Biobased Janus molecule for the facile preparation of water solutions of few layer graphene sheets

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A biobased Janus molecule was used to prepare water solutions of nano-stacks made by few layer graphene. The Janus molecule was 2-(2,5-dimethyl-1H-pyrrol-1-yl)-1,3-propanediol (serinol pyrrole, SP), a serinol derivative obtained through the neat reaction of 2-amino-1,3-propanediol with 2,5-hexanediol, with atom efficiency of about 85%. SP contains the pyrrole ring, suitable for the interaction with carbon allotropes and hydroxy groups, that can easily interact with polar surroundings. Adducts, with about 14% by mass of SP, were prepared by reacting SP with a high surface area nano-sized graphite (HSAG), with about 35 graphene layers stacked in crystalline domains. Green methods were adopted, such as ball milling (HSAG–SP–M) and heating. Infrared spectra revealed peaks due to both HSAG and SP and additional peaks that could be attributed to the adduct. Both thermal and mechanical reactions left substantially unaltered the order in the graphitic layers and the interlayer distance, as shown by X-ray diffraction patterns. The relative intensity of the G and D band in the Raman spectrum was not modified by the thermal reaction, whereas enhancement of the D peak was observed after ball milling. Stable water solutions of HSAG–SP–M were prepared in a concentration range from 0.1 to 1 mg mL$^{-1}$. Centrifugation allowed isolation of adducts with few stacked graphene layers, as revealed by high resolution transmission electron microscopy. An image of the adduct showed an organic layer tightly adhered to the carbon surface. SP appears a suitable molecule for the easy functionalization of carbon allotropes, such as nano-stacks of graphene layers, without substantially affecting the bulk crystalline organization and promoting the separation of the aggregates into stacks containing a low/very low number of graphene layers.

1. Introduction

Since the discovery of fullerenes, nano-sized carbon allotropes have been the subject of huge amounts of research activity. Carbon nanotubes (CNTs), both single and multi-walled, or graphitic nanofillers made by few layers of graphene are studied as they dramatically improve properties such as electrical and thermal conductivity, mechanical reinforcement, thermal and flame resistance, of both thermoplastic and elastomeric matrices. In recent years, attention has been in particular focussed on graphene, in consideration of its huge surface area, exceptional mechanical strength and high carrier mobility.

Single or few layers of graphene can be obtained through micromechanical cleavage, epitaxial growth and chemical vapor deposition of graphene films, liquid phase exfoliation. Much interest has been directed, particularly in view of large scale production, to the oxidation of graphite to graphite oxide (GO) followed by thermal or chemical reduction. Such method gives also the chance of preparing GO. The precise structure of GO is still unknown. However, the adopted oxidation methods lead to extensive modification of the graphene layers. It was reported that hydroxy and epoxide groups are mainly present on the surface of basal planes and carbonyl and carboxyl groups are bounded to the edges. GO can be dispersed in polar media and can be the platform for further functionalizations. The oxidation–reduction method is however characterized by several drawbacks. The oxidation requires strong acids, explosive oxidation agents, harsh reaction conditions and long reaction times. Thermal reduction occurs at high temperature and under controlled conditions, efficient chemical reducing agents could be toxic, hazardous and expensive, such as, for instance, those based on hydrazine. Besides that, it is increasingly acknowledged that the complete restoration of sp$^2$ hybridization of graphene carbons can be hardly achieved.

Aim of this work was the introduction, on graphene layers, of oxygen containing functional groups, in particular of hydroxy groups, without substantially affecting the sp$^2$ hybridization of the carbon atoms, thus keeping structural order inside graphene sheets. The achievement of this goal could allow the easy preparation of solutions in polar media (e.g. in water) of
graphene layers with almost unchanged thermal and electrical conductivity, without the need of further chemical treatments. The peculiar objective of the work was to perform the modification of graphene layers through simple reactions inspired to the principles of green chemistry, ideally by using biobased chemicals.

To pursue these objectives, 2-amino-1,3-propanediol, known as serinol (S), was identified as the starting building block. Serinol can be directly obtained from renewable sources\(^\text{83}\) and is commercially available, arising from the reaction of easily available chemicals such as glycerol. The chemoselectivity of its amino and hydroxyl groups was exploited\(^\text{84,85}\) to perform Paal–Knorr reaction\(^\text{68,69}\) of S with 2,5-hexanedione (HD) obtaining 2-(2,5-dimethyl-1H-pyrrol-1-yl)-1,3-propanediol, shown in Fig. 1 and hereinafter indicated as serinol pyrrole (SP).

