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From biowaste to magnet-responsive materials for water remediation from polycyclic aromatic hydrocarbons

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| 20 | Abstract |
| 21 | Composted urban biowaste-derived substances (BBS-GC) are used as carbon sources for the |

preparation of carbon-coated magnet-sensitive nanoparticles obtained via co-precipitation method 22 23 and the subsequent thermal treatment at 550 °C under nitrogen atmosphere. A multitechnique approach has been applied to investigate the morphology, magnetic properties, phase composition, 24 thermal stability of the obtained magnet-sensitive materials. In particular, pyrolysis-induced 25 modifications affecting the BBS-GC/carbon shell were highlighted. The adsorption capacity of such 26 bio-derivative magnetic materials for the removal of hydrophobic contaminants such as polycyclic 27 aromatic hydrocarbons was evaluated in order to verify their potential application in wastewater 28 remediation process. The promising results suggest their use as a new generation of magnet-29 responsive easily-recoverable adsorbents for water purification treatments. 30

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32 Keywords: Adsorption; Biomass valorization; Iron oxides; Magnetic nanomaterials; Pyrolysis.

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34 **1. Introduction**

In recent years, a wide number of studies has been dedicated to the valorization of biowaste and 35 biowaste-derived substances in order to prepare novel smart products, thus reaching a reduction of 36 the industrial and/or urban residues, re-entering them in the economic cycle in a more sustainable 37 and environmental-friendly development of our society (Fava et al., 2015; G. Magnacca et al., 38 2015; Deng et al., 2016; Nisticò, 2017a). In particular, low-cost starting materials from natural 39 sources (i.e., clavs and minerals as well as biopolymers), industrial wastes and agriculture residues 40 (such as lignin and humic/fulvic acid-like substances, chitin and chitosan, alginate and waste 41 slurry), are considered as an important source of fine chemicals in substitution of the more 42 traditional petroleum-derived ones (Bailey et al., 1999; Avetta et al., 2014; Vakili et al., 2014; 43 Bautista et al., 2015; Choi et al., 2015a; Choi et al., 2015b; Manna et al., 2015; Nisticò et al., 2015; 44 Franzoso et al., 2016). Among these low-cost bio-based sources, lignocellulose-derived materials 45 and chitin/chitosan seem to be highly attractive resources for the future generation of chemicals 46 47 (Tran et al., 2015).

Shape- and size-controlled smart magnetic nanoparticles (mNPs) consisting in a magnet-sensitive 48 49 core made of iron oxides (i.e., magnetite/maghemite) stabilized by one or more covering layer(s), currently attracted the interests of worldwide experts due to their unique magnet-responsive 50 properties which favor their uses in several technological applications, such as drug-delivery or 51 contaminated wastewater (bio)remediation processes (Cesano et al., 2016; Lata and Samadder, 52 2016; Zhao et al., 2016; Lu et al., 2007; Nisticò, 2017b). In this context, various protective and/or 53 stabilizing materials were investigated for inducing particular surface properties at the mNPs. 54 Typically, both oxides (mainly silica) and/or organic (surfactants, (bio)polymers, and carbons) 55 coatings were considered as possible covering-materials, thus combining the advantages of both 56 components selected (namely, the magnet-sensitive Fe-containing oxides core and the functionality 57 at the surface) (Lu et al., 2007; Li et al., 2013; Nardi et al., 2013). 58

As already evidenced in our previous works (Nisticò et al., 2015; Franzoso et al., 2017; Nisticò et 59 al., 2018), Bio-Based Substances (BBS-GC, where GC stands for green compost) are 60 supramolecular aggregates made of organic macromolecules with a highly branched and complex 61 lignin-derived structure. Such substances are isolated from urban composted biowastes after more 62 than 180 days of composting, and as a consequence of the starting material complexity (i.e., urban 63 public park trimming and home gardening residues). BBS-GC contain several functional groups 64 (the BBS-GC chemical structure and functionalities are fully reported in Table S1, whereas their 65 thermal stability is reported in Figure S1) (Nisticò et al., 2015). The macromolecular nature of 66 these BBS, very similar to humic/fulvic substances with several O-containing functional groups, 67 suggested their potential use as stabilizing/protective agents for the greener synthesis of mNPs with 68 both adsorption properties and enhanced photoactive response in advanced oxidation processes 69 (AOPs) for the remediation of contaminated water (Magnacca et al., 2014). 70

Polycyclic aromatic hydrocarbons (PAHs) are among the most ubiquitous environmental contaminants (Vane et al., 2013). Due to their resistance to biodegradation and bioaccumulation, the presence of the PAHs in surface water, groundwater, and drinking water, poses a major risk to human health and wildlife. Different techniques have been developed for removing PAHs from contaminated water, among which physical remediation by means of selective adsorbents is one of the most frequently used strategies due to its high effectiveness and low operational cost (Pérez et al., 2008; Dave and Ghaly, 2011).

