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**Biochars intended for water filtration: A comparative study with activated carbons of their physicochemical properties and removal efficiency towards neutral and anionic organic pollutants**

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(Article begins on next page)

1 **Biochars intended for water filtration: a comparative study with activated carbons of their**  
2 **physicochemical properties and removal efficiency towards neutral and anionic organic**  
3 **pollutants**

4  
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24 **Abstract**

25 Seven biochars (BCs) obtained from pyrolysis or gasification of different vegetal feedstocks were  
26 thoroughly characterized in comparison with three commercial activated carbons (ACs) routinely  
27 used in drinking water treatment plants. BCs and ACs characterization included the determinations  
28 of ash, iodine and methylene blue adsorption indexes, and the release of metals and polycyclic  
29 aromatic hydrocarbons, which were performed according to international standards applied for  
30 adsorption media to be used in drinking waters. Total specific surface area, micropore and mesopore  
31 specific surface area, pH of the point of zero charge, and the release of polychlorinated biphenyls  
32 were also determined in all chars. Principal component analysis and cluster analysis were performed  
33 in order to summarize the complex set of information deriving from the aforementioned  
34 characterizations, highlighting the BC most similar (BC6 from high temperature gasification of  
35 woody biomass) and most different (BC7 from low-temperature pyrolysis of corn cob) from ACs.  
36 These BCs were studied for their adsorption in ultrapure water towards diiodoacetic acid (an emergent  
37 disinfection by-product), benzene, and 1,2-dichlorobenzene, in comparison with ACs, and results  
38 obtained were fitted by linearized Freundlich equation. Overall, BC6 showed higher sorption  
39 performances compared to BC7, even though both BCs were less performing sorbents than ACs.  
40 However, the sorption properties of BCs were maintained also in real water samples collected from  
41 drinking water treatment plants.

42

43 **Keywords:** pyrolysis, gasification, filtration media, PAH and PCB release, metal release, drinking  
44 water plant

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## 50 **1 Introduction**

51 Pollution by anthropic activities still affects the quality of water resources. According to the European  
52 Water Framework Directive (2000/60/EU), waters must achieve good ecological and chemical status,  
53 to protect human health, water supply, natural ecosystems and biodiversity (European Commission,  
54 2010).

55 The accomplishment of the environmental objectives set by the European policies can be achieved  
56 through proper integrated water management policies and technological approaches, which must be  
57 sustainable from both environmental and economic points of view. Within this framework, water  
58 treatments play significant and crucial roles in the supply of safe waters intended for human  
59 consumption.

60 Adsorption is an effective and economically feasible approach routinely integrated in drinking water  
61 production (Ali and Gupta, 2006) for the removal of micropollutants occurring *per se* in raw waters  
62 and/or as a result of disinfection processes, thereby eliminating the toxicity induced by these  
63 molecules in treated water (Han and Zhang, 2018; Han et al., 2021). Within this context, adsorption  
64 is generally based on the use of activated carbon (Jiang et al., 2017; Perrich, 2018; Jiang et al., 2020),  
65 even though innovative sorbents, such as mesoporous silica (Kyzas and Matis, 2015; Rivoira et al.,  
66 2016) or other waste-derived ceramic materials (Jana et al., 2016; Bruzzoniti et al., 2018) have been  
67 proposed as alternative adsorption media. Low-cost materials like biochar (BC) have recently  
68 received attention for their physicochemical characteristics including the porous structure, which is  
69 similar to that of activated carbons. Biochar is the solid by-product of the thermal conversion of a  
70 wide range of feedstocks, such as agricultural wastes (Ali and Gupta, 2006; Colantoni et al., 2016),  
71 wood residues (Wang et al., 2013), manure (Cao and Harris, 2010), and sludge (Méndez et al., 2017).  
72 Due to its properties, biochar has found application as animal feed additives (McHenry, 2010) and  
73 soil amenders (Singh et al., 2010), as well as for the adsorption of micropollutants from aqueous  
74 matrices (Palansooriya et al., 2020).