The reaction was performed in the absence of solvents and catalysts, with yield of about 95% and atomic efficiency of about 85%. SP contains two hydroxyl groups, that promote the compatibility with polar surroundings such as water, eco-friendly solvents as alcohols, ethers and esters and polar polymer matrices. SP contains as well pyrrole ring, that could favour \(\pi-\pi\) interaction with the aromatic rings of carbon allotropes. SP appears as a \textit{Janus Bifrons} molecule, suitable for the preparation of solutions of graphitic nanofillers in polar media.\(^\text{65,67,70,71}\)

Reaction was performed between SP and a nano-sized graphite with very high surface area (HSAG), higher than 300 m\(^2\) g\(^{-1}\), commercially available and obtained through milling. HSAG was characterized by a relatively low number of layers stacked in the direction orthogonal to graphene planes (about 30) and by a high shape anisotropy, defined as the ratio between the crystallites size in directions orthogonal and parallel to the layers.\(^\text{72}\) The interaction of SP and HSAG was promoted by mechanical and thermal energy. Adducts were studied by means of elemental, thermogravimetric (TGA), infrared (IR), wide angle X-ray diffraction (WAXD), transmission electron microscopy (TEM) analysis and Raman spectroscopy. Dispersions of HSAG–SP adducts in water were studied through dynamic light scattering (DLS) and UV spectroscopy. HSAG–SP adducts isolated through centrifugation of supernatant solutions were analyzed through high resolution TEM (HR-TEM). Electrical conductivity of HSAG–SP adducts was investigated on sheets deposited on paper.

2. Experimental

2.1. Materials

Reagents and solvents commercially available were purchased and used without further purification: 2,5-hexanedione (Merck-Schuchardt), 2-amino-1,3-propanediol (kindly provided by Bracco), acetone (Aldrich), high surface area graphite (HSAG) with Synthetic Graphite 8427® as trademark was purchased from Asbury Graphite Mills Inc., with a minimum carbon mass % of 99.5, a surface area of 330 m\(^2\) g\(^{-1}\), the number of stacked layers in the orthogonal direction to graphitic planes is about 35.\(^\text{72}\)

2.2. Synthesis of 2-(2,5-dimethyl-1H-pyrrol-1-yl)-1,3-propanediol (SP)

2-(2,5-Dimethyl-1H-pyrrol-1-yl)-1,3-propanediol was prepared following the reaction scheme shown in Scheme 1.\(^\text{67}\)

A mixture of hexan-2,5-dione (41.4 g; 0.36 mol) and serinol (30.0 g; 0.33 mol) was poured into a 100 mL round bottomed flask equipped with magnetic stirrer. The mixture was then stirred, at room temperature, for 6 h. The resulting compound 4a,6a-dimethyl-hexahydro-1,4-dioxa-6b-azacyclopenta[cd]pentalene (HHP) was characterized through \(^1\)H NMR and the yield was estimated to be 99%. \(^1\)H NMR (400 MHz, DMSO-\(d_6\), \(\delta\) in ppm): 1.28 (s, 6H); 1.77 (m, 2H); 1.93 (m, 2H); 3.60 (m, 4H); 3.94 (q, 1H).\(^\text{67,68}\)

The product mixture obtained from the synthesis of HHP was kept under vacuum for 2 h and then heated to 180 °C for 50 min. After distillation under reduced pressure at 130 °C and 0.1 mbar, SP was isolated as yellow oil with 96% yield (see Fig. 1 and Scheme 1). The global yield of the two step synthesis was therefore about 95%. \(^1\)H NMR (400 MHz, DMSO-\(d_6\), \(\delta\) in ppm): 2.16 (s, 6H, –CH\(_3\) at C-2,5 of pyrrole moiety); 3.63 (m, 2H, CH\(_2\)OH); 3.76 (m, 2H, CH\(_2\)OH); 4.10 (quintet, 1H, at C-3 of diol); 4.73 (t, 2H, CH\(_2\)OH); 5.55 (s, 2H, C-3,4 of pyrrole moiety).\(^\text{67,68}\)

2.3. Synthesis of HSAG–SP adducts

HSAG–SP adducts were prepared by reacting HSAG and SP with the help of either mechanical (HSAG–SP-M) or thermal (HSAG–SP-T) energy.

2.3.1. Synthesis of HSAG–SP-M adduct. In a 100 mL round bottomed flask was put in sequence graphite (5 g, 66 mmol) and acetone (15 mL). The suspension was sonicated for 15 min, using

![Scheme 1](image-url)  Reaction pathway for the preparation of the serinol derivatives HHP and SP.
a 2 L ultrasonic bath. After this time, a solution of SP (5.87 g, 66 mmol) in acetone (15 mL) was added. The resulting suspension was sonicated for 15 min. The solvent was removed under reduced pressure. The black powder of graphite/SP was treated using a planetary ball mill S100 from Retsch, having the grinding jar moving in a horizontal plane, with a volume of 0.3 L. The grinding jar was loaded with 6 ceramic balls having a diameter of 20 mm. 10.87 g of graphite/SP adduct were put into the jar, that was allowed to rotate at 300 rpm, at room temperature, for 6 h. After this time, the mixture was placed in a Büchner funnel with a sintered glass disc. It was repeatedly washed with distilled water (6 × 100 mL) obtaining 9.8 g of black powder.

