Chemical and structural properties of reduced graphene oxide—dependence on the reducing agent

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ABSTRACT

Graphene oxide (GO) prepared from graphite powder using a modified Hummers method and reduced graphene oxide (rGO) obtained from GO using different reductants, i.e., sodium borohydride, hydrazine, formaldehyde, sodium hydroxide and L-ascorbic acid, were investigated using transmission electron microscopy, X-ray diffraction, Raman, infrared and electron spectroscopic methods. The GO and rGOs' stacking nanostructure (flake) size (height x diameter), interlayer distance, average number of layers, distance between defects, elementary composition, content of oxygen groups, C sp3 and vacancy defects were determined. Different reductants applied to GO led to modification of carbon to oxygen ratio, carbon lattice (vacancy) and C sp3 defects with various in-depth distribution of C sp3 due to oxygen group reduction proceeding as competing processes at different rates between interstitial layers and in planes. The reduction using sodium borohydride and hydrazine in contrary to other reductants resulted in a larger content of vacancy defects than in GO. The thinnest flakes rGO obtained using sodium borohydride reductant exhibits the largest content of vacancy, C sp3 defects and hydroxyl group accompanied by the smallest content of epoxy, carboxyl and carbonyl groups due to a mechanism of carbonyl and carboxyl group reduction to hydroxyl groups. This rGO similar diameter to GO seems to result from a predominant reduction rate between the interstitial layers. The thicker flakes of a smaller diameter than in GO are obtained in rGOs prepared using remaining reductants and result from a higher rate of reduction of in plane defects.
Introduction

Wide varieties of methods for graphene oxide (GO) reduction have been recently utilized [1–5]. These methods include thermal, microwave, photo, chemical, electrochemical and solvothermal reduction. The most commonly applied chemical reduction routes use a wide variety of chemical compounds. Strong reducing agents are hydrazine and its derivatives (hydrazine hydrate and dimethylhydrazine), metal hydrides (sodium hydride, sodium borohydride), hydroiodic acid, etc. Other, weak reducing agents include ascorbic acid, hydroquinone, pyrogallol, hot strong alkaline solutions (KOH, NaOH), hydroxylamine and urea [1 and references included]. Reduction procedures of GO using different conditions, i.e. vapor, aqueous solutions of different concentrations and time, result in preparation of reduced graphene oxide (rGO) of variety of chemical and structural properties.

The main criteria for estimating the effect of reduction are C/O ratio, optical properties, structural defects and electrical conductivity. The electrical conductivity of a monolayer graphene depends on the carrier transport and therefore rGO chemical properties, where the oxygen groups attached to the charge carrying plane are more important in contrary to oxygen groups attached to the edges of the charge carrying planes. Reduction of chemically prepared GO results in rGO arranged in graphene stacking nanostructures (flakes) of various average height and diameter with graphene interlayers separated by a distance depending on the degree of the reduction, which implicates the average number of layers in graphene flakes [5]. Therefore, the conductivity of rGO depends not only on C/O ratios, the number of still existing oxygen groups and their ratios, but also on structural properties of the obtained rGO flakes and layers like size of flakes, type and content of defects in graphene layers.

Description of structural and chemical properties of graphene materials basing on results obtained using X-ray photoelectron spectroscopy (XPS), X-ray excited Auger electron spectroscopy (XAES), reflection electron energy loss spectroscopy (REELS), Raman spectroscopy, X-ray diffraction (XRD), transmission electron microscopy (TEM), Fourier transformed infrared spectroscopy (FTIR), atomic force microscopy (AFM), etc., has been often reported [1–19]. The FTIR and XPS analyses are usually applied for probing the chemical properties of atoms, carbon hybridizations and oxygen groups, where FTIR analysis information is related to the bulk, and XPS information depth is limited to the surface, i.e. few nanometers. By the recently developed complementary approaches of Raman spectroscopy [8–19] of few micrometers information depth, the surface AFM [8] and REELS [5] methods provide the information on the structural and chemical properties like level of disorder, dopants, the number of layers, the linear dispersion of electronic energy, the crystal orientation, the defects and strain [8–19].

Although, numerous procedures of reduction of GO using various reducing agents have been proposed [1 and references within], there are still many points to clarify like how oxygen groups are reduced, how lattice defects are being formed and reconstructed and what are the mechanisms of reduction and carbon lattice reconstruction using various chemical preparations and compounds. An attempt to reveal these differences was made using FTIR, Raman and surface XPS, XAES and REELS studies utilizing different in-depth sensitivity. The particular aim of the present work was to obtain information on differences in effectiveness of the chemical reduction of GO flakes performed in aqueous solutions of various chemical compounds, i.e. strong and weak reductants like sodium borohydride (NaBH₄), hydrazine (N₂H₄), formaldehyde (CH₂O), sodium hydroxide (NaOH), L-ascorbic acid (C₆H₈O₆), as well as L-ascorbic acid (C₆H₈O₆) prepared at room temperature (RT). The study presents the detailed comparison of the structural and chemical properties of the reduced GO samples, however is not concerned with investigating the kinetics of the reduction reaction.