75 Internationally recognized standards detailing the physicochemical characteristics of biochars to be  
76 used for agricultural applications have been recently released (European Biochar Foundation (EBC)).  
77 Similar standards have not yet been established on the characteristics required for biochars to be used  
78 in water purification. However, international standards are available that require compliance with  
79 specific limits for certain physical and chemical parameters in adsorbent materials (Comite Europeen  
80 de Normalisation (CEN), 2004), and particularly in activated carbons (Comite Europeen de  
81 Normalisation (CEN), 2009), used for the treatment of drinking water.

82 In the last years, biochars obtained from a very wide range of experimental conditions (e.g. feedstock,  
83 thermo-chemical process and pretreatment of biomass and/or post-treatment of biochar) were  
84 investigated as sorbent media for water purification issues, highlighting their promising adsorption  
85 properties towards a large variety of organic and inorganic contaminants (Gwenzi et al., 2017; Wang  
86 et al., 2020). However, it should be emphasized that, with few exceptions (Del Bubba et al., 2020) it  
87 is not verified whether the biochars prepared in the various experimental conditions comply with the  
88 requirements set out in the aforementioned standards, relating to the adsorbent materials intended to  
89 be used for the filtration of drinking water. Moreover, in most cases, ultrapure water is used to  
90 investigate sorption capabilities of biochars, whilst it would be advisable to perform these studies  
91 using real aqueous matrixes. Last but not least, except in rare cases (Del Bubba et al., 2020), no  
92 comparison has been made with the adsorption capacity of standard activated carbons (ACs). All  
93 these aspects represent obvious limitations in the reliable evaluation of the applicability of biochars  
94 for water treatment (Castiglioni et al., 2021).

95 Based on the considerations mentioned above, the aim of this research was to investigate the  
96 physicochemical properties, the regulated leachable substances, and the removal performances of  
97 seven biochars (commercially available or synthesized for the purpose), obtained from pyrolysis or  
98 gasification of vegetal biomass, in comparison with three commercially available vegetal ACs used  
99 in an Italian drinking water facility, at different age of operation. Data obtained were chemometrically  
100 treated through principal component analysis, allowing for selecting the most promising biochars to

101 be further investigated by adsorption tests. In a first phase of this study, adsorption capabilities were  
102 tested in ultrapure water, whereas afterwards the sorption capacity was evaluated on a restricted group  
103 of biochars in water samples collected at intermediate treatment stages of a potabilization plant. In  
104 all cases ACs were also tested as reference comparative materials.

105 Diiodoacetic acid (DIAA), benzene, and 1,2 dichlorobenzene, were selected as model pollutants  
106 commonly monitored in drinking water facilities. Specifically, DIAA is a model emerging  
107 disinfection by-product (Bruzzoniti et al., 2019b) never investigated before for its sorption by  
108 biochars. Moreover, 1,2 dichlorobenzene can also originate from disinfection treatments during the  
109 potabilization process (Lahaniatis et al., 1994; Hou et al., 2012) and its monitoring in tap water is  
110 recommended by the World Health Organization guidelines (taste threshold value  $1 \mu\text{g L}^{-1}$ ) (World  
111 Health Organization, 2017), while benzene is regulated by the Directive 2020/2184 regarding the  
112 quality of water intended for human consumption ( $1 \mu\text{g L}^{-1}$ ). It should also be noted that benzene and  
113 1,2 dichlorobenzene are volatile organic carbons (VOCs) still detected in some industrial districts  
114 (Martínez et al., 2002) and are therefore also important from the wastewater treatment viewpoint.