Conventional adsorbents selected for their high oleophilic properties in contaminated water remediation are polymeric foams in the shape of rolls and/or sheets (typically made of polypropylene, polystyrene, and polyurethane) (Teas et al., 2001; Dave and Ghaly, 2011). Additionally, as reported in the literature, high-surface-area activated carbons and graphitic carbons can be a very interesting alternative as adsorbents in wastewater treatments (Lee et al., 2006; Cesano et al., 2012; Magnacca et al., 2018). Hu et al. (2014) demonstrated the use of compressible

carbon nanotube-graphene hybrid aerogels having hydrophobicity and oleophilicity for oil sorption.
However, the use of such materials, especially in open-space contaminated areas, presents some
limitations, one above all the difficulties in their recovering. Therefore, a step forward in this field
is the preparation of easily-recoverable carbon-coated magnet-sensitive materials (Wang et al.,
2006; Lu et al., 2007; Hao et al., 2014; Cesano et al., 2016).

Magnetic materials consisting of iron oxides (i.e magnetite/maghemite) particles stabilized by functional carbon shells, received a great attention as easily-recoverable magnet-responsive smart materials for wastewater remediation (Hao et al., 2014; Mehta et al., 2015). Recently, in our previous study (Nisticò et al., 2017), another biowaste-derived substrate (namely, chitosan) was selected as starting material for the preparation of carbon-coated magnet-sensitive composite materials for water remediation from PAHs, with promising results.

Therefore, the aim of this study is the preparation and physicochemical characterization of carbon-95 96 coated magnetite/maghemite NPs synthesized through iron salts co-precipitation process in the presence of BBS as initial stabilizers and carbon-source. Pyrolysis under nitrogen atmosphere is 97 98 carried out at mild condition (550 °C), in order to convert the BBS into a carbon layer without 99 altering the structure and properties of magnetite. Morphology, thermal stability and magnetic properties of such composites mNPs were fully assessed. Adsorption experiments involving the 100 water remediation from a PAHs mixture by direct contact with such BBS-derived carbon-materials 101 and subsequent easily-recovery by magnetic separation are also examined. 102

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104 2. Material and methods

105 2.1 Materials

106 Anhydrous ferric chloride FeCl_3 (CAS 7705-08-0, purity \geq 98%, Fluka Chemika) and ferrous 107 sulphate heptahydrate FeSO_4 ·7H₂O (CAS 7782-63-0, purity \geq 99.5%, Fluka Chemika) were 108 selected as iron oxide precursors. Bio-Based Substances (BBS-GC) were obtained following a

previously reported procedure (Nisticò et al., 2015) starting from urban composted biowastes (i.e., 109 urban public park trimming and home gardening residues after 180 days of composting) sampled 110 from the process lines of ACEA Pinerolese Industriale S.p.A. waste treatment plant in Pinerolo 111 (Italy) in an advanced system that comprises specific technological facilities, developed by ACEA 112 *Pinerolese Industriale S.p.A*, and under European validation. Naphthalene (NAP; $\log K_{ow} = 3.37$), 113 acenaphthene (AC; $\log K_{ow} = 3.98$), acenaphthylene (ACL; $\log K_{ow} = 4.07$), fluorine (FL; $\log K_{ow} =$ 114 4.18), anthracene (ANT; $\log K_{ow} = 4.46$), phenanthrene (PHE; $\log K_{ow} = 4.50$), pyrene (PY; $\log K_{ow} =$ 115 4.88) and fluoranthene (FN; log $K_{ow} = 4.90$) were purchased from Sigma-Aldrich (purity > 99.0%). 116 The aqueous solutions for adsorption experiments were prepared using ultrapure water Millipore 117 Milli-QTM and all chemicals were used without further purification. 118

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120 **2.2 Preparation of BBS-derived carbon magnet-sensitive materials**

121 The starting BBS-stabilized magnetite NPs were synthesized following the procedure reported in the literature (Franzoso et al., 2017; Nisticò et al., 2018). In detail, 3.7 g of FeCl₃ and 4.17 g of 122 $FeSO_4$ ·7H₂O (molar ratio Fe(III)/Fe(II) = 1.5) were dissolved in 100 mL of deionized water and 123 heated up to 90 °C. Once the temperature was reached, two solutions were added in sequence: a) 10 124 mL of 25 vol.% ammonium hydroxide, and b) 50 mL of a previously prepared BBS aqueous 125 solution. Three solutions with different content of BBS were employed: 1, 2, and 3 wt.%, 126 respectively. The mixture was mechanically stirred at 90 °C for 30 min and then cooled down to 127 room temperature (RT). Such dispersions of BBS-stabilized magnetite NPs were purified by 128 washing twice with deionized water, deposited onto glass Petri dishes, and oven-dried at 80 °C 129 overnight. BBS-derived magnetic-materials were then thermally treated in a quartz tube under 130 nitrogen atmosphere (N₂ flux of 250 mL min⁻¹) for 1 h at 550 °C (from RT to the target 131 temperature, heating ramp 5 °C min⁻¹). The resulting materials were manually crumbled in a mortar 132 for the further experiments. In analogy to our previous work (Franzoso et al., 2017), depending on 133

the different BBS amount (1, 2 and 3 wt.%), the obtained NPs were coded with the acronym MB1, MB2 and MB3, respectively. Thermally-treated samples were further coded with pyT, where py stands for pyrolysis, and T refers to the temperature of pyrolysis (550 °C). Magnetite (M0), as obtained following the co-precipitation procedure without the BBS addition, was taken as a neat magnetite/maghemite reference.