2.3.2. Synthesis of HSAG–SP-T adduct. In a 100 mL round bottom flask was put in sequence graphite (5 g, 66 mmol) and acetone (15 mL). The suspension was sonicated for 15 min, using a 2 L ultrasonic bath. After this time, a solution of SP (1.116 g, 6.6 mmol) in acetone (5 mL) was added. The resulting suspension was sonicated for 15 min. The solvent was removed under reduced pressure. The black powder of graphite/SP (6.10 g) was poured into a 100 mL round bottomed flask equipped with magnetic stirrer and was heated at 130 °C for 6 h. After this time, the mixture was placed in a Büchner funnel with a sintered glass disc. It was repeatedly washed with distilled water (3 × 20 mL), obtaining 5.86 g of black powder.

2.4. Characterization of HSAG–SP adducts

2.4.1. Thermogravimetric analysis. TGA tests under flowing N2 (60 mL min⁻¹) were performed with a Mettler TGA SDTA/851 instrument according to the standard method ISO9924-1. Samples (10 mg) were heated from 30 to 300 °C at 10 °C min⁻¹, kept at 300 °C for 10 min, and then heated up to 550 °C at 20 °C min⁻¹. After being maintained at 550 °C for 15 min, they were further heated up to 900 °C and kept at 900 °C for 30 min under flowing air (60 mL min⁻¹).

2.4.2. Elemental analysis. Elemental analysis was performed with a Thermo FlashEA 1112 Series CHNS-O analyzer, after pretreating samples in an oven at 100 °C for 12 h.

2.4.3. FT-IR. FT-IR spectra were recorded between 450 and 4000 cm⁻¹ by using a Perkin-Elmer FT-IR Spectrum One equipped with Universal ATR Sampling Accessory with diamond crystal.

2.4.4. Raman spectroscopy. Raman spectra of powder samples deposited on a glass slide were recorded by using a Horiba Jobin Yvon Labram HR800 dispersive Raman spectrometer equipped with Olympus BX41 microscope and a 50× objective. The excitation line at 632.8 nm of a He/Ne laser was kept at 0.5 mW in order to prevent possible samples degradation. The spectra were obtained as the average of four acquisitions (scan time: 30 s for each acquisition) with a spectral resolution of 2 cm⁻¹.

2.4.5. Wide angle X-ray diffraction. Wide-angle X-ray diffraction (WAXD) patterns were obtained in reflection, with an automatic Bruker D8 Advance diffractometer, with nickel filtered Cu-Kα radiation. Patterns were recorded in 10–100° as the 2θ range, being 2θ the peak diffraction angle. Distance between crystallographic planes was calculated from the Bragg law. The Dhk correlation length, in the direction perpendicular to the hkl crystal graphitic planes, was determined applying the Scherrer equation.

\[
D_{hk\ell} = K\beta_{hk\ell} \cos \theta_{hk\ell} / b
\]

where: K is the Scherrer constant, λ is the wavelength of the irradiating beam (1.5419 Å, Cu-Kα), βhk\ell is the width at half height, and θhk\ell is the diffraction angle. The instrumental broadening, b, was determined by obtaining a WAXD pattern of a standard silicon powder 325 mesh (99%), under the same experimental conditions. The width at half height, \(β_{hk\ell} = (B_{hk\ell} - b)\), was corrected, for each observed reflection with \(β_{hk\ell} < 1°\), by subtracting the instrumental broadening of the closest silicon reflection from the experimental width at half height, \(B_{hk\ell}\).

2.5. Water solutions of HSAG–SP adducts

2.5.1. Preparation. Water solutions of HSAG–SP adducts at different concentrations (1 mg mL⁻¹; 0.5 mg mL⁻¹; 0.3 mg mL⁻¹; 0.1 mg mL⁻¹) were prepared. Each solution was sonicated for 10 min using an ultrasonic bath (260 W) and subsequently UV-vis absorption was measured. The solution (10 mL) of each sample was put in a Falcon™ 15 mL Conical Centrifuge Tubes and centrifuged at: 2000 rpm for 10 minutes and at 9000 rpm for 5, 30, 60 and 90 minutes. Supernatants obtained after each centrifugation processes were removed and analyzed. UV-vis absorptions and DLS analysis were measured immediately after each centrifugation and after 1 week stored.