Information depth of electron spectroscopic methods

A convenient measure of surface sensitivity in electron spectroscopic methods (XPS, XAES, REELS) is the information depth, \( ID \), defined as the maximum depth, normal to the surface, from which useful signal information of a specified percentage, \( P \), of signal originates [20, 21]. Therefore, \( ID \) can describe the sampling depth of this specified percentage of signal. The averaged information depth called a mean escape depth, \( MED \), in XPS and XAES and mean
penetration depth, MPD, in REELS is defined as an average depth normal to the surface from which the specified particles escape [22, 23]. Assuming single scattering of electrons, these values can be evaluated from the equations given as follows [20, 21].

\[
ID(P)_{XPS} = \frac{1}{100} \ln \left( \frac{1}{1 - P/100} \right), \quad ID(P)_{REELS} = \frac{\cos \theta_{out} \cos \theta_{out} \ln \frac{1}{1 - P/100}}{\cos \theta_{in} + \cos \theta_{out}},
\]

\[
MED_{XPS} = \frac{\cos \theta_{out}}{\cos \theta_{in} + \cos \theta_{out}}, \quad MPD_{REELS} = \frac{\cos \theta_{in} \cos \theta_{out}}{\cos \theta_{in} + \cos \theta_{out}},
\]

where \( \lambda \) is the inelastic mean free path, IMFP, of electrons scattered inelastically dependent on the kinetic energy, KE, of the electrons and on the material, \( \cos \theta_{in} \) and \( \cos \theta_{out} \) are angles of incidence and emission of electrons with respect to the surface normal of the specimen and \( P \) is a specified percentage of the signal considered. The values of \( ID(P = 99\%) \), \( MED \), \( MPD \) for the applied geometry of analysis and values of IMFPs in graphite published by Shinotsuka et al. [24] evaluated for the investigated transitions are listed in Table 1. The considered theoretical values of IMFPs [24] are in agreement with the experimentally determined values of IMFPs for bi-, tri- and multilayer graphene as found by Xu et al. [25].

The in-depth profile of graphene layers from REELS spectra is evaluated using an exponential interpolation of intensity of reflected inelastically scattered electrons, where the signal can be attenuated according to equation [26]:

\[
I = I^{\infty} (1 - \exp(-z/\lambda \cos \theta_{out})), \quad (2)
\]

where \( I \) is the signal intensity from the overlayer of thickness \( z \), \( I^{\infty} \) is the signal intensity from the overlayer material of infinite thickness [5], \( \lambda \) is the IMFP [24] and \( \theta_{out} \) is the electron emission angle with respect to the surface normal.

**Experimental**

**Sample preparation**

The following samples were investigated: a commercially available graphite (denoted as Gr) of 99.0% purity (AcrosOrganics, USA, 325 mesh) and expanded graphite (denoted as Gr-exp), graphene oxide (denoted as GO) and reduced graphite oxide samples prepared from GO using different reductants, i.e. sodium borohydride (NaBH₄), hydrazine (N₂H₄) [5], sodium hydroxide (NaOH), formaldehyde (CH₂O) and L-ascorbic acid (C₆H₈O₆). These reduced graphene oxide samples were denoted as rGO-NaBH₄, rGO-N₂H₄, rGO-NaOH, rGO-CH₂O and rGO-C₆H₈O₆.

The GO sample was prepared from commercially available graphite using a modified Hummers method [27]. The reduction using NaBH₄, NaOH, C₆H₈O₆ and CH₂O reductants proceeded in 100 ml water suspension of GO (0.6 wt.%) mixed with 50 ml 1 M water solution of NaBH₄, NaOH, C₆H₈O₆, and C₃H₂O₆, whereas reduction of GO using N₂H₄ proceeded using 50 ml of 50% solution of hydrazine hydrate. The GO and reducing agent solutions were boiled under a reflux for 30 min. Then, all the reaction mixtures were cooled to RT, washed in deionized water till the pH of filtrate stabilized to ca. 7–8, and then they were dried. Additional reduction procedure using L-ascorbic acid (C₆H₈O₆) at RT was proceeded under stirring of GO and C₆H₈O₆ in water solution for 30 min and then washing in deionized water till pH filtrate stabilized to ca. 7–8 and drying.

**HR-TEM, EDX, PEELS and XRD apparatus**

A high resolution transmission electron microscope (HR-TEM), Tecnai 20F X-Twin, equipped with an electron source, a cathode with field emission gun (FEG), EHT = 200 keV, a camera Orius and a high-angle annular dark-field (HAADF) detector, was used. This TEM was equipped with energy-dispersive X-ray spectrometer (EDX) with the energy resolution of 134 eV (EDAX RTEM SN9577+) and parallel electron energy loss spectrometer (PEELS) with the energy resolution of 0.8 eV. The quantification was performed according to the modified standardless/thin foil method. The preparation of samples proceeded in the following steps: sonication for 5 s of a few milligrams of sample in ethanol (99.8% anhydrous) using ultrasounds, applying a drop of the solution of 5 μl on a carbon coated copper mesh with holes (Lacey type Cu 400 mesh, Plano), evaporating the solvent at room temperature and then investigating the remaining dried powder stuck on the copper mesh.
Powder X-ray diffraction (pXRD) patterns were obtained by using an X’Pert Pro diffractometer, with Cu Kα radiation (λ = 1.5406 Å) and with a X’Celerator detector.

**Raman apparatus**

A Raman micro-spectrometer, Senterra, Bruker Optik with a 532 nm laser was used. The measurement parameters are the following: laser power of 2 mW, acquisition time of 360 s, number of scans 2 and analyzed range of 800–3000 cm⁻¹.