## 115 **2 Materials and methods**

### 116 **2.1 Reagents**

117 For the determination of adsorption indexes, the following reagents, supplied by Merck (Kenilworth,  
118 NJ, USA), were used: iodine solution (0.1 N), sodium thiosulfate solution (0.1 N), zinc iodide starch  
119 solution, hydrochloric acid (37%), potassium hexacyanoferrate (>99%), methylene blue, anhydrous  
120 acetic acid (>99.8%). Ammonia solution (28%), dichloromethane and 2-propanol were from VWR  
121 International (Radnor, PA, USA). For the evaluation of extractable metals, an ICP multi-element  
122 standard solution IX ( $100 \text{ mg L}^{-1}$  of As, Be, Cd, Cr (VI), Ni, Pb, Se, Tl) from Merck was used.

123 For the determination of extractable polycyclic aromatic hydrocarbons (PAHs), the 16 compounds  
124 listed by EPA were purchased from Sigma Aldrich (Darmstadt, Germany). For the analysis of  
125 extractable polychlorinated biphenyls (PCBs), the compounds were purchased from LGC Standards  
126 (Milan, Italy). They were non dioxin-like PCBs: 3,3'-dichlorobiphenyl (PCB 11), 4,4'-

127 dichlorobiphenyl (PCB 15), 2,4,4'-trichlorobiphenyl (PCB 28), 2,2',5,5'-tetrachlorobiphenyl (PCB  
128 52), 2,2',4,5,5'-pentachlorobiphenyl (PCB 101), 2,2',3,4,4',5-hexachlorobiphenyl (PCB 138),  
129 2,2',4,4',5,5'-hexachlorobiphenyl (PCB 153), 3,3',4,4',5,5'-hexachlorobiphenyl (PCB 169),  
130 2,2',3,4,4',5,5'-heptachlorobiphenyl (PCB 180), 2,3,3',4,4',5,5'-heptachlorobiphenyl (PCB 189);  
131 and dioxin-like PCBs: 3,4,4',5-tetrachlorobiphenyl (PCB 81), 2,3',4,4',5-pentachlorobiphenyl (PCB  
132 118), 2',3,4,4',5-pentachlorobiphenyl (PCB 123), 2,3',4,4',5,5'-hexachlorobiphenyl (PCB 167).

133 Labelled isotope compounds for PCBs ( $2 \text{ mg L}^{-1}$ ) and for PAHs ( $5 \text{ mg L}^{-1}$ ), Wellington Laboratories  
134 (Ontario, Canada), were used as internal and surrogate standards in order to obtain calibration curves  
135 and extraction recoveries, respectively. The  $^{13}\text{C}$  surrogate solutions of PAHs contained:  
136 [ $^{13}\text{C}_6$ ]benzo(a)anthracene [ $^{13}\text{C}_6$ -BaA], [ $^{13}\text{C}_6$ ]chrysene [ $^{13}\text{C}_6$ -Chr], [ $^{13}\text{C}_6$ ]benzo(b)fluoranthene [ $^{13}\text{C}_6$ -  
137 BbFl], [ $^{13}\text{C}_6$ ]benzo(k)fluoranthene [ $^{13}\text{C}_6$ -BkFl], [ $^{13}\text{C}_4$ ]benzo(a)pyrene [ $^{13}\text{C}_4$ -BaP],  
138 [ $^{13}\text{C}_6$ ]indeno(1,2,3-cd)pyrene [ $^{13}\text{C}_4$ -Ind], [ $^{13}\text{C}_6$ ]dibenzo(a,h)anthracene [ $^{13}\text{C}_6$ -DBA], and  
139 [ $^{13}\text{C}_{12}$ ]benzo(g,h,i)perylene [ $^{13}\text{C}_{12}$ -BP]. The  $^{13}\text{C}$  surrogate solution of PCBs contained:  $^{13}\text{C}_{12}$ -PCB28,  
140  $^{13}\text{C}_{12}$ -PCB52,  $^{13}\text{C}_{12}$ -PCB118,  $^{13}\text{C}_{12}$ -PCB153, and  $^{13}\text{C}_{12}$ -PCB180.