2.5.2. UV-vis spectroscopy. The suspensions of adduct (3 mL) were placed by pipette Pasteur, in quartz cuvettes of 1 cm optical path (volume 1 or 3 mL) and analyzed by a spectrophotometer Hewlett Packard 8452A Diode Array Spectrophotometer. Resets the instrument with pure solvent and has one UV spectrum from 200–340 nm. It was recorded a white the solvent employed. The UV-visible spectrum reported intensity the absorption as a function of the wavelength of the radiation between 200 and 750 nm.

2.5.3. Dynamic light scattering (DLS). The nanoparticle size in HSAG–SP solutions was analyzed with a Zetasizer Dynamic Light Scattering system (Malvern Instrument Ltd) at room temperature using a 1 ¼ 632.8 nm He–Ne laser. Each centrifuged solution, prepared using a concentration of 1 mg mL⁻¹, was analyzed by DLS: 3 mL of solution were placed using a pipette Pasteur, in quartz cuvettes of 1 cm optical path and analyzed immediately after centrifugation and after 10, 60 and 100 min of storage.

2.5.4. Transmission electron microscopy. Water solutions (1 mg mL⁻¹) of adducts were centrifuged using an ALC-Centrifuge 4206. TEM analysis of supernatants was performed with a Zeiss EM900 microscope operating at an accelerating voltage of 80 kV. Few drops of acetone diluted suspension of the sample were deposited on a carbon film supported on a standard Cu grid and air-dried for several hours before analysis.

2.5.5. High-resolution transmission electron microscopy (HR-TEM). HR-TEM investigations on adducts’ samples, isolated from the supernatant solutions after centrifugation, were carried out with a Philips CM 200 field emission gun.
microscope operating at an accelerating voltage of 200 kV. Few drops of the water solutions were deposited on 200 mesh lacey carbon-coated copper grid and air-dried for several hours before analysis. During acquisition of HR-TEM images, the samples did not undergo structural transformation. Low beam current densities and short acquisition times were adopted. To estimate the number of stacked graphene layers and the dimensions of the stacks visible in HR-TEM micrographs, the Gatan Digital Micrograph software was used.

3. Results and discussion

The synthesis of SP, summarized in Scheme 1, is described in detail in the experimental part and has been discussed elsewhere. It is worth underlining the high atomeconomy of the reaction (82%), with water as the only co-product of the first step. The high yield, about 95%, was easily reproduced in all the preparations performed in this research. This led to the atomic efficiency of about 78% for the preparation of the SP sample used in this work.

The interaction of HSAG with SP was promoted with the help of mechanical or thermal energy, as described in the experimental part. The adopted procedures are summarized in the block diagram of Fig. 2.

In brief, HSAG and SP were first premixed. HSAG–SP mixture was either ball milled in a jar for 6 h or heated in a flask for 6 hours at 130 °C. The products taken either from the jar or from the flask were washed several times with water, until SP was undetectable in the washing water. The HSAG/SP molar amount (in the case of HSAG, the moles of aromatic rings were considered) was explored in the range from 2 : 1 to 10 : 1. Adducts prepared by using either mechanical or thermal energy were named HSAG–SP-M and HSAG–SP-T, respectively. In the present work, were characterized HSAG–SP-M and HSAG–SP-T samples having almost the same amount of SP, determined from thermogravimetric and elemental analysis.

TGA analysis was performed under nitrogen on HSAG, SP and on the HSAG–SP adducts. Thermographs are shown in Fig. 3. In Table 1, are shown data of mass loss for HSAG and for the adducts.

SP achieves its boiling point at about 230 °C. Curve of HSAG shows decomposition profile made by two main steps, that can be attributed to: decomposition of oxygen-containing and alkenylic groups for T in the range from 150 °C to 700 °C and combustion with oxygen (see experimental part) at T > 700 °C.

Fig. 2 Block diagram for the preparation of HSAG–SP adducts: with the help of mechanical (HSAG–SP-M) or thermal energy (HSAG–SP-T).

Fig. 3 TGA curves of HSAG (a), SP (b), HSAG–SP-T (c), heated HSAG–SP-M (d).
Mass loss due to water removal, that could be expected at $T < 150 \degree C$, can not be detected.

Larger mass loss in the temperature range from 150 to 500 \degree C is shown by HSAG–SP–T and HSAG–SP–M. Degradation occurs at a temperature much higher than the one observed in the SP curve, indicating that residual SP is not present on HSAG. Both the adducts reveal an appreciable mass loss at temperatures below 150 \degree C. Interestingly, the larger amount of mass loss at $T > 150 \degree C$ is accompanied by larger mass loss at $T < 150 \degree C$. It could be hypothesized that HSAG–SP adducts that contain larger amount of oxygenated functional groups hold larger amount of water. The onset temperature of the last step degradation is pretty similar for HSAG and for the adducts, indicating that the reaction with SP did not lead to lower thermal stability of HSAG.