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140 **2.3 Physicochemical characterization**

Scanning electron microscopy (SEM) analyses were carried out by using a ZEISS EVO 50 XVP 141 microscope with LaB_6 source, equipped with detectors for secondary electron collection and energy 142 dispersive X-ray Spectroscopy (EDS). SEM micrographs were obtained after sputtering samples 143 with a 10-15 nm thick gold film. Atomic force microscopy (AFM) and magnetic force microscopy 144 (MFM) measurements were performed by using a modified Nanosurf Easyscan2 AFM instrument, 145 equipped with a 10 µm scan-head, high performance anti-vibration platform in an acoustically 146 insulated enclosure and Faraday cage. AFM and MFM images were obtained in a dual-pass mode: 147 the first scan was for the morphological imaging in the intermittent contact mode, while the second 148 149 scan was operated at constant-height (H) above the same surface with a magnetic probe (Multi75M-G, Budget Sensors; resonant frequency = 75 kHz, force constant = 3 N m^{-1}) having a tip radius of 150 about 60 nm by monitoring the shifting of the phase and of the amplitude signals. The magnetic tip 151 was magnetized by an external magnet (magnetization along the tip-axis) and tested on a magnetic 152 grid prior to measurements. Magnetization measurements were carried out performing experiments 153 in a LakeShore 7404 vibrating sample magnetometer. The hysteresis loop of the samples was 154 registered at RT cycling the magnetic field from -20000 to 20000 Oe. X-ray diffraction (XRD) 155 patterns were obtained by using an X'Pert PRO MPD diffractometer from PANalytical, equipped 156 with Cu anode, working at 40 kV and 30 mA, with a Bragg-Brentano geometry, in a flat sample-157 holder. The acquisition was performed in a 0.02° interval steps, with 45 s step⁻¹. Fourier transform 158

infrared (FTIR) spectra were registered in transmission mode using a Bruker Vector 22 spectrophotometer equipped with Globar source, DTGS detector, and working with 128 scans at 4 cm^{-1} resolution in the 4000-400 cm⁻¹ range. Samples were analyzed as self-supporting pellets by dispersing the mNPs in KBr (1:20 weight ratio).

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164 **2.4 Adsorption experiments**

2.4.1 Single component PAH system: preliminary adsorption experiments were performed using 165 ANT as reference PAH substrate. Kinetic experiments were carried out contacting the adsorbent 166 (500 mg L^{-1}) with an ANT solution (50 µg L^{-1}) at pH 6 and sampling at fixed times up to a 167 maximum of 24 hours. After contact, an aliquot of 5 mL was recovered, the solid magnetically 168 separated from the solution and the supernatant analyzed. The effect of BBS amount contained in 169 MB samples on adsorption phenomenon was studied considering [ANT]=40 μ g L⁻¹, 170 $[MBpy550]=20 \text{ mg L}^{-1}$, Contact time=3 hours. The ANT concentration effects on adsorption were 171 studied at three different adsorbent dosage (10 mg L⁻¹, 20 mg L⁻¹ and 50 mg L⁻¹). A series of Pyrex-172 flasks were filled with 100 mL of ANT solutions (40-120 µg L⁻¹) and MB3py550 adsorbent were 173 added under continuous stirring. The flasks were capped and shaken in darkness during 3 hours. 174 After contact, an aliquot of 5 mL was recovered, the mNPs magnetically separated from the 175 solution and the supernatant analyzed by fluorescence spectroscopy. All adsorption experiments 176 were carried out in triplicate at $20 \pm 1^{\circ}$ C. ANT concentrations were determined by fluorescence 177 measurements obtained by means of a Hitachi F-7000 fluorescence spectrometer. The excitation 178 wavelength selected was 250 nm, slit widths of excitation and emission were 2.5 nm, and 1 cm path 179 length quartz cuvettes were used. 180