141 Volatile organic compounds, namely benzene ( $100 \text{ } \mu\text{g L}^{-1}$  in methanol), and 1,2-dichlorobenzene  
142 ( $100 \text{ } \mu\text{g L}^{-1}$  in methanol), were purchased from Ultra Scientific Italia (Bologna, Italy). DIAA was  
143 supplied by Chemical Research (Rome, Italy). Ultrapure water was obtained by an EMD Millipore  
144 Milli-Q Direct Water Purification System (Millipore, Bedford, MA, USA).

## 145 2.2 Biochar and activated carbon samples

146 The seven BCs considered in this study were donated for the purpose by different companies, which  
147 produce the materials for commercial scopes or for their internal use through well-established  
148 procedures. The three ACs were supplied by a local potabilization plant at different age of operation:  
149 AC1: new activated carbon; AC2: regenerated activated carbon; AC3: regenerated activated carbon  
150 in use at the plant. The status of operation, and the characteristics of the feedstock and the thermal  
151 process used to produce the ten chars are summarized in **Table 1**.

152 Before being characterized and used in isotherm studies, all the char samples were repeatedly washed  
153 with ultrapure water according to the ASTM D-5919-96 method.

### 154 2.3 Biochar and activated carbon characterization

155 The chars investigated in this study were physicochemically characterized through the determination  
156 of ash content, pH of the point of zero-charge ( $\text{pH}_{\text{pzc}}$ ), physisorption analysis, iodine and methylene  
157 blue adsorption indexes ( $\text{I}_2\text{In}$  and  $\text{MBIn}$ ), as well as for water-extractable substances of environmental  
158 concern. The procedures adopted for the aforementioned determinations are briefly described below,  
159 whilst full details are provided in the *Supplementary Material* section. For the determination of the  
160 parameters described below, standard methods were used when available.

#### 161 2.3.1 Ash content

162 The ash content was determined according to the ASTM International D 2866-11 (American Standard  
163 Test Method (ASTM), 2018), which refers to the analysis of ACs.

#### 164 2.3.2 Water-extractable substances

165 Metals (namely Sb, As, Cd, Cr, Pb, Hg, Ni and Se), PAHs and PCBs were extracted according to the  
166 EN 12902 standard (Comite Europeen de Normalisation (CEN), 2004). After extraction, metals were  
167 determined by an Elan 6100 ICP-MS (Perkin Elmer, Waltham, Massachusetts, USA), whereas PAHs  
168 and PCBs were preconcentrated by solid-phase extraction (SPE) and analysed by GC-MS, as  
169 elsewhere described (Bruzzoniti et al., 2019a; Rivoira et al., 2019). Quality controls were performed  
170 to verify the recovery efficiency of PAHs and PCBs during the preconcentration step, using labelled  
171 standards, as specified in the *Supplementary material* (see **Tables S1** for BCs and **S2** for ACs).

#### 172 2.3.3 pH of the point of zero charge

173 The pH of the point of zero-charge ( $\text{pH}_{\text{pzc}}$ ) was determined using the pH drift method, widely adopted  
174 for the evaluation of the surface charge of biochars and ACs (Del Bubba et al., 2020).

#### 175 2.3.4 Adsorption indexes

176 The determination of  $\text{I}_2\text{In}$  and  $\text{MBIn}$  was performed according to the definitions indicated by CEFIC  
177 for ACs (Conseil Européen des Fédérations de l'Industrie Chimique (CEFIC), 1986).



178 2.3.5 *Physisorption analysis*

179 Physisorption analysis of biochars and ACs was performed via nitrogen adsorption and desorption  
180 experiments using a Porosity Analyser Thermo Fisher Scientific (Milan, Italy) model  
181 SORPTOMATIC 1990 according to the American Society for Testing and Materials specifications  
182 (American Society for Testing and Materials, 2012, 2017). In further detail, the specific surface area  
183 (SSA) and micropore surface area (MiSSA) were determined respectively by the Brunauer–Emmett–  
184 Teller (BET) method and by t-plot method, whereas mesopore surface area (MeSSA) was measured  
185 by the Barrett-Joyner-Halenda (BJH) method applied to desorption data.