**FTIR apparatus**

The FTIR spectra in the 70–4000 cm⁻¹ range were recorded in vacuum spectrometer with Fourier transformation, Vertex 70 V, Bruker Optik, at T = 22 °C, p = 10⁻¹ Pa, resolution 4 cm⁻¹ and number of scans 100. Before the sample measurement, a “vacuum spectrum” was recorded and subtracted automatically as background during registration of spectra for the investigated samples. The sample was mixed with KBr at a ratio of 1/300 mg and then compressed at 7 MPa cm⁻² to form a pellet, and the transmission spectrum was recorded in the range of 750–1900 cm⁻¹.

**XPS apparatus**

The XPS measurements were carried out in an ultra-high-vacuum (UHV) using the ESA-31 electron spectrometer (homemade) [28]. The spectrometer is equipped with a hemispherical electron energy analyzer of a high relative energy resolution of 0.5% without retardation (the retarding ratio, k, can be changed from 2 to 100), an electron gun (LEG62-VG Microtech), a homemade X-ray excitation source (Al Kα X-rays hv = 1486.67 eV) and an Ar⁺ ion source of AG21 (VG Scientific). The XPS spectra were measured in the fixed retarding ratio (FRR) mode (k = 4, 8, 16) at a photon incidence and electron emission angles of 70° and 0°, respectively, with respect to the surface normal of the specimen. The REELS spectra were measured using a primary electron energy of 4 keV (k = 41, generally keeping the analyzer pass energy at about 50–100 eV) and the electron beam current intensity of a few nA at electron incidence and emission angles of 50° and 0° with respect to the surface normal of the specimen, respectively.

The prepared powder of graphite, GO and rGO samples was placed on a holder using an UHV tape and investigated without any thermal and/or chemical pretreatment.

**Results and discussion**

**Structural properties of graphene flakes by HR-TEM, EDX and PEELS**

The EDX spectra of exemplary rGO-NaBH₄ and rGO-CH₂O samples (Fig. S1) reveal in the bulk the presence of C, O and Na. The PEELS spectra show differences in the intensity of elastic peak and inelastic loss peaks in the energy loss region of about 50–400 eV, as well as inelastic energy loss values indicating structural variations between the presented samples.

The TEM images of exemplary rGOs (Fig. 1) show transparent parts confirming graphene layers structure with dark parts due to the overlapping layers.
Graphene interlayer distance and size of flakes by XRD

The XRD (002) and (10) patterns of the investigated samples (Fig. 2) were fitted using Pearson7 function (Fig. S2). The resulting fitting parameters, i.e. Bragg’s angle of (002) and (10) patterns and the respective full-width at half maximum (FWHM) values, are listed in Table S1. The structural parameters, i.e. the average interlayer distance, \( d \), height of the stacking nanostructures, \( H \), number of layers in the flakes, \( n \), and diameter of the flakes, \( D \), evaluated using Bragg’s and Scherrer’s equations are listed in Table 2. For evaluating the value of \( H \) from (002) pattern and \( D \) from (10) pattern, the constants of 0.9 and 1.84 were considered, respectively.

The structural parameters of GO, rGOs vary depending on the reductant (Table 2). The interlayer distance is 0.721 nm (GO) and 0.446–0.345 nm (rGOs). These values confirmed elsewhere, i.e. 0.335 nm (graphite), 0.335–0.4 nm (rGO) and 0.77–0.9 nm (GO) increase with humidity [5, 29–31]. The rGO reduced using NaBH\(_4\) shows considerably large flake diameter of the smallest thickness and average number of graphene layers. These average number of layers in different rGO flakes increases in the following order: rGO-NaBH\(_4\) < rGO-N\(_2\)H\(_4\) < rGO-C\(_6\)H\(_8\)O\(_6\)--RT < rGO-C\(_6\)H\(_8\)O\(_6\) < rGO-CH\(_2\)OH < rGO-NaOH.

Structural properties of graphene flakes by Raman spectroscopy

The Raman spectrum is considered to depend on: (i) clustering of the sp\(^2\) phase, (ii) bond disorder, (iii) the presence of sp\(^2\) rings or chains and (iv) the sp\(^2\)/sp\(^3\) ratio, where these factors act as competing forces influencing the shape of the Raman spectra. Classification according to these features was proposed by Ferrari and Robertson [12] as an amorphization trajectory consisting of three stages ranging from graphite to tetrahedral amorphous carbon (ta-C) or defected diamond, i.e. (1) graphite to nanocrystalline graphite (nc-Gr), (2) nanocrystalline graphite to amorphous carbon (a-C) and (3) a-C to 100% sp\(^3\) ta-C or defected diamond. The classification and interpretation of the Raman spectra require consideration of intensity and width (FWHM) of the Raman modes. The main features of Raman spectra of graphene show characteristic major bands assigned to the first-order \( D \) (\( \sim 1320–1350\) cm\(^{-1}\)), \( G \) (\( \sim 1570–1605\) cm\(^{-1}\)), \( D' \) (\( \sim 1620\) cm\(^{-1}\)), \( D + D' \) (\( \sim 2900\) cm\(^{-1}\)) modes and the second-order 2D mode (2640–2680 cm\(^{-1}\)) (Fig. 3). The \( D \) mode is an in-plane breathing vibration of the six-membered carbon rings and is considered as a disorder-induced band. Therefore, the \( D \) band reflects defects like destroyed carbon hexagons, and it is not present for a perfect graphite structure in contrary to amorphous structures, for which its intensity is high. The intensity and FWHM of \( D \) band depend on disorder and the type of edges, where the intensity is higher for armchair than for zigzag edges. The \( G \) mode is due to the in-plane stretching vibration of carbon atom pairs and is observed for all carbon structures containing sp\(^2\) bonds, both aromatic carbon and other sp\(^2\) structures. The 2D allowed mode is the most intense for an ideal single-layer graphene. A low intensity \( D \) band accompanied by a presence of 2D band indicates a high-quality graphene. The main features of the evolution of Raman spectrum in stage 1 are appearance of \( D \) mode, increasing of \( I(D)/I(G) \) ratio following Tuinstra and Koenig (TK) relations [13], appearance of \( D' \) mode, disappearance of \( D \) and 2D mode doublet structure, appearance of \( D + D' \) mode and at the end of stage 1, overlapping of \( G \) and \( D' \) modes. The ratio of intensity of \( D \) to \( G \) mode is inversely proportional to the size of effective crystallite size in the direction of the graphite plane, \( L_a \), or graphitic cluster, i.e.:

\[
I(D)/I(G) = C(\lambda)/L_a
\]

where \( \lambda \) is Raman wavelength and \( C(515.5\text{ nm}) = 4.4\) nm [12]. In the stage 2, the position of \( G \) mode decreases, the TK relations fails and \( I(D)/I(G) \) decreases. The broad feature from about 2300 cm\(^{-1}\) to 3200 cm\(^{-1}\) modulated by 2\( D \), \( D + D' \) and 2\( D' \) modes appears instead of a well-defined second-order peaks. For stage 2, the new relation was proposed by Ferrari and Robertson [12]:

\[
\frac{I(D)}{I(G)} = C'(\lambda)L_a^2
\]

with \( C'(514) \sim 0.55\) nm\(^{-2}\).

The stage 3 is beyond the data presented and therefore is omitted in discussion.

In all stages of amorphization trajectory, the structural defects exist lowering the symmetry of the infinite crystal and they include: (i) the point-like defects, (ii) cluster defects and (iii) boundary or edge defects. The experimental investigation of defects [17] in Ar bombarded graphene and graphite with
different grain size like: (i) vacancy defects produced by the deformation of the carbon lattice bond, (ii) on-site defects, which describe out-of-plane atoms bonded to carbon atoms (namely sp$^3$ hybridization) and (iii) boundary or edge defects, provided detectable $D$ and $D'$ modes modification. The experimental data showed that $I(D)/I(D')$ is maximum ($\sim 13$) for defects associated with sp$^3$ hybridization, it decreases for vacancy-like defects ($\sim 7$) and reaches a minimum for boundary-like defects in graphite ($\sim 3.5$) [17]. Other work by Lucchese et al. [18] studying disorder and defects in graphene caused by Ar ion bombardment provided quantification of defects, i.e. the average distance between point-like defects, $L_D$, and density of defects, $n_D (L_D \sim 1/\sqrt{n_D})$ by fitting the observed $I(D)/I(G)$ versus $L_D$ using a phenomenological model. The values of $I(D)/I(G)$ have a non-monotonic dependence on $L_D$, increasing with increasing $L_D$ up to $L_D \sim 4$ nm in stage 2, and then decreasing for $L_D > 4$ nm (stage 1) and being in agreement with a well-established graphitization trajectory for carbon materials proposed by Ferrari and Robertson [12]. Such behavior suggests the existence of two disorder-induced competing mechanisms contributing to the Raman $D$ band. The phenomenological model considers that the impact of a single ion in the graphene sheet causing modifications on two length scales, here denoted by $r_A$ and $r_S$ (with $r_A > r_S$), which are the radii of two circular areas measured from the impact point and subscript $A$ stands for “activated,” whereas the subscript $S$ stands for “structurally-defective,” with values of $r_S = 1$ nm and $r_A = 3$ nm [19]. This model extended by Concado et al. [19] to other Raman lines is valid for $L_D > 10$, i.e. for a low-density defects materials.

The measured shapes, positions, relative intensities of $D$, $G$, $2D$, $D' + D'$ modes and FWHM of Raman
spectra are characteristic for carbon materials (Fig. 3, Table 3, Fig. S3–S5). According to the above, both graphite samples can be classified to stage 1, whereas GO and rGOs to stage 2. The values of $I_a$ evaluated according to Eqs. (3) are listed in Table 3. Therefore, graphite samples exhibit a low defect density, whereas GO and rGO exhibit a high defect density. The stage 1 Gr and Gr-exp samples show spectra of a high intensity and small FWHM of $G$ mode indicating dominating graphite structure of low defect density. These samples are characterized by the
The stage 2 of amorphization trajectory includes GO and rGO samples, which characteristic features are high intensities and width of FWHM of D and G modes, very weak intensity of 2D mode and values of I(D)/I(G) close to 1. The variation of positions and FWHM of D, G and 2D modes (Fig. S3-S5) confirms that the investigated samples have different thickness, C sp³ content, graphitic clusters size, distance between defects and their density (Fig. S5), crystallinity (Fig. S4, Table 3) [8–19]. The FWHM and intensity values of D and G mode indicate the distance between point-like defects smaller than 3 nm [19], for which phenomenological evaluation of distance between defects proposed by Concado et al. [19] is not valid. The rGOs exhibit higher crystallinity than GO, whereas the highest crystallinity is shown by graphite and expanded graphite (Fig. S4). As reported elsewhere for rGOs [12], increasing frequency of D band position and decreasing frequency of G band position accompanied by decreasing FWHM of D and G bands (Fig. S3) is related to decreasing C sp³ content and increasing Ld. While increasing Ld value provides decreasing position and FWHM of 2D mode [20], the increasing number of layers leads to an opposite effect [10]. Therefore, the 2D band in Raman spectra exhibits these two effects (Fig. S3).