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2.4.2 Mixture of PAHs: adsorption experiments with a mixture of eight PAHs (NAP, ACL, AC,
FL, PHE, ANT, FN and PY) on MB3py550 were performed. PAHs mixture solution was prepared

from a dilution of PAHs mixture stock solution (20 mg L^{-1} of each PAH in methanol) with 184 ultrapure water and agitation during 48 h at RT in amber borosilicate glass containers. Final 185 methanol concentrations were kept under 0.1% of the total solution volume to avoid co-solvent 186 effects. The initial concentrations of each PAH ranged from 2.2 to 44 μ g L⁻¹. Concentrations of 187 each PAH after 24 h of contact with the adsorbent (50 mg L⁻¹ of MB3py550) were determined by 188 gas chromatography mass spectrometer (GC-MS) using the following procedure: 200 mL of 189 sample extracted after the magnetic separation was eluted through a Strata RP-18 cartridge; then, 190 PAHs were recovered with 6 mL of dichloromethane. The fraction was concentrated to 1 mL under 191 a nitrogen stream, transferred to a glass vial, and quantitated by GC-MS. A 30 m HP-5 MS fused 192 silica column (0.25 mm i.d., 0.25 µm film thickness) was used. Column temperature was 38 °C for 193 1 min; it increased up to 300 °C with ramping at 6 °C min⁻¹ and was held 5 min at 300 °C. Samples 194 were run in the electron impact mode at 70 eV and in the selected ion monitoring (SIM) mode with 195 a 4.04 s scan time over a 50–450 amu range resolution. Each compound was recognized by a target 196 ion and two qualifiers. 197

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2.4.3 Water-soluble fraction of light crude oil dissolved in freshwater (WSF): Adsorption 199 experiment with crude oil dissolved in water was performed in order to evaluate the adsorption 200 capacity of the MB3py550 toward a real complex matrix. Light crude oil, obtained from Rincon de 201 los Sauces, Neuquen, Argentina, stored at 4 °C, was used to prepare the WSF. A solution composed 202 of crude oil (300 μ g mL⁻¹ in methanol) and freshwater at a ratio of 1:100 (v/v) was stirred for 72 h 203 at 20 °C in the dark. WSF was separated using the bottom drain and was stored refrigerated at 4 °C 204 for 24 h. In the adsorption experiment, 10 mg of MB3py550 were added to a 200 mL WSF sample 205 and the mixture was placed on a magnetic shaker and allowed to equilibrate for 24 h at 20°C, in the 206 dark. Then, the 150 mL of supernatant was magnetically separated and spiked with recovery 207 standards (1-chloro-octadecane). Hydrocarbons concentrations in the supernatant, expressed as total 208

petroleum hydrocarbons (TPHs: C6-C35), diesel range (DRO: C10-C25) and residual range (ORO: 209 >C25-C35) was determined by gas chromatography flame ionization detector (CG-FID) following 210 the procedure described below: 150 mL of sample was eluted through Strata RP-18 cartridge using 211 6 mL of hexane and 3 mL of dichloromethane. The fraction was concentrated to 1 mL under a 212 nitrogen stream, transferred to a glass vial and quantified by GC-FID. An Agilent (Miami, USA), 213 6890 gas chromatograph equipped with a ZB-5 fused silica capillary column (30 m \times 0.25 mm i.d., 214 0.25 µm film thickness) and a flame ionization detector (FID) was used. Analyses were conducted 215 in splitless mode using nitrogen as a carrier gas at a flow rate of 2 mL min⁻¹. The column 216 temperature was programmed from 60 °C to 300 °C at a rate of 6 °C min⁻¹ and held at 300 °C for 5 217 min. Quality control and quality assurance protocols to ensure the accuracy and precision of 218 analyses included: employing blanks and reference material and duplicate samples. The average 219 recoveries based on matrix spiked samples were 70-110% and limit of detections are determined to 220 be $13 \ \mu g \ L^{-1}$ for TPHs and $17 \ \mu g \ L^{-1}$ for PAH fraction. 221

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Regeneration and reuse of BBS-derived carbon magnet-sensitive materials: Each cycle 223 2.5 of reuse consisted of two stages: sorption and regeneration. For the sorption, 50 mg of MB3py550 224 was contacted with 100 mL of 80 μ gL⁻¹ ANT solution during 3 h. After magnetic separation, an 225 aqueous aliquot was withdrawn for the determination of ANT equilibrium concentration and the 226 remained supernatant was discarded. Later, for the regeneration stage, 10 mL of methanol was 227 added to the MB3py550 and the dispersion was shaking for 10 min. The solid was magnetically 228 separated and the regenerated MB3py550 was then reused for subsequent ANT sorption 229 experiments again. 230

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232 **3. Results and discussion**

233 **3.1 Morphological and magnetic characterization**

Considering the results of the adsorption study reported below, the main part of the characterization studies described in the following concerns the sample showing the best performance in terms of pollutant removal, i.e. MB3py550 and its precursor MB3.

The morphology of the MB3 and MB3py550, are shown in **Figure 1a-d**. From the low-resolution SEM images (**Figure 1a,c**), it is clear that both samples are in the form of microagglomerates having irregular shape and exposing a complex surface. At higher magnification (**Figure 1b,d**), more evident aggregated nanoparticles with sizes in the 20-80 nm range (at the adopted resolution), are illustrated. From **Figure 1b**, it can be also noticed that salts or by-products (evidenced in the figure by the white arrow) can be present in the sample after water washing and drying, while a more homogeneous aspect is obtained for the thermally treated samples (**Figure 1d**).