186 2.3.6 *Quality control of biochar and activated carbon characterization*

187 Quality control of char characterization measurements was carried out by comparing the results here  
188 obtained for AC1 and AC2 with those reported in the technical specifications of the two commercial  
189 materials.

190 2.4 Adsorption studies on DIAA and VOCs

191 Adsorption tests were performed on DIAA and VOCs in ultrapure water (pH = 6.5±0.1) using the  
192 micro-isotherm technique for adsorbates at ppb concentrations, as established by ASTM D5919-96  
193 standard (American Standard Test Method (ASTM), 1996). Aliquots of 40 mL and 100 mL,  
194 containing fixed amounts of DIAA and VOCs, respectively (DIAA: 5 µg L<sup>-1</sup> for BCs and 20 µg L<sup>-1</sup>  
195 for ACs; VOCs: 5 µg L<sup>-1</sup> or 20 µg L<sup>-1</sup> for BCs and 20 µg L<sup>-1</sup> for ACs) were put in contact with  
196 different amounts of BCs/ACs varying approximately between 0.02 and 0.5 g. The mixture was  
197 stirred in an orbital shaker for 24 hours. The solution was then filtered through a mixed cellulose ester  
198 membrane (0.45 µm). Control experiments using the same aforementioned concentrations of target  
199 analytes were also conducted without the addition of the adsorbent materials, in order to estimate  
200 their removal due to mechanisms other than sorption (e.g. volatilization and degradation).

201 Experimental data were fitted by the Freundlich isotherm model (Foo and Hameed, 2010):

202 
$$\log \frac{X}{M} = \log K_F + \frac{1}{n} \log C_e$$

203 Where  $X/M$  is the ratio of the amount of analyte adsorbed per mass unit of sorbent ( $\text{mg g}^{-1}$ ),  $K_F$  is the  
204 constant of the Freundlich isotherm equation ( $\text{mg}^{1-1/n} \text{L}^{1/n} \text{g}^{-1}$ ) related to adsorption capacity,  $C_e$  is  
205 the equilibrium concentration ( $\text{mg L}^{-1}$ ), and  $1/n$  is the exponent of non-linearity.

## 206 2.5 Analytical determination of DIAA and VOCs

207 Residual DIAA concentrations were determined by ion chromatography coupled with triple-stage  
208 quadrupole mass spectrometry as elsewhere described (Bruzzoniti et al., 2019b). Residual VOC  
209 concentrations were determined by GC-MS after SPE. Details of both procedures are given in the  
210 *Supplementary Material Section*.

## 211 2.6 Water sample collection and characterization

212 Water samples were withdrawn from two potabilization plants located in the Piedmont region (North  
213 Italy) which treat the same raw water. One sample (labelled as DSB) was taken at the outlet of the  
214 dynamic separation basins for the removal of slurry from clarified waters (in which coagulant,  
215 hypochlorite and chlorine dioxide solutions are dosed), before entering the activated carbon beds. The  
216 other sample (labelled as CB) was taken at the outlet of a clarification basin (in which coagulant only  
217 is added), before entering the activated carbon beds. The water samples were characterized for pH  
218 and total organic carbon (TOC). TOC was determined using a Shimadzu TOC-V-CSH analyser, by  
219 the differential method, i.e. analysing both total carbon (TC) and total inorganic carbon (TIC) through  
220 separate measurements and calculating TOC by subtracting TIC from TC.

## 221 2.7 Data analysis

222 Least squares regressions and related analyses of variance (ANOVA) were performed with Excel  
223 2016 (Microsoft, Redmond, WA, USA). Principal component analysis (PCA) and cluster analysis  
224 (CA) were carried out using the Minitab statistical software package, version 17.1.0 (Minitab Inc.,  
225 State College, PA, USA). All data plots were performed using Excel 2016.