As reported in the Experimental part, pristine HSAG had 99.5% as carbon content. The nitrogen content of HSAG–SP adducts was determined by means of elemental analysis. Table 2 shows the experimental values and the amount of SP, theoretically estimated.

The adducts contain comparable amount of SP, 13% and 14% by mass. HSAG–SP–T was obtained by adopting 10/1 as the HSAG/SP molar ratio, whereas 2/1 as molar ratio was used for the preparation of HSAG–SP–M. Studies have not been performed on the yield of the reaction between HSAG and SP as a function of the preparation procedure. As mentioned above, samples discussed in this work were selected as they show comparable amount of SP.

Functional groups present on HSAG–SP adducts were investigated by means of IR spectroscopy. Fig. 4 shows the IR spectra of pristine HSAG (a), SP (b) and of HSAG/SP mixture (HSAG/SP = 2/1 as molar ratio) (c). Such mixture was prepared by simply mixing HSAG and SP, without giving either thermal or mechanical energy. Spectra of HSAG–SP–T and HSAG–SP–M adducts are in Fig. 4d and in Fig. 4e, respectively.

In the SP spectrum (Fig. 4b), band in the region from 3370 cm$^{-1}$ to 3330 cm$^{-1}$ can be reasonably attributed to hydrogen bonded OH groups, the pyrrole ring is evidenced by the collective vibration mode of C=C/C–C–C stretches, located at about 1530 cm$^{-1}$ and by other bands at 1395 and 1490 cm$^{-1}$, related to vibrational modes involving the pyrrole unit. The above bands due to pyrrole ring can be taken as fingerprint of SP. These bands are still clearly detectable in the spectrum of the mixture (Fig. 4c) where they do not show frequency shifts with respect to pristine SP (Fig. 4b). Spectra of both washed adducts (Fig. 4d and e) show bands that cannot be attributed to HSAG. In particular, they are: bands in the region of sp$^3$ CH stretching at about 2900 cm$^{-1}$ and features in the region characteristic of the C–C stretches of aromatic rings, namely the bands at 1590 and 1470 cm$^{-1}$, peaks at 1383 cm$^{-1}$ and 956 cm$^{-1}$ which could be attributable to vibrations involving the diole function and the alkenyl groups of reacted SP molecules. These findings confirm what already observed through elemental and TGA analysis, that means the formation of a stable HSAG–SP adduct. Moreover, they also seem to suggest the occurring of a reaction or at least of an interaction able to modify the nature and/or the surrounding of the chemical bonds in the pyrrole ring of SP. This aspect is under investigation. As working hypothesis, Diels–Alder reaction between dangling double bonds of graphene layers and the pyrrole rings is taken into consideration.

Organization at the solid state of HSAG and of HSAG–SP adducts was investigated by means of WAXD. Fig. 5 shows WAXD patterns of HSAG (a), HSAG–SP–M (b), HSAG–SP–T (c).

Assignment of reflections due to pristine HSAG is shown in Fig. 5a. Patterns of HSAG–SP–T and HSAG–SP–M reveal that the (002) reflection remains at the same 2$\theta$ value as in the pattern of pristine HSAG. This finding indicates that SP was not intercalated in the interlayer space. The number of stacked layers was calculated by applying the Scherrer equation (see experimental section) to the (002) reflection. Such number was found to be: 35 for HSAG, 24 for HSAG–SP–M, and 29 for HSAG–SP–T. It

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**Table 1** Mass loss for HSAG, HSAG–SP–M and HSAG–SP–T, from TGA analysis$^a$

<table>
<thead>
<tr>
<th>Sample</th>
<th>$T &lt; 150 \degree C$</th>
<th>$&lt; T &lt; 300 \degree C$</th>
<th>$&lt; T &lt; 500 \degree C$</th>
</tr>
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<tbody>
<tr>
<td>HSAG</td>
<td>0</td>
<td>0</td>
<td>3.1</td>
</tr>
<tr>
<td>HSAG–SP–M</td>
<td>0.8</td>
<td>2.7</td>
<td>7.8</td>
</tr>
<tr>
<td>HSAG–SP–T</td>
<td>0.8</td>
<td>2.7</td>
<td>7.7</td>
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</table>

$^a$ Residual mass loss was above 700 \degree C. Total mass loss was 100%.

**Table 2** Mass% of nitrogen and of SP

<table>
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<tr>
<th>Adduct</th>
<th>Mass% of Nitrogen</th>
<th>Mass% of SP</th>
</tr>
</thead>
<tbody>
<tr>
<td>HSAG–SP–T</td>
<td>1.96</td>
<td>14.03</td>
</tr>
<tr>
<td>HSAG–SP–M</td>
<td>1.97</td>
<td>13.00</td>
</tr>
</tbody>
</table>

$^a$ From elemental analysis. $^b$ Calculated on the basis of the nitrogen content.