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Figure 1. Low- and high-resolution SEM images of both MB3 (a,b), and MB3py550 (c,d). The arrow in section b) evidences the presence of some synthesis by-product. AFM topography of MB3py550 (e), MFM phase shift images (f,g,h) of region e) obtained at various lift heights (H = 60 nm, 80 nm and 100 nm, respectively).

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Aggregated nanoparticles, 20-50 nm in sizes, are shown in the topography image of AFM/MFM 263 (Figure 1e). Besides the AFM topography shown in Figure 1e, the phase shifting of the same 264 region obtained in a second-pass lifted acquisition was operated at different heights (H = 60 nm, 80 265 nm, 100 nm) by using the same tip probe for analyzing the magnetic interactions and minimizing 266 topographic effects. The intensity of the negative phase shifting, that is typically associated with the 267 attractive interactions between the tip and the sample (dark regions in Figure 1f-h), was found to 268 depend on the adopted heights: the higher resolution was obtained at H = 60 nm and the magnetic 269 characteristics of the more prominent nanoparticles were found at H = 80 nm, while at higher 270 heights the signal-to-noise ratio is minimized. 271

To better investigate the magnetic properties, an extensive characterization was performed. The 272 magnetic properties of all samples are reported in **Table S2**, whereas magnetization curve profiles 273 274 of both reference magnetite/maghemite and the MB3-series of samples are reported in Figure S2. In general, all bare samples (not thermally treated) reveal superparamagnetic behaviors, with 275 276 negligible remanence (M_r) and very low coercitivity (H_c below 10 Oe). As already reported in our previous work (Franzoso et al., 2017), the values of saturation magnetization (M_s) of bare BBS-277 stabilized NPs were respectively 56 (MB1), 36 (MB2), and 30 (MB3) emu g⁻¹, thus suggesting a 278 different amount of magnetite/maghemite per gram of sample. Moreover, the decrease in terms of 279 M_s for all the BBS-stabilized samples compared to the neat magnetite M0 (64 emu g⁻¹) is mainly 280 due to the presence of the BBS shell, which induces the reduction of the surface magnetic moments 281 (Kim et al., 2003; Sun et al., 2009; Cesano et al., 2015; Bianco Prevot et al., 2017; Franzoso et al., 282 2017; Nisticò, 2017b). 283

Conversely, the pyrolysis treatment performed at 550 °C mainly affects the remanence (which moves from almost zero up to 3.4-5.2 emu g^{-1}) and the coercitivity (which increases from less than

10 up to 53-82 Oe). However, saturation magnetization value still remains below the neat
magnetite/maghemite M0 value.

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289 **3.2** Pyrolysis-induced effects on BBS-derived magnet-sensitive NPs

In order to unveil the chemistry behind the pyrolysis treatments, both XRD and FTIR analyses were performed at different experimental conditions. In particular, the iron oxide phases forming the magnet-responsive inorganic core of the NPs were identified through XRD. For the sake of brevity only the MB3-series of samples are displayed in **Figure 2.a**.



Figure 2.a. XRD patterns of the neat magnetite M0 (A, solid black curve), MB3 (B, solid red curve), and MB3py550 (C, solid blue curve). Main reflections due to magnetite phase are highlighted. Non-labeled peaks refer to by-products signals. **b.** FTIR spectra in the 4000-500 cm⁻¹ range relative to neat magnetite M0 (A, dotted black curve), MB3py550 (B, solid red curve) MB3 (C, solid black curve) and BBS-GC (D, dotted red curve).

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By considering the bare MB3 sample (**Figure 2.a**, diffractogram B), all the crystalline planes highlighted in the figure at $2\theta = 30.1^{\circ}$ (220), 35.4° (311), 43.0° (400), 53.9° (422) 57.2° (511), and 62.6° (440) are consistent with both magnetite (card number 00-019-0629 from ICCD Database) and maghemite (card number 00-039-1346 from ICCD Database) (Cesano et al., 2015). No relevant

reflections are expected from BBS since its XRD pattern presents only one broad amorphous 315 contribution centered at ca. $2\theta = 25^{\circ}$ and few negligible signals due to its ash content (Figure S3). 316 All the extra signals (not labeled) are consistent with the presence of by-products of the co-317 precipitation reaction carried out in ammonia aqueous solution, i.e., ammonium-containing salts as 318 ammonium chloride (card number 01-073-0363 from ICCD Database) in analogy to our previous 319 work (Nisticò et al., 2017). Pyrolysis treatments carried out at 550 °C favored the 320 abatement/disappearance of these ammonia-derived salts, maintaining the magnetite/maghemite 321 crystalline phases (Figure 2.a, diffractogram C). 322

Pyrolysis-induced changes affecting the organic shell and its interaction with the magnetic core of the NPs were followed by means of FTIR analysis in the 4000-500 cm⁻¹ range. Infrared spectra of both reference magnetite (M0) and BBS, together with those of the MB3-series are collected in **Figure 2.b**.