# 226 3 Results and discussion

## 227 3.1 Characterization of chars

228 *3.1.1 Ash content*

229 In biochars, ash percentages were found in the quite wide range of 6-49% (**Table 2**), with the lowest  
230 value achieved for BC7, which derived from corn cob under pyrolysis treatment at 450°C (**Table 1**).

231 Ash content in materials intended for water filtration is regulated by EN 12915-1 standard, which sets  
232 a limit of 15%, since a high ash content in filtering media is expected to reduce adsorption activity  
233 (Inyang and Dickenson, 2015). Hence, as regards this parameter, only BC2, BC3, and as previously  
234 mentioned BC7, are allowed to be used as sorbent materials in potabilization facilities.

235 The data obtained here can be interpreted based on the characteristics of biomass and thermal  
236 conversion processes through which the chars were obtained. According to literature, ash  
237 concentration of chars is mainly influenced by the type of feedstock, being woody biomass the one  
238 providing a lower ash content, compared to other feedstocks, such as non-woody vegetal biomass and  
239 animal waste (Tomczyk et al., 2020). However, the type of thermal conversion process (i.e. pyrolysis  
240 or gasification) and the temperature and contact time conditions adopted in the process may also play  
241 a role in determining the ash concentration, which obviously depends on the amount of char obtained.

242 In this regard, it should be remarked that biochar yield is a function of the type of thermal conversion  
243 process (i.e. pyrolysis or gasification) and the temperature and contact time conditions adopted in the  
244 process, being the highest yields obtained with pyrolysis conducted at low temperature and high  
245 contact time (slow pyrolysis) (Inyang and Dickenson, 2015). Hence, it is evident that, if the same  
246 feedstock is used, the ash concentration will be higher in gasification processes than in pyrolysis  
247 (Fryda and Visser, 2015). Moreover, increasing ash percentages will be obtained with increasing  
248 temperature (Rafiq et al., 2016) and higher ash concentrations will be found under fast pyrolysis  
249 conditions (Brewer et al., 2009). Based on these considerations, it makes sense that BC2, BC3, and  
250 BC7, all deriving from slow pyrolysis processes (**Table 1**), showed ash percentages much lower than  
251 BC5 and BC6, which were conversely obtained under gasification conditions using a same patented  
252 process and plant. The very high ash concentration found in BC4 (about 29%) compared to BC3, both  
253 produced with the patented PYREG® pyrolysis process under the same experimental conditions,

254 could be attributed to the different nature of the feedstocks employed, i.e. non woody vegetal biomass  
255 for BC4 and woody waste biomass for BC3 (**Table 1**). Finally, the unexpected high ash content of  
256 BC1 probably depends on the peculiar characteristics of the woody waste used as feedstock, which  
257 derives from the cutting of a forest planted for the phytoremediation of a soil contaminated by  
258 different chemicals, including heavy metals. Virgin activated carbon (AC1) showed a lower ash  
259 content (7%), in agreement with the high standard quality requested by the potabilization plant in its  
260 specifications. Higher ash percentages were obviously found in regenerated and in-use ACs (i.e. AC2  
261 and AC3).