### Chemical groups in graphene flakes by FTIR spectroscopy

The FTIR measurement procedures (transmittance of KBr pellets) do not provide quantitative determination of surface (or volumetric) concentration of chemical moieties (Fig. 4). The comparison of band intensity for various samples cannot provide clear information about their chemical structure because of the arbitrary background subtraction and band extension. A stoichiometric structure of GO, heterogeneity of the oxygenated functional groups and their interaction lead to overlapping of the characteristic bands and affect their position and intensity. However, the recorded spectra indicate: (i) the presence (or absence) of the chemical structures and (ii) changes in relative intensities of the respective bands caused by reduction procedures informing on chemical structures transformation.

In all FTIR spectra (Fig. 4), two main range of absorption bands 1750–1450 cm⁻¹ and 1300–950 cm⁻¹ characteristic for C = O and C-O moieties, respectively, can be noticed [6, 7]. The left side of the spectra starts with more or less shaped band near 1720 cm⁻¹ attributed to the stretching vibration of C = O moiety in carboxylic (conjugated and/or non-conjugated) or carbonate systems (acid, ester, anhydride, dioxolan). The bands at 1635 cm⁻¹ and 1565 cm⁻¹ can be assigned to aromatic carbon–carbon bonds, carbonyl moieties in various chemical surroundings (quinone-, ketone-, aldehyde-like, amide-like), carbon–oxygen ion-radical structures and conjugated systems (diketone, keto-esters, keto-enol and quinone-hydroquinone structures). The adsorbed/intercalated water provides deformation vibration band δ(HOH) located near 1630 cm⁻¹ overlapping with C–OH mode at 1620 cm⁻¹. The next set of the overlapping peaks, which form an absorption band in 1300–950 cm⁻¹ region, can be attributed to C-O moieties existing in a different structural environment. In this spectral region, the presence of C=O–C symmetric stretching vibration bands in ether-, ester- lactone-, pyrone-, furane-like structures should be taken into consideration. Also, phenol and hydroxyl molecular groups exhibit characteristic mode located near 1070 cm⁻¹ (C–OH vibration), while the epoxide group provides a mode near 1290 cm⁻¹ (asymmetric C=O–C stretching).

The graphite (Gr) spectrum was arbitrary extended and small amount of oxygen bonded with carbon in the form of carboxylic and ether groups, as well as strongly adsorbed water can be observed. The peak assigned to in-plane stretching of aromatic rings ν(C = C)) with frequency 1580 cm⁻¹ is visible (only partially overlapped). The oxidative treatment of graphite (GO samples) gives large enhancement of C = O and C-O bands described by the above mentioned oxygen-containing functional groups—the increase in relative intensity of these peaks and covering the aromatic peaks can be observed. The difference in the shape of the spectra recorded for Gr-exp and GO-foil—markedly the increase in relative intensity of the band in C–O–C region (near 1200 cm⁻¹)—can be explained by stronger dehydration of GO-foil with creation of anhydride structures (lactone). The GO reduction using different procedures modifies the FTIR spectra. After reduction with sodium hydroxide in rGO-NaOH, a relative decrease in the peak ascribed to carboxylic moieties (1720 cm⁻¹) and a relative increase of the band associated with the presence of hydroxyl groups
Similar changes can be observed for GO reduced with formaldehydes (rGO-CH$_2$O). The GO reduction with other weak reduction agent—L-ascorbic acid at RT (rGO-C$_6$H$_8$O$_6$-RT) and (rGO-C$_6$H$_8$O$_6$) changes the FTIR spectra due to extensive adsorption of organic molecules. Mainly the presence of the overlapped bands of aliphatic chains and hydroxyl functional groups can be observed. The reduction of GO with sodium borohydride (sample rGO-NaBH$_4$) completely dehydrates carbon materials (disappearance of a peak near 1630 cm$^{-1}$) creating anhydride structures (lactone, cyclic esters) without removing carbon–oxygen functionalities. In samples reduced using sodium borohydride and L-asorbic acid, a decrease of alkene stretching vibration (1535 cm$^{-1}$) and a presence of a strong amide stretching vibration (1565 cm$^{-1}$) in contrary to GO and rGOs reduced using formaldehyde and sodium hydroxide are observed. The smallest content of water is observed in rGO-NaBH$_4$, rGO-C$_6$H$_8$O$_6$-RT and rGO-C$_6$H$_8$O$_6$ in contrary to GO, rGO-NaOH and rGO-CH$_2$O.

**Average number of layers in graphene flakes by REELS**

The REELS spectrum of carbon nanomaterials reflects the structural features in intensity and energy loss values of reflected electrons losing their kinetic energy on valence electrons forming $\pi$ and $\pi + \sigma$ bonds [32, 33]. The values of intensity and energy loss are characteristic for the type of bonds (C sp$^2$ and C sp$^3$ hybridizations), and content of these hybridizations at the surface (S) and in the bulk (B) varies with chemical and structural properties of carbon nano-materials, i.e. graphite and diamond [32], single, bi-, triple-, multilayer graphene and graphene oxide [34–37]. For fitting the spectroscopic data reflecting the interaction of matter with electromagnetic radiation, the most common pseudo-Voigt profile, i.e. convolution of Lorentzian and Gaussian symmetric components, is used since Lorentzian refers to distribution of decaying oscillations and Gaussian to...