The signals at 575 and 620 cm⁻¹ evidenced in M0, MB3 and MB3py500 are due to Fe-O stretching 327 vibrations and consistent with the presence of iron oxides (Figure 2.b, blue box) (Cesano et al., 328 2015; Franzoso et al., 2017; Nisticò et al., 2017), whereas the presence of BBS in MB3 is 329 confirmed by the presence of the signal at ca. 1600 cm⁻¹ due to BBS-carboxylate stretching mode 330 (Bianco Prevot et al., 2017) (Figure 2.b, green box) and 1110 cm⁻¹ which can be assigned to C-O 331 stretching vibration. The interaction between BBS and the iron oxide core of the NPs is confirmed 332 by the presence of the very sharp band at 1400 cm⁻¹ which is an evidence of the direct interaction 333 between BBS-carboxylate functionalities and the magnetite surface (i.e., carboxylate-iron bond 334 stretching mode) (Ou et al., 2009), and by the shift of the absorption band assigned to C-O 335 stretching mode attributable to the organic matter (mainly polysaccharides and other BBS-derived 336 substances) adsorbed onto the iron oxide surface (Figure 2.b, red box). The thermal treatment 337 carried out at 550 °C (Figure 2.b, curve B) induced the significant reduction of all the infrared 338 signals representative of the BBS-GC coating. On the other hand, the presence of the absorption 339

band at 1620 cm⁻¹ typical of the C=C stretching mode, suggesting the incipient conversion of the BBS into graphite-like materials. The iron oxide phase signals at 575 cm⁻¹ and 620 cm⁻¹ remained unaltered, confirming what already evidenced by XRD results.

These characterization results of MB3py550 and its precursor MB3 were similar to those previously obtained in another synthesis batch (Nisticò et al., 2018), which confirms the reproducibility of the synthesis method.

346

347 **3.3 Adsorption studies**

Figure 3 shows the kinetic of adsorption of ANT toward both MB3 and MB3py550. The 348 equilibrium was reached after approximately 3 h for MB3py550. Interestingly, MB3 showed some 349 retention capacity toward ANT (70% of removal after 22 h of contact). This could be explained 350 considering that ANT is capable of interacting with the π electrons associated with the aromatic 351 moieties in the BBS matrix. However, a higher ANT % removal was obtained for MB3py550 (ca. 352 99%), which is expected due to the increased hydrophobicity of the material after pyrolysis and 353 consequent formation of graphite-like structures, thus confirming the results obtained from FTIR 354 spectra. In all cases, pyrolized materials showed higher ANT uptake than non-pyrolyzed materials 355 (data not shown). It is well documented in the literature that π -electron systems of the graphite-like 356 materials can be engaged in $\pi - \pi$ electron-donor-acceptor (EDA) interactions with a series of π -357 donors (e.g., benzene, toluene, PAHs) which can enhance their removal capacity toward aromatic 358 359 compounds (Zhu and Pignatello, 2005; Keiluweit and Kleber, 2009).



361 362

Figure 3. Kinetics of ANT adsorption on MB3 (black squares, black solid line) and MB3py550 (red circles, red solid line) nanoparticles. [ANT]=50 µg L⁻¹, [MB3py550]=500 mg L⁻¹.

The ANT removal capacity of the three pyrolyzed nanomaterials with different amounts of BBS-GC (i.e. MB1py550, MB2 py550 and MB3 py550) was studied. The best uptake of ANT was observed for MB3py550 (**Figure S4**), for this reason this adsorbent was selected for the investigation of the potential application of these type of materials in different experimental conditions.

In order to optimize the experimental conditions, and in particular to find out the minimum amount of adsorbent needed for an effective ANT removal, sorption experiments were performed by varying the amount of MB3py550 (10, 20 and 50 mg L⁻¹) for different ANT concentration (40, 80 and 120 μ g L⁻¹). The ANT removal capacity increased along with the adsorbent amount increase achiving more than 95% removal for all ANT concentration tested with 50 mg L⁻¹ of MB3py550 and for 40 μ g L⁻¹ ANT with 20 mg L⁻¹ of adsorbent (**Figure 4**).



378

Figure 4. Effect of initial concentration of ANT and [MB3py550] dosage on ANT removal.
Contact time = 3 h. All experiments are performed at both 20 ± 1° C and pH 6.

