 V
bias the relatively higher abundance of the N1 type nitrogen
is obvious.

Figure 5: C1s, N1s and O1s lines of the SLGO samples treated at different bias (untreated black, 0 V blue, 50 V red, 100 V green, 200 V magenta) (colour online).

Figure 6: Decomposition of the N1s lines of the HOPG (a), GR (b) and SLGO (c) samples treated at 100 V bias.
Table 1: Assignment of N1s components of the plasma treated nano-carbon materials.

<table>
<thead>
<tr>
<th>Component</th>
<th>Energy (eV)</th>
<th>Assignment</th>
</tr>
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<tbody>
<tr>
<td>N1</td>
<td>398.3 ± 0.3</td>
<td>sp² N in pyridine type ring</td>
</tr>
<tr>
<td>N2</td>
<td>399.7 ± 0.3</td>
<td>sp² N in diazine/triazine type ring structure</td>
</tr>
<tr>
<td>N3</td>
<td>400.9 ± 0.3</td>
<td>N substituting C in graphite plane</td>
</tr>
</tbody>
</table>

4 Conclusions

It was demonstrated that three different carbon materials (GR, SLGO, HOPG) could be conveniently modified using low-pressure RF plasma activated nitrogen. 4-10 atomic % of nitrogen could be incorporated into the surface at short 10 min treatment time. The amount of nitrogen can be varied by the negative sample bias of 0-200 V. The implanted nitrogen (and the oxygen impurity) is incorporated in the first few atomic layers of the samples.

Identical chemical bonds of the nitrogen were detected on the surface of all these samples. The three different chemical bonding states of the nitrogen was delineated and assigned to C=N–C (imine, pyridine) type (N1), to sp² N in melamine-type ring structure (N2) and to N substituting carbon in a graphite-like environment (N3). The relative abundance of these three types of N does not vary significantly with the applied bias.

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References