**Fig. 4** IR spectra of HSAG (a), SP (b), HSAG/SP = 2/1 (c), HSAG–SP–M adduct (d), heated HSAG–SP–T adduct (e).
appears that the functionalization reaction promotes the reduction of the crystalline order in the direction orthogonal to the graphene layers. 100 and 110 reflections indicate the in plane correlation lengths. The in plane order was estimated by applying the Scherrer equation to the 110 reflection: the correlation length was found, for HSAG and for the adducts, in a range from 26.5 nm to 28 nm. This result seems to suggest that the reaction of HSAG with SP, also via ball milling, does not substantially alter the in plane order.

Structural characterization of HSAG and HSAG-SP adducts was performed via Raman spectroscopy, a tool widely employed for the study of carbonaceous materials. In Raman spectrum, two lines, named D and G and located at 1350 cm\(^{-1}\) and 1590 cm\(^{-1}\) respectively, reveal the presence of graphitic sp\(^2\)-phase. In particular, the G peak is characteristic of bulk crystalline graphite (graphene), whereas the D peak appears in the presence of either disorder or confinement (e.g. by edges) of the graphitic layers. Structural defects can be holes, sp\(^3\) or sp carbon atoms, free radicals, distortions from planarity, graftered functional groups. Moreover, it is worth reminding that graphitic layers have finite dimensions and irregular boundaries with, for example, dangling bonds. All these defects give rise to Raman scattering in the D region with a frequency that depends on the excitation wavelength. Some of the authors performed ball milling of graphitic powders, in order to vary the average crystallite size and the amount of defects. Raman spectrum showed evident D line, whose intensity increased with the grinding time. Fig. 6 shows Raman spectra of HSAG (Fig. 6a) HSAG-SP-T (Fig. 6b) and HSAG-SP-M (Fig. 6c). Analysis was performed on a good number of spots and spectra can be considered representative of the investigated samples.

In all the spectra, G and D bands are present, with similar intensity. Interestingly, larger intensity can be observed for the D band in the spectrum of the adduct prepared through ball milling, whereas relative intensity of G and D peak appears substantially unaltered after the thermal treatment of HSAG with SP. Hence, there are not indications that the reaction of HSAG with SP appreciably alter per se the structure of the graphitic layers, whereas the milling process appears to play the major role. The interpretation proposed by some of the authors to comment Raman spectra of ball milled graphite can be applied also in the present work. Graphitic samples have a core sufficiently far from the edge, with electronic and vibrational properties that can not be distinguished from those of infinite and ideal graphene layers. A confined crown region close to the edge is electronically perturbed and local electronic and vibrational structures, affected by confinement effects, are different from those of the bulk materials. At the very edges, many types of molecular disorder are possible, making the system different from the ideal sp\(^2\) carbon layer. In consideration of findings from WAXD analysis, the enhancement of the D band in the Raman spectrum of HSAG-SP-M could be prevalently attributed, at least as working hypothesis, to larger amounts of defects in the crown region and at the edges.

On the basis of the results discussed so far and in the light of what available in the scientific literature, it seems possible to comment that HSAG-SP adducts are nano-stacks of confined graphene, with peripheral functional groups containing oxygen atoms, in particular hydroxyl groups.

As it should be expected, considering the presence of such functional groups, water dispersions of HSAG-SP were easily prepared (as described in the experimental part). Such dispersions were analyzed by UV-vis absorption. Dispersions freshly prepared and after 30 min centrifugation at 2000 rpm of HSAG-SP-M adducts are shown in Fig. 7 (Fig. 7b and c, respectively), together with a suspension of pristine HSAG (Fig. 7a). Decantation can not be observed for the dispersions based on HSAG-SP adducts.

UV-vis absorption analysis was performed on water dispersions of HSAG-SP-M adduct with 0.1, 0.3, 0.5 and 1
mg mL$^{-1}$ as adduct concentration. In Fig. 8a it is shown that the absorbance monotonously increases with the adduct concentration. Plot of UV-vis absorbance as a function of the adduct concentration (not reported in this manuscript) reveals a linear correlation, suggesting that water solutions of HSAG–SP-M were prepared. The stability of the solution with 1 mg mL$^{-1}$ as the adduct concentration was investigated by taking UV-vis spectra on a samples stored for 1 week and on samples centrifuged at 9000 rpm for 30 minutes and 1 hour respectively. In Fig. 8b, it is evident that the same absorbance was measured for the suspensions freshly prepared and stored for 1 week, confirming that the adduct did not settle down, even after long time. Moreover, it is possible to observe that only the centrifugation at 9000 rpm led to appreciable reduction of absorbance, that nonetheless remains at a remarkable level.