**Table 3** The parameters of $D$, $G$ and $2D$ bands of Raman spectra and the effective crystallite size in the direction of the graphite plane, $L_a$, for the investigated samples

| Sample          | $D$ band | $G$ band | $2D$ band | $I(D)$/$I(G)$ | $I(2D)$/$I(G)$ | $L_a$(nm) |
|-----------------|----------|----------|-----------|---------------|---------------|-----------|
|                 | Pos. (cm$^{-1}$) | FWHM (cm$^{-1}$) | Pos. (cm$^{-1}$) | FWHM (cm$^{-1}$) | Pos. (cm$^{-1}$) | FWHM (cm$^{-1}$) |
| GO-foil         | 1337     | 119      | 1594      | 82            | 2687          | 243       | 0.991  | 0.4650 | 1.34    |
| rGO-NaBH$_4$    | 1343     | 148      | 1587      | 97            | 2701          | 229       | 0.980  | 0.8422 | 1.34    |
| rGO-C$_6$H$_8$O$_6$-RT | 1342     | 110      | 1585      | 73            | 2691          | 206       | 1.017  | 0.7831 | 1.36    |
| rGO-C$_6$H$_8$O$_6$-RT | 1341     | 100      | 1585      | 79            | 2689          | 152       | 1.028  | 0.8410 | 1.37    |
| rGO-NaOH        | 1342     | 60       | 1585      | 68            | 2687          | 93        | 1.242  | 0.7210 | 1.5     |
| Gr              | 1340     | 65       | 1583      | 61            | 2683          | 101       | 1.133  | 0.8097 | 1.44    |
| Gr-exp          | 1342     | 45       | 1572      | 24            | 2710          | 72        | 0.214  | 0.4315 | 20.56   |
|                 | 1345     | 42       | 1575      | 24            | 2712          | 67        | 0.041  | 0.4604 | 107.32  |

**Figure 4** The FTIR spectra recorded from the investigated samples.
distribution of velocities leading to Doppler broadening. The REELS spectra recorded for the investigated carbon nanomaterials resulted as fitted Gaussian functions of surface and bulk C sp²/sp³ components (Fig. S6) as reported elsewhere [5]. Results of this fitting presenting the electron kinetic energy loss values typical for surface (S) and bulk (B) C sp² and C sp³ components, these components’ percentage contributions and the ratio of C sp²/B/C sp³S are listed in Table 4. For approximating the intensity ratio of π + σ energy loss peak components from C sp²B bulk to C sp³S surface as a function of graphene layer in-depth profile, z, Eq. (2) is applied. For a single layer graphene, the ratio of C sp²B/C sp³S is zero, whereas for graphite, it results from the experimental REELS spectra fitting (Fig. S6). The average number of layers, n, within the in-depth profile, z, of graphene flakes (Fig. 5) was evaluated using interlayer distances from XRD (Table 2). The values of n resulting from REELS (Table 4) and XRD (Table 2) for GO and rGOs flakes are consistent, as also reported previously [5].

Surface elemental content, C sp²/sp³ hybridizations and oxygen groups in graphene flakes

The quantification of the surface atomic content of GO and rGOs was carried out from the area under C 1 s, O 1 s, N 1 s, Na 1 s, Si 2p and S 2p photoelectron peaks after using Tougaard background subtraction [38]. The atomic content was evaluated from the XPS MultiQuant software [39, 40] accounting for Scofield photoionization cross sections [41], electron elastic scattering and analyzer transmission function. The results are presented in Table S2 and Fig. 6.

Samples of GO and rGO show contaminations of N, Na, S and Si (Table S2, Fig. 6a). For rGOs, the increasing content of C and decreasing content of O are observed with increasing number of layers in flakes (Fig. 6). The C 1 s and O 1 s spectra after Tougaard inelastic background subtraction [38] were fitted as suggested by Kevtun et al. [42] to pseudo-Voigt functions (combination of Gauss and Lorentz asymmetric and symmetric components) using the XPSPeakfit software [43]. The C1s spectra components binding energy (BE) values for different chemical forms, i.e. C sp²/C sp³ hybridizations and oxygen groups (hydroxyl – C-OH, epoxy – C-O-C, carbonyl – C = O and carboxyl – C-OOH), were applied as reported elsewhere for different carbon nanomaterials like nanodiamonds, graphite, carbon nanotubes, graphene oxide and graphene [42,44–48 and references within]. The vacancy defects were considered at about 283.9 eV as suggested by Barinow et al. [48]. The O 1 s BE of carbon–oxygen groups was selected in the range of experimental [5, 49, 50] and calculated values [50]. The resulting fitted spectra are shown in Fig. S7, S8. The values of C 1 s and O 1 s BEs for carbon lattice vacancies, C sp²/sp³ hybridizations and carbon–oxygen groups resulting from the above fitting procedure shown in Table S3 and S4 are in agreement with a literature data [5, 42, 44–50]. The evaluated oxygen content resulting from C 1 s and O 1 s spectra fitting is in a reasonable agreement (Fig. S9).