381

In order to compare our results with those reported in literature for different nanomaterials, the 382 sorption capacity of ANT, determined as the difference between initial concentration of ANT and 383 the equilibrium concentration of ANT per unit mass of adsorbent, was calculated for each 384 MBSS3py550 dosage and each initial ANT concentration tested in Figure 4. The highest sorption 385 capacity of ANT on MB3py550 was 4.5 mg g⁻¹ corresponding to 20 mg L⁻¹ of adsorbent dosage and 386 ANT concentration of 40 μ g L⁻¹. In terms of sorption capacity of ANT, MB3py550 is superior to 387 mesostructured silica nanoparticles (1.6 mg g^{-1}) (Topuz and Uyar, 2017), similar to electrospun 388 nanofibrous membranes (4.1 mg g⁻¹) (Dai et al., 2011) and lower than single-walled carbon 389 nanotubes multiwalled carbon nanotubes (Yang et al., 2006). However, being superparamagnetic, 390 the MB3py550 have the unique capability of being magnetically treatable. Moreover, a comparison 391 with previous results obtained with chitosan as a source of carbon for preparing carbon-coated 392 magnetic nanoparticles, shows that MB3py550 have a sorption capacity of ANT almost one order 393

394 of magnitude higher (Nisticò et al., 2017), which valorizes the use of BBS-GC in this type of 395 sorbents.

To evaluate the removal capacity of MB3py550 towards different PAHs, a mixture of NAP, ACL,

AC, FL, PHE, ANT, FN and PY was contacted with 50 mg L^{-1} of adsorbent (**Figure 5.a**).

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Figure 5.a. Adsorption experiments of PAHs on MB3py550. [PAHs] = 2.2 to 44 μ g L⁻¹, [MB3py550] = 50 mg L⁻¹, Contact time = 24 h. **b.** Uptake of the water-soluble fraction of crude oil by MB3py550 nanoparticles. Conditions: [TPH] = 0.94 mg L⁻¹, [MB3py550] = 50 mg L⁻¹, Contact time = 24 h. All experiments are performed at 20 ± 1 °C and pH 6. Error bars represent standard deviation of three independent experiments.

412

The results show different removal capacities depending on the structure of PAHs; in particular, 413 high % removal (> 80%) was obtained for FL, ANT, PHE, PY and FN, whereas lower retention was 414 observed for NAP, AC and ACL. The removal capacity increased along with the number of 415 aromatic rings in the polycyclic structure and the PAHs hydrophobicity, suggesting that the 416 pyrolyzed nanomaterials are substantially good in the removal of PAHs from aqueous solutions. 417 Moreover, a clear trend of the removal capacity with the PAHs hydrophobicity (K_{ow}) was also 418 observed (see K_{ow} values in the experimental section). These results can be explained considering 419 that the PAHs sorption occurs following two mechanisms: in the first one the π - π interactions allow 420

the adsorption of the molecules with the more extended aromaticity, whereas in the second one thehydrophobicity of the material attracts PAHs to the adsorbing surface.

Finally, the removal capacity of BBS-derived carbon magnet-sensitive adsorbent toward a sample 423 water spiked with crude oil was studied. The experiment was performed by putting in contact the 424 water-soluble fraction (WSF) with MB3py500 (50 mg L^{-1}) at pH 6 and 20 ±1 °C. The composition 425 of the WSF used in the experiments was: C12–C25 alkanes (0.78 mg L^{-1}) and C26–C36 alkanes 426 (0.16 mg L⁻¹). It is worth mentioning that the PAHs present in the WSF sample belong to the C26-427 C36 fraction. Figure 5.b shows a higher uptake of heavier fraction hydrocarbons carried out by 428 MB3py550 (ca. 70% removal), thus meaning that the adsorption capacity increased along with the 429 hydrocarbon hydrophobicity as was also observed for the PAHs adsorption test. These promising 430 results encourage the use of urban wastes as starting materials for the preparation of sustainable 431 substrates to be successfully applied in the removal of hydrocarbons from contaminated waters. 432

The ANT removal during six continuous cycles of regeneration and reuse are shown in Figure 6. It was found that no loss of ANT adsorption capacity was observed on the regenerated MB3py550 after six cycles. This reusability of the adsorbent is one of the most important features for their promising applications in environmental detoxification, specifically continuous flow-through systems could be designed for contaminated water treatments with high ANT removal efficiency.

438



440

Figure 6. Cycles of reuse (adsorption shown) of MB3py550 in the removal of ANT. $[ANT]_0 = 80$ 442 $\mu g L^{-1}$, $[MB3py550] = 500 m g L^{-1}$, contact time = 3h.

443

444 **4.** Conclusions

Urban composted biowaste-derived substances with a lignin-like structure have proven to be an effective carbon-source for the production of easily-recoverable, low-cost carbon-coated magnetsensitive NPs, produced through pyrolysis treatments performed in nitrogen at 550 °C. Adsorption experiments testing the removal of PAHs from contaminated water evidenced that the samples thermally treated are effective due to the π - π conjugation between the PAHs and the π -electron system of the pyrolyzed material. The successful adsorption results suggest their possible application in wastewater remediation processes from PAHs-induced pollution.