Average size of particles of HSAG and HSAG–SP adducts, in water dispersions, was determined by means of DLS (as reported in the experimental part). Table 3 shows data for as prepared dispersions and for the supernatant dispersions after 30 min centrifugation at 9000 rpm.

Values obtained by DLS measurements represent the hydrodynamic radius of HSAG particles in the dispersions. It appears that particles of HSAG–SP-M have lower average size than particles of HSAG. The average size further decreases in the supernatant suspension. On the contrary, the average size appears to increase, with respect to HSAG, when the HSAG–SP adduct is prepared by heating. Also the average size in the supernatant suspension is larger than in HSAG. It seems thus that the functional groups can favour the aggregation of HSAG particles that experience thermal treatment. The average size of HSAG–SP-M aggregates in solution was investigated at different times ($t = 0, 10, 60, 100$ min), after centrifugation at 9000 rpm for 60 min. It was found to be about 130 nm at $t = 0$ and to remain constant for about 10 nm, achieving a plateau value of about 200 nm after 20 min. This finding seems to confirm the tendency to aggregate of HSAG–SP adducts. The dependence of the particles average size on the centrifugation time was determined for the HSAG–SP-M adduct: after 5, 30, 60 and 90 min of centrifugation, 260 nm, 150 nm, 135 nm and 120 nm were respectively measured.

TEM analysis of supernatant suspensions of HSAG, HSAG–SP-T and HSAG–SP-M adducts, after 60 min of centrifugation, was performed after their deposition on a carbon film. Analysis was repeated on significant number of samples. Representative TEM micrographs are shown in Fig. 9. HSAG in Fig. 9a is characterized by graphite stacks randomly arranged, with lateral size that appears lower than that of HSAG–SP-T (Fig. 9b) and HSAG–SP-M (Fig. 9c). This is probably due to the fact that only HSAG aggregates of small size can be

![Fig. 7 Water dispersions of HSAG (a), HSAG–SP-M: freshly prepared (b), after 30 min centrifugation at 2000 rpm (c).](image)

![Fig. 8 Dependence of UV-vis absorbance on concentration of HSAG–SP-M adduct in water (a) and after centrifugation at 9000 rpm (b).](image)

**Table 3** Average size of HSAG, HSAG–SP-T and HSAG–SP-M in water dispersions, determined through DLS

<table>
<thead>
<tr>
<th>Starting dispersion</th>
<th>Supernatant dispersion$^a$</th>
</tr>
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<tbody>
<tr>
<td>HSAG</td>
<td>500</td>
</tr>
<tr>
<td>HSAG–SP-M</td>
<td>300</td>
</tr>
<tr>
<td>HSAG–SP-T</td>
<td>1500</td>
</tr>
</tbody>
</table>

$^a$ After 30 minutes centrifugation at 9000 rpm.
suspended in water. Micrographs of both adducts revealed stacks apparently made by a low number of layers in the case of HSAG–SP-T (Fig. 9b) and by very few layers in the case of HSAG–SP-M (Fig. 9c). These findings suggest that the milling procedure promoted larger exfoliation of HSAG–SP adducts and, as a consequence, the formation of modified stacks of graphene layers that found it easier to remain dispersed in water, with respect to the stacks of HSAG–SP-T adducts. The difference among the average size of graphene stacks determined by DLS and the lateral size of the sheets observed in TEM micrographs seems to suggest the extreme flexibility of HSAG–SP adducts.

The structure of HSAG–SP-M adduct was investigated by performing HR-TEM analysis on samples isolated from the supernatant solutions, after centrifugation for 10 min at 2000 rpm and for 5 and 60 min at 9000 rpm. Fig. 10 shows HR-TEM micrographs at lower and higher magnifications.

Micrographs at lower magnification in Fig. 10a, c and e reveal that the lateral size of HSAG–SP-M adducts is of the same order of magnitude in samples isolated after centrifugations at different rpm and for different times. This indicates that the milling step does not cause appreciable breaking of the graphitic layers. Moreover, it appears that the SP modifier is able to bring into water solution also nano-graphites with pretty large lateral size. Micrographs at higher magnification allow to visualize the stacks of graphene layers that are disposed with a lateral side perpendicular to the beam. Thanks to this disposition of the nano-stacks, it is possible to estimate the number of stacked graphene layers. Fig. 10b shows stacks isolated after centrifugation for 10 min at 2000 rpm: are visible stacks (indicated in the boxes) of about 3.5–4.2 nm, made by about 10–12 stacked graphene layers. As revealed by the exam of a good number of micrographs, such stacks are the most abundant ones in a population of stacks that contain either little larger or little lower number of layers. Fig. 10d shows a stack (indicated in the box) isolated after centrifugation for 5 min at 9000 rpm: is of about 2.8 nm and contains about 8 stacked graphene layers. Such a stack was very frequently observed in the analyzed micrographs after centrifugation for 5 min. In Fig. 10f it is shown the adduct isolated after centrifugation at 9000 rpm for 60 min. It seems that a layer of organic substance (indicated by the arrow) is tightly adhered to the carbon allotrope: voids can not be observed between the carbon surface and the organic substance. It appears that the organic substance, that has to be assumed based on SP, is also on the graphene face. The fractionation of HSAG–SP-M adduct, as a function of centrifugation, appears to be due to the different numbers of stacked layers, rather than to different lateral size, and to the different amount of SP modifier.