### 262 *3.1.2 Water-extractable substances*

263 The thermal conversion process that transforms biomasses into chars may lead to the formation of  
264 unwanted organic and inorganic hazardous species, depending on the original composition of the  
265 feedstocks. Among them, PAHs (Wang et al., 2017), PCBs (European Biochar Foundation (EBC))  
266 and heavy metals (Lievens et al., 2009) can be present in biochars, thus introducing possible limitation  
267 in the use of the chars themselves. The EN 12915-1 normative regulates the presence of water  
268 extractable pollutants in materials to be applied for water treatments, setting a threshold concentration  
269 limit for the sum of six PAH compounds (i.e. fluoranthene, benzo(b)fluoranthene,  
270 benzo(k)fluoranthene, benzo(a)pyrene, benzo(g,h,i)perylene and indeno-(1,2,3-cd)-pyrene) at 0.02  
271  $\mu\text{g L}^{-1}$ . In addition, the EN standard imposes limits to the presence of As ( $10 \mu\text{g L}^{-1}$ ), Cd ( $0.5 \mu\text{g L}^{-1}$ )  
272  $^1$ ), Cr ( $5 \mu\text{g L}^{-1}$ ), Hg ( $0.3 \mu\text{g L}^{-1}$ ) Ni ( $15 \mu\text{g L}^{-1}$ ), Pb ( $5 \mu\text{g L}^{-1}$ ), Sb ( $3 \mu\text{g L}^{-1}$ ) and Se ( $3 \mu\text{g L}^{-1}$ ). As  
273 regards PCBs, no limit is currently established by the EN standard. However, it should be mentioned  
274 that PCB concentrations are regulated in biochars to be used for soil conditioning and feed additives  
275 (European Biochar Foundation (EBC); International Biochar Initiative, 2015).

276 Results obtained for leachable PAHs showed that all the chars fulfil the limits set by EN 12915-1  
277 regulation. In detail, the regulated PAHs were detected in BC1, BC2, BC3, BC4, and BC7 (all  
278 deriving from pyrolysis), with the sum of their concentrations ranging from  $1.6 \text{ ng L}^{-1}$  (BC4) to  $13.3$

279 ng L<sup>-1</sup> (BC7). Conversely, for ACs and the other BCs, the concentrations of PAHs included in the EN  
280 standard were below detection limits (**Table S3** of the *Supplementary material*).

281 EPA PAHs other than the ones included in the EN standards were also determined (see **Table S3**),  
282 highlighting that PAHs with 2-3 aromatic rings were generally more abundant than those with higher  
283 molecular weight, as also observed elsewhere (Lyu et al., 2016). BC7, which was produced under  
284 pyrolysis at the lowest temperature (450°C), was the material providing by far the highest total  
285 leachable PAH concentration (826 ng L<sup>-1</sup>), while biochars obtained under gasification conditions  
286 showed the lowest PAHs release (11-12 ng L<sup>-1</sup>). These BCs were also the ones providing respectively  
287 the highest and the lowest benzo(a)pyrene equivalent concentrations (BaPy<sub>eq</sub>), based on toxic  
288 equivalence factors (TEFs) available in literature (Berardi et al., 2019). PAHs occurrence in BCs can  
289 be explained based on re-polymerization phenomena of the radical hydrocarbon fragments formed  
290 during the thermal process, which are favoured by the absence of oxygen. Moreover, the presence of  
291 PAHs depends also on the conversion temperature adopted, which plays a main role in PAH  
292 formation up to about 500°C, but also in their degradation beyond this value (Parker et al., 2014; Lyu  
293 et al., 2016). An influence of the biomass composition in the presence of leachable PAHs can also be  
294 evidenced from the comparison of BC3 (321 ng L<sup>-1</sup>) and BC4 (26 ng L<sup>-1</sup>), which were obtained under  
295 exactly the same pyrolysis conditions, but with completely different feedstocks.

296 The concentrations of extractable PCBs found in the BCs and ACs leachate are reported in **Table S4**  
297 of the *Supplementary material*. BC1 was the only char exhibiting the presence of all the PCBs  
298 investigated in its leachate, with total concentration of about 96 ng L<sup>-1</sup>. Conversely, all the other BCs  
299 showed PCB15 as the only leachable chlorinated biphenyl, the concentration of which was in the low  
300 ng L<sup>-1</sup> range (0.56-2.8 ng L<sup>-1</sup>) in BC2, BC3, and BC4 and about one order of magnitude higher (21.7-  
301 29 ng L<sup>-1</sup>) in BC5 and BC6. PCBs occurrence in the BC leachate could be ascribed to the thermal  
302 transformation of chloride, originally contained in feedstocks (Reed) and in this regard, it should be  
303 recalled that BC1 has been prepared with wood waste deriving from a multi-contaminated soil. To

304 the best of our knowledge, this is the first study investigating PCBs in BC leachates, thus preventing  
305 any comparison with literature data.