452

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- 465

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| | ACCEPTED MANUSCRIPT | | | | | | |
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| 596 | | | | | | | |
| 597 | Supplementary Material | | | | | | |
| 598 | | | | | | | |
| 599 | From biowaste to magnet-responsive materials for water remediation from polycyclic | | | | | | |
| 600 | aromatic hydrocarbons | | | | | | |
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| | | | | | | | |

615

616 **Table S1**. BBS-GC chemical composition and functional groups.

| Metal analysis | | | | | | |
|------------------------------|-----------------|-----------------|-----------------|---------------|--------------------------|-----------------------------|
| Si (%) | Fe (%) | Al (%) | Mg (%) | Ca (%) | K (%) | Na (%) |
| 12.14±0.07 | 1.03 ± 0.02 | 0.59 ± 0.01 | 1.67 ± 0.25 | 4.86±0.61 | 1.18 ± 0.07 | 0.06 ± 0.01 |
| Microelements | | | | | | |
| Cu (ppm) | Ni (ppm) | Zn (ppm) | Cr (ppm) | Pb (ppm) | Cd (ppm) | Hg (ppm) |
| 73±1 | 100±3 | 157±13 | 49±1 | 43±2 | < 0.02 | < 0.02 |
| General characteristics Char | | | | Characteristi | cs in aqueous solu | ution 3g/L ^{a)} |
| Moisture (%) | Ashes (%) | C (%) | N (%) | рН | Conducibility (µS/cm) | Surface tension (N/m) |
| 3.69 | 31.2 | 39.94±0.35 | 4.82±0.14 | 9.02 | 429 | 56.8 |

Concentration values as mole fraction of total C for functional groups and C types in BBS-GC by

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| | | | NMR analysi | s ^{b)} | | |
|-------|------|------|-------------|-----------------|------|------|
| Aliph | NR | COOH | OR | Ph | PhOH | Kt |
| 0.31 | 0.07 | 0.12 | 0.20 | 0.16 | 0.06 | 0.02 |

^{a)} maximum solubility is 100 g/L in water at ca. 60 °C. solutions are stable in one week at different temperatures (tests conducted at 4 °C, 25 °C and 40 °C) with water hardness \leq 40 °F. ^{b)} legends: aliph = aliphatic groups, NR = amino groups, COOH = carboxylic acids, OR = alkoxy groups, Ph = aromatic phenyl groups, PhOH = phenoxy groups, Kt = ketones.

621

622 **BBS-GC thermal characterization**

623 Thermo-gravimetric analysis (TGA) was carried out by means of a TA Q600 (TA Instruments).

TGA tests were performed in order to monitor the BBS matrix degradation. The thermal analysis

625 was performed with a heating ramp of 10 °C/min from RT to 800 °C, either in nitrogen or in air

atmosphere. Two replicas were registered for each treatment condition.



627

Figure S1. TG curves of BBS-GC reference material heated either in nitrogen (solid black line) or
in air (solid red line) atmosphere. The black dotted line refers to the temperature investigated for the
pyrolysis treatments (namely, 550 °C).

631

The TGA analysis of BBS carried out in nitrogen (inert) atmosphere presents two main weight losses: the first one due to the water evaporation (centered at ca. 100 °C), whereas the second one is mainly due to the degradation phenomena involving the BBS organic structure (in the 250-600 °C range). A carbonaceous residue (containing some ashes), which represented ca. the 50 wt.% of the starting sample, was collected at 800 °C. Vice versa, the TGA analysis of BBS performed under air (oxidant) atmosphere evidences the complete oxidation of the BBS organic shell, leaving an

638 inorganic residue, corresponding to the ash content in the sample, not far from the value of 31 wt.%

639 reported in **Table S1**.

Table S2. Magnetic properties of samples registered at RT.

| Samples | Saturation M_s (emu/g) | Remanence M_r (emu/g) | Coercivity H _c (Oe) |
|----------|--------------------------|-------------------------|--------------------------------|
| M0 | 64 | 1.0 | 10 |
| MB1 | 53 | 0.1 | 2 |
| MB2 | 36 | 0.6 | 8 |
| MB3 | 30 | 0.2 | 4 |
| MB1py550 | 45 | 5.2 | 82 |
| MB2py550 | 44 | 4.7 | 82 |
| MB3py550 | 51 | 3.4 | 53 |



Figure S2. Magnetization curves relative to neat magnetite and BBS-stabilized (either as prepared
and pyrolyzed) samples. Legend: M0 (black triangles, dotted black curve), MB3 (red circles, solid
red curve), and MB3py550 (blue circles, solid blue curve).



Figure S4. Effect of BBS-stabilized magnetite NPs used as adsorbent on ANT removal. [ANT] = $40 \ \mu g \ L^{-1}$, [MBpy550] = $20 \ mg \ L^{-1}$, Contact time = 3 h. Error bars represent standard deviation of three independent experiments.

HIGHLIGHTS

- Biowaste derived substances were used for production of magnet sensitive
 adsorbent.
- Carbon-coated magnetite NPs were produced through pyrolysis treatments.
- Biowaste-derived substances coated iron oxides are successfully used for PAHs removal.
- Thermally treated NPs show high adsorption capacity for PAHs.
- The reusability of NPs after six cycles is found to be excellent.
- Results encouraged the application of these NPs in wastewater remediation.