It was seen in HR-TEM micrograph in Fig. 10b that the graphene nano-stacks are mostly made by about 10 graphene layers, that means quite a lower number with respect to that estimated on pristine HSAG via WAXD analysis (about 30). These findings allow to comment that reaction of HSAG with SP, through ball milling, and the successive centrifugation of water solutions of the HSAG–SP adduct lead to isolate nano-stacks

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**Fig. 9** TEM micrographs of water supernatant after 60 min centrifugation at 9000 rpm: HSAG (a), HSAG–SP-T (b) and HSAG–SP-M (c).

**Fig. 10** Micrographs of HSAG–SP-M adducts isolated from supernatant solutions, after centrifugation for 10 min at 2000 rpm (a and b), 5 min at 9000 rpm (c and d) and for 60 min at 9000 rpm. Micrographs are: low magnification bright field TEM (a, c and e), HR-TEM images (b, d and f).
made by few layers of graphene, whose number appears to depend on the adopted experimental conditions.

4. Conclusions

The neat reaction of serinol with 2,5-hexanedione led to the preparation of 2-(2,5-dimethyl-1H-pyrrol-1-yl)-1,3-propanediol, with yield of about 95% and atomic efficiency of about 78%.

SP, that contains pyrrole ring, was able to form stable adducts with HSAG, a nano-graphite with high surface area and high shape anisotropy. Preparation of HSAG–SP adducts was carried out in the absence of organic solvents or catalysts, assisted by mechanical energy in a ball milling jar or by thermal energy, by simply heating the reaction flask. Adducts with 13–14% by mass of SP were prepared.

Functionalization reaction was successful. More than 10% by mass of non graphitic, nitrogen containing substance was detected by TGA. Functional groups, that could be attributed to SP, were revealed by IR analysis. After the reaction with SP, the same interlayer distance between graphitic layers and the same in plane order were maintained, suggesting that functionalization occurred on the edges and/or on the faces of graphite aggregates. The presence in IR spectrum of peaks other than those due to SP and HSAG leads to hypothesize that a chemical reaction occurred, with the formation of covalent bonds on the adduct, that could justify the stability of the adducts. Raman spectra of pristine HSAG and HSAG–SP adducts showed evident G and D bands. Their relative intensity was not substantially altered after the thermal reaction of HSAG with SP, suggesting that such reaction does not lead to appreciable structural modification of the graphene layers. This finding is in line with what revealed by WAXD analysis. Instead, the D band became prevailing after the ball milling treatment, that appears to play the major role in affecting the structure of the carbon allotrope.

Water solutions were prepared with HSAG–SP-M, from 0.1 to 1 mg mL⁻¹ as adduct concentration. Such solutions, whose UV absorbance followed the Lambert–Beer law, were stable for weeks at rest. Centrifugations were performed at different rpm and for different times, isolating HSAG–SP-M samples from the supernatant solutions. HR-TEM analysis showed that the lateral size of graphitic flakes was very similar in all the samples, whereas the number of stacked layers decreased down to at least about 8 by increasing the rpm (9000), even at short time (5 min) of centrifugation. The HR-TEM picture of HSAG–SP-M adduct isolated at 9000 rpm for long centrifugation time revealed that an organic layer, reasonably based on SP, was tightly adhered to the carbon surface.

The HSAG–SP adducts prepared in this work are nano-stacks of confined graphene layers, with oxygen and nitrogen containing functional groups that do not appreciably alter the solid state organization of the carbon allotrope. Thanks to such functional groups, water solutions of the graphene nanostacks can be easily formed. Moreover, centrifugation technique allow to isolate, from the supernatant solutions, HSAG–SP samples with few graphene layers.

SP is a Janus molecules and appears suitable to form stable adducts with carbon allotropes, such as HSAG, and to promote the separation of graphitic aggregates into stacks containing a low/very low number of graphene layers. Results here reported pave the way for the easy functionalization of carbon allotropes. Moreover, the easy preparation of stable dispersions in polar media allows to envisage a large variety of applications.

Acknowledgements

Dr Valeria Cipolletti (Politecnico Milano) is gratefully acknowledged for X-ray analysis and “PRIN Research Project 2010–2011” for the financial support.

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