306 The concentrations of metals determined in BC leachates are illustrated in **Table S5** of the  
307 *Supplementary material*, where the limits set by UNI EN 12915-1 are also reported. BCs generally  
308 complied with the release limits set by the aforementioned regulation, with the only exception of  
309 BC1, which exceeded the limit for Cr, with an observed concentration ( $17.3 \mu\text{g L}^{-1}$ ) three times higher  
310 than this limit, in agreement with the considerations previously reported. As regards ACs, only AC3  
311 exceeded limit for Se ( $5 \mu\text{g L}^{-1}$  versus  $3 \mu\text{g L}^{-1}$ , see **Table S5**), suggesting that a possible saturation  
312 of adsorption sites occurred during service.

### 313 *3.1.3 pH of the point of zero charge*

314 The values of  $\text{pH}_{\text{pzc}}$  were characterized by a high variability (7.0-12.0), in agreement with the wide  
315 range of the production conditions, including the type of feedstock that, as already reported in  
316 literature, significantly influence this property (Ippolito et al., 2020). For BCs, a significant linear  
317 correlation with positive slope was found by plotting ash concentrations as a function of  $\text{pH}_{\text{pzc}}$  values  
318 ( $R^2 = 0.780$ ,  $P < 0.05$ ). This correlation can be ascribed to the ash composition typically reported in  
319 literature (Ippolito et al., 2020), which mainly consists in metals present in the hydroxide form, thus  
320 promoting the increase of the pH in solution. ACs did not follow this trend, as the higher the amount  
321 of ash, the lower the  $\text{pH}_{\text{pzc}}$  observed. This finding is in agreement with the increasing amount of  
322 chemicals other than organic carbon adsorbed by ACs during operation, which however do not  
323 influence the alkalinity of material surfaces.

324 The measurement of the  $\text{pH}_{\text{pzc}}$  values allowed us to make some considerations about the net surface  
325 charge of the chars, which mainly depends on the surface functional groups of the material and is  
326 extremely useful to explain the adsorption behaviours of BCs towards ionized or ionisable  
327 compounds. Most BCs (with the only exception of BC7) exhibited  $\text{pH}_{\text{pzc}}$  higher than pH values of  
328 drinking water collected before entering AC filters of the aforementioned potabilization plants, which  
329 ranged between 7.6 and 7.7. Hence, BC1-BC6 are expected to exhibit positively charged surfaces

330 when they were applied to the treatment of these waters, whereas BC7 is supposed to be negatively  
331 charged (Li et al., 2017).

#### 332 3.1.4 Adsorption indexes of chars

333 The adsorption efficiency of the BCs was evaluated in comparison with ACs through the  
334 determination of  $I_2In$  and  $MBIn$  (**Table 2**).

335  $I_2In$  is commonly considered as related to the presence in the structure of micropores (average  
336 diameter less than 2 nm) and should be therefore informative for the removal efficiency of small-size  
337 organic water pollutants (Del Bubba et al., 2020). Conversely,  $MBIn$  should be associated to the  
338 abundance of mesopores (average diameter in the range 2-50 nm) and thus considered as a useful  
339 indicator of adsorption capacity towards medium-large sized organic pollutants (Del Bubba et al.,  
340 2020). For BCs, the  $I_2In$  was found in the range 77-197 mg  $I_2$  g<sup>-1</sup>, with BCs obtained from gasification  
341 (i.e. BC5 and BC6) showing the highest values (156