Comparison of carbon–oxygen groups content is provided in Fig. 7a–b. For all carbon nanomaterials the C 1 s BE of C sp² component is 284.5 ± 0.1 eV (Table S3). Samples of graphite show at the surface C–OH and C = O groups and traces of an adsorbed water. Samples of GO and rGO show additionally C–O–C and C=OOH groups and a larger content of water at O 1 s BE of 535.2 ± 0.1 eV, 536.4 ± 0.3 eV, 538.3 ± 0.3 eV, attributed to adsorbed water, liquid water and liquid and gas phase water (overlapping peaks), respectively, as reported elsewhere [51–54]. Reduction of GO decreases the amount of carbon–oxygen groups C–OH > C–O–C > C = O > C-OOH and vacancy and C sp³ defects (Table S3). This is observed with increasing number of graphene layers in flakes (Fig. 7a). The ratio of rGO to GO carbon–oxygen groups decreasing for C–OH and increasing for C-OOH > C = O > C–O–C accompanied by dehydration (Figs. 4, 7a, Table S4) with number of layers suggests different rate of oxygen groups reduction influencing the thickness of rGO flakes (Fig. 7b).

The dependence of number of graphene layers on the in-depth distribution of C sp³ defects resulting from evaluation using: (i) the first derivative of Auger C KLL spectrum [55] from which the parameter D is calculated [46] (C sp³ C KLL) (Fig. S10), (ii) C 1 s spectra fitting (C sp³ C 1 s) (Table S5), REELS spectra fitting of (iii) surface (C sp³ REELS-surface) and (iv) bulk (C sp³ REELS-bulk) components (Table 4) is provided in Fig. 8a. The GO-rGOs show surface C sp³ enrichment. The content of C sp³ decreases with a depth since an average information depth increases as following: REELS-surface (outer surface layer) < C
KLL (1–2 layers) < REELS-bulk ~ C 1 s (2–5 layers) (Table 1). The smallest difference between the outer layer and 1–2 layers below obtained for rGOs of the thinnest flakes indicates the most efficient penetration of a reducing agent between the layers, i.e. case of rGO-NaBH4 and rGO-N2H4 (Fig. 8b).

**Influence of reducing agents on structural and chemical properties of graphene flakes**

Dependence of different physicochemical properties of rGO flakes on the number of layers is presented in Figs. 6–9. The REELS indicates that the thickness of graphene flakes in GO and rGOs increases in the order of GO-foil-fresh < rGO-NaBH4 < rGO-N2H4 < rGO-C6H8O6 < rGO-CH2O < rGO-NaOH. This is accompanied by increasing graphitic cluster size, \( l_{w} \), from 1.34 nm to 1.5 nm (Table 2) and C/O ratio (Fig. 6), distance between defects and decreasing defect density, decreasing interlayer distance, content of C sp\(^3\) hybridizations and vacancy defects (Fig. 9), carbon–oxygen groups and water situated between the interstitial layers (Fig. 7, Table S3, S4). The reduction using various reductants proceeds with different rate, where oxygen group content decreases in the following order: C–OH [C–O–C [C=O [C-OOH (Fig. 7a).

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**Figure 5** Dependence of C sp\(^2\)B/C sp\(^2\)S ratio on the penetration depth for electron signal information depth \( ID (P = 99\%) \), IMFP in graphite from Shinotsuka [24] for evaluating the average number of layers resulting from REELS and Raman spectra.
G modes in Raman spectra provide evidence on average distance between defects $L_D < 3$ nm [19]. For rGOs, the XPS results indicate decreasing density of C sp$^3$ and vacancy defects with increasing number of layers in graphene flakes (Fig. 9a). The estimate of $L_D$ from C sp$^3$ and vacancy defects (Table S3) provides for GO the value of $L_D < 1$ nm (C sp$^3$ defects) and $L_D \sim 15$ nm (vacancy defects), whereas for rGOs, $L_D \sim 1$–2 nm (C sp$^3$ defects) and 4–15 nm (vacancy defects). The oxygen content obtained from XPS (Table S2) in a range of 26.1 at.% (GO-foil-fresh) to 8.6 at.% (rGO-NaBH$_4$) provides an estimate of distance between defects of $L_D^* \sim 1$ nm and $L_D^* \sim 3$ nm, respectively, which remains in agreement with the Raman spectroscopy data.

The modification of the flake size of rGOs, i.e. height and diameter, depends on the applied reductants (Fig. 9b). The rGO-NaBH$_4$ of the thinnest flakes (Table 4) and flake diameter similar to GO (Table 2) shows the largest number of surface carbon–oxygen groups and vacancy defects of larger content than in GO (Table S3, Fig. 7a, Fig. 9a). Also, no water is present between the interstitial sites (Fig. 4) and the ratio of rGO to GO group content decreases in the following order: C–OH > C=O > C–O–C due to a mechanism of NaBH$_4$ reduction converting C = O and C-OOH to C–OH groups as proposed by Samulski [56]. The rGO-N$_2$H$_4$, rGO-C$_6$H$_8$O$_6$, rGO-NaOH, rGO-CH$_2$O of thicker flakes and smaller diameters than GO exhibit a similar content of vacancy defects than GO, a smaller content of C sp$^3$ defects and a content of oxygen groups decreasing in an order: C-OOH > C = O > C–OH > C-O–C. The binding energies for carbon–oxygen groups in graphene and defected...
graphene, where edges may be considered as defected graphene, do not vary distinctly [57]. However, a predominant rate of reduction between the interstitial site over that in plane results in thin flakes of a large diameter, whereas a higher rate of reduction in plane leads to smaller diameter flakes of larger number of layers. The reduction of GO at RT conditions in contrary to boiling conditions (e.g. exemplary rGO-C₆H₈O₆-RT sample) results in a larger number of C sp³ and vacancy defects, carbon–oxygen groups, interstitial water and smaller average number of layer in flakes. The boiling reduction conditions lead to reparation of the vacancy defects and provide a higher reduction rate between the interstitial sites. Aging of GO (sample GO-foil-fresh and GO-foil) provides a larger number of layers in flakes, which could be influenced by additional oxidation and adsorption of water leading to graphene layers stacking.

