Electrochemical exfoliation-streamline method for synthesis of nitrogen doped graphene

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Graphene was discovered in the early 21st century, but has already proven itself in many applications – energy, medicine, electronics, food and sports, and more. Functionalization of nanostructured carbon materials with both non-metallic and metallic atoms is possible in various ways, imparting enhanced or new properties to the starting material, even catalytic activity. A method of electrochemical exfoliation was used to obtain the graphene sheets and simultaneously functionalize them with nitrogen. To ensure N-doping the process is done in a NaN₃ electrolyte solution which provides less quantity of oxygen groups that tend to block defect sites on the graphene, compared with such solvents as NaNO₂. Two graphite electrodes are inserted into the electrolyte and a pulse power of 0–10V is applied. The solution containing the obtained material is filtered through a 0.1 µm filter and dried. The material is characterized using SEM, XRD and XPS. In the XPS characterization graphene oxide is used as a reference material.

Keywords: nitrogen-doped graphene, sodium azide, sodium nitrite, electrochemical exfoliation

INTRODUCTION

The goal of the paper is to explain the electrochemical exfoliation process for the synthesis of N-functionalized graphene as well as the characterization of the acquired material.

Beyond its unique set of physico-chemical properties, graphene can be considered as a robust, atomic scale scaffold from which other 2D materials can be derived through the replacement/attachment of foreign atoms and functional groups [1–3]. Among the possible doping agents, nitrogen (N) has drawn a considerable amount of attention because its atomic radius is comparable to that of carbon and contains five valence electrons available to form strong covalent bonds. The conjugation between the nitrogen lone-pair electrons and the graphene π-system modifies graphene physical and chemical properties. The substitution of carbon atoms by nitrogen ones influences the atomic charge distribution on the graphene scaffold and creates ‘active sites’ thus significantly increasing the electrochemical activity of nitrogen-doped graphene, known as N-graphene (NG) [4].

Graphene, as a 2D carbon nanomaterial in which sp2 hybridized carbon atoms are aligned in
a honeycomb lattice, is a relatively new material with a wide range of applications due to its chemical stability, electric and thermal conductivity and large surface area [3–7]. Recent studies reveal that substitutional doping of graphene with heteroatoms N can tailor its electrical conductivity, wettability and electro-active surface area to facilitate charge-transfer and electrolyte–electrode interactions, and may give rise to its electrochemical performance [7, 8].

As mentioned before, the functionalization of graphene improves upon its base qualities. In the case of nitrogen doping resolve at least three types of C–N bonds (defect sites) can form during the functionalization process [7, 9]: quaternary N (or graphitic N), pyridinic N and pyrrolic N (Fig. 1) [10]. Usually, pyridinic N bonds with two C atoms at the edges or defects of graphene contribute one p electron to the π system. Pyrrolic N refers to N atoms that contribute two p electrons to the π system, although unnecessarily bond into the five-membered ring, as in pyrrole. Quaternary N is the N atoms that substitute for C atoms in a hexagonal ring. In these nitrogen types, pyridinic N and quaternary N are sp2 hybridized and pyrrolic-N is sp3 hybridized [2, 3, 6, 10].

Nitrogen doped graphene can further be used in batteries, sensors, supercapacitors, fuel cells and more [2, 6, 7].

**EXPERIMENTAL**

Commercially available chemicals have been used. High-purity graphite rods (Alfa Aesar, 99.9995%), commercially available chemicals (analytical grade)-NaN₃, NaNO₂, HNO₃ and 2-propanol were used as supplied without additional purification. Distilled water was used for electrolyte preparation and washing.

Electrochemical exfoliation of graphite was carried out in a 800 ml beaker using two high-purity graphite rods (HOPG) and a potentiostat (212, ISSP) situated as shown in Fig. 2. The synthesis procedure was carried out in a pulse mode of electrolysis-polarization of electrode by +10 and 0 V (both throughout 30 s) with multiple changing of polarization potential. The synthesis process was carried out in 27 h maximum time. One molar aqueous solution of NaN₃ and NaNO₂ was used as electrolyte with different procedures and results. In the case of NaNO₂ about 75 ml of HNO₃ was added at the beginning of the process. The obtained dispersion of the graphene material was stirred using ultrasound treatment in a UP200St ultrasonic processor (hielsher) for 1 min and then filtered through series of filters with different pore sizes (0.45, 0.2 and 0.1 µm). During filtration the material was rinsed thoroughly with distilled water and dried in an oven at 50°C.

For XPS measurements an ESCALAB Xi, Thermo Fisher spectroscope was used. For SEM images a Helios 5 UX, Thermo Fisher Scientific microscope was used. For XRD measurements a MiniFlex 600, Rigaku spectroscope was used.
The dry material was then wrapped in filter paper and washed with a mixture of water and 2-propanol using a Soxhlet extractor therefore getting rid of the unwanted leftover synthesis solution. After letting the process run for 15 h, the material was dried and, if needed, transferred to the appropriate solvent and dispersed using ultrasound treatment for 1 min.

RESULTS AND DISCUSSION

During the exfoliation process a change in the colour of the electrolyte from colourless to grey, then to black, could be observed, indicating that graphene sheets were created during the process [3].

During the electrolysis process azide decomposes into Na⁺ and N₃⁻ ions as well as sodium nitrite decomposes into Na⁺ and NO₂⁻ ions. The N₃⁻ ion changes further: N₃⁻ → 3/2N₂+e⁻ and in water N₃⁻+3H₂O+2e⁻ → N₂+NH₃+3OH⁻. The nitrite ion reacts with water creating nitric acid and nitric oxide: NO₂⁻+H₂O → HNO₃+NO and in further reactions N₂ and NH₃ are created. The N₃⁻ and NO₂⁻ ions intercalate in the graphite layers, and when they decompose, they simultaneously split the graphite layers and functionalize the graphene [3].

Figures 3, 4 and 5 show the X-ray photoelectron spectroscopy (XPS) measurements of graphene.
oxide GO synthesized by a modified Hummer’s method \([11]\) and NG from electrochemical exfoliation in NaN\(_3\) and NaNO\(_2\), respectively.

XPS is the standard quantitative spectroscopy technique to study the nitrogen doping effect in graphene. In the XPS spectrum of N-graphene, at about 400 and 284 eV the peaks appear that resemble N1s and C1s, respectively. The ratio of peak intensity between N1s and C1s is employed to determine the nitrogen content in N-graphene.

In Fig. 4 we can examine GO to determine that graphene has indeed been produced considering Fig. 3.

![Fig. 4. XPS spectra of NG synthesized using NaN\(_3\)](image)

![Fig. 5. XPS spectra of NG synthesized using NaNO\(_2\)](image)
that graphitic carbon (Csp2 hybridized) can be viewed (284–285 eV) in all of the scans [2, 8, 12].

The N1s spectrum is used to determine the nitrogen configurations. Usually the N1s spectrum can be deconvoluted to several individual peaks that are allocated to different C–N bonds: pyridinic-N (400 eV), pyrrolic-N (401–402 eV), quaternary-N (402–403 eV) and N-oxides of pyridinic N (408–410 eV) [4–6, 8, 9, 12, 13]. As a consequence of the sample being charged, the readings could change, as we see in Fig. 4, where all of the peaks have moved.

Figure 6 shows the X-ray spectroscopy (XRD) measurements of the solution after the electrochemical exfoliation in NaN₃. We found that the solution contained mainly NaHCO₃ [14] and we can assume that all NaN₃ has dissociated. We can assume that the C⁴⁺ ion presence tells us that the C–C bonds in graphene have been split and the lack of nitrogen in the solution tells us that there is a big likelihood that C–N bonds have been created.

Figure 7 shows the XRD measurements of graphene that was synthesized using NaNO₂ electrolyte solution. We can see a peak at around 26–28 degrees and that is consistent with the peak of nitrogen-doped graphene [5, 6, 12, 13].

Figure 8 is a scanning electron microscope (SEM) image taken of nitrogen doped graphene.

**Fig. 6.** XRD spectra of NaN₃ solution after synthesis

**Fig. 7.** XRD spectra of NG synthesized in NaNO₂

![Graph](image-url)
that has been synthesized using NaN$_3$. We can see the single layer of graphene that has been formed (a) which confirms that the graphite has been exfoliated into graphene.

Furthermore, we see small particles of graphene at the size of 10 nm and even barely visible particles as small as 1 nm (b), in the image they can be seen as small white spots on a bigger graphene piece.

Figure 8 is a SEM image taken of nitrogen doped graphene that has been synthesized using NaN$_2$. In the image (a) we see 50 nm graphene sheets that have not come apart fully. In the image (b) we see a similar situation, only the sheets are smaller at around 3 nm if we look at the smallest sheets.

In conclusion, the process and results of electrochemical preparation of multilayer graphene functionalized with nitrogen (N-MLG) via exfoliation of graphite electrodes in the presence of azide and nitrite anions in a pulse mode of electrolysis have been proven. Sodium azide and nitrite provide not only exfoliation of graphite into graphene but also simultaneously functionalize the graphene with nitrogen ions as a part of azide and nitrite ion decomposition. The X-ray diffractometer analysis confirmed that all of the nitrogen had left the solution after synthesis and C$^{+4}$ ion was present which indicated that the nitrogen probably had intercalated into the graphene sheets. The X-ray photoelectron spectroscopy confirmed the nitrogen groups in the material and GO was used as a reference material. The scanning electron microscope images confirmed that few-layer graphene was acquired as well even though it could not have been
filtered due to its small size. Further studies are needed to improve upon the synthesis method.

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Fig. 9. SEM images of NG synthesized in NaNO₂: (a) sheet stack at a size ~50 nm, (b) sheet stacks at a size of ~3 nm

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Santrauka
Elektrocheminės eksfoliacijos būdas panaudotas sintezuojant grafeno lakštus ir kartu dopuojant juos azotu. Darbe aprašomos šios sintezės sąlygos, užtikrinančios mažesnio kiekio deguonies turinčių funkcinių grupių susidarymą. Susintetintos medžiagos charakterizuotos skenuojančios elektroninės mikroskopijos, rentgeno spindulių difrakcijos ir fotoelektroninės spektroskopijos metodais.