Summary of results and conclusions

The bulk and surface properties of GO prepared from graphite using the modified Hummers method and rGOs obtained from this initial GO using a range of

![Grapha](image1.png)

**Figure 8** Comparison of a C sp³ content resulting from Auger C KLL spectra parameter D evaluation (Table S5), C 1 s spectra fitting (Table S3) and REELS spectra fitting (Table 4) the samples are indicated in the order of increasing number of layers and b difference in C sp³ percent resulting from C KLL, C 1 s and REELS-surface and REELS-bulk components. Notation ”*”—sample from Ref [5].

![Graphb](image2.png)

**Figure 9** Dependence of a the atomic percent of vacancy and C sp³ defects (Table S3) obtained from XPS spectra and b ratio of height and diameter values of rGOs to GO flakes (Table 2) on average number of layers in graphene flakes obtained from XRD and REELS (Table 4).
reducing agents were characterized by applying a set of complementary methods. The dependence of rGOs chemical and structural properties on the preparation procedures applying different reductants was investigated. Novelty and significance of these study rely on: (i) investigating the effect of few reducing agents like conventional and “green” reductants in similar conditions, (ii) characterizing GO and rGOs structural and chemical bulk and surface properties, (iii) providing correlation between rGOs bulk and surface properties due to the applied reductants, (iv) applying various complementary methods of analysis such as XRD, EDX, HR-TEM, PEELS, Raman spectroscopy, REELS, XPS and XAES, (v) confirming reliability of different methods for evaluating the selected physicochemical properties.

The GO and rGOs showed different interlayer distances in the range of ca. 0.446–0.345 nm with the largest interlayer distance for GO (0.721 nm). The GO and GO flakes of different size, i.e. diameter and height containing various average number of layers, were obtained. The number of layers in flakes resulting from XRD and REELS analyses was in agreement. For rGOs, the number of layers in flakes increased in the following order: \( \text{NaBH}_4 < \text{N}_2\text{H}_4 < \text{C}_6\text{H}_5\text{O}_6 < \text{RT} < \text{CH}_3\text{O} < \text{NaOH} \) from 3.4 ± 2 nm to 12.4 ± 2.1 nm. This was accompanied by increasing graphitic cluster size from 1.34 nm to 1.5 nm, distance between defects and decreasing defect density. The Raman and XPS data provided in GO-rGOs a consistent value of distance between defects, \( \text{LD} \), from 1 to 3 nm. The content of oxygen groups decreased in the order of \( \text{C–OH} > \text{C–O}–\text{C} > \text{C} = \text{O} > \text{C–OOH} \) and the ratio of these oxygen groups in rGO to GO showed decrease for C–OH groups and increase for C–OOH > C = O > C–O–C groups. This provides an evidence on the importance of C–OH groups present in planes and edges to obtain thin graphene flakes. The GO showed the most inhomogeneous distribution of oxygen groups and C sp\(^3\) hybridizations at the surface and in the bulk. In contrary to GO, the rGOs obtained using various reductants resulted in decreasing differences between in-depth distribution of the C sp\(^3\) and carbon–oxygen groups with increasing number of layers. The applied reductants provided rates of reduction between the interstitial sites, which are different from the rates in the planes. The thinnest flakes were obtained for rGO-NaBH\(_4\) characterized by the largest density of vacancy and C sp\(^3\) defects, content of C–OH group, inhomogeneous in-depth distribution of C sp\(^3\), where the reduction proceeded predominantly between the interstitial sites.

**Acknowledgement**

The authors (B.L., L.K, J.T.) acknowledge the support of the bilateral project between the Polish Academy of Sciences and the Hungarian Academy of Sciences. The authors (B.L., N.R., L.K.) acknowledge the support of the Horizon 2020 MSCA-COFUND agreement No. 711859 and the financial resources for science in the years 2017–2021 awarded for the implementation of an international co-financed project (3549/H2020/COFUND/2016/2). The author (J.T.) acknowledges the support by the European Regional Development Fund and Hungary in the frame of the project GINOP-2.2.1-15-2016-00012. The author (G.T.) acknowledges the support of the National Science Centre Poland for financial support (Grant “MINIATURA” No. 2017/01/X/ST5/01248). The author A.M thanks the National Centre for Research and Development for support of the project LIDER/33/0117/L-9/17/NCBR/2018. The author L.S. would like to thank for the support of the project NCBiR (2018-2012) POIR.01.01.01-00-0802/17-00 (Polski Bazalt S.A.).

**Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no conflict of interest.

**Electronic supplementary material:** The online version of this article (https://doi.org/10.1007/s10853-020-05461-1) contains supplementary material, which is available to authorized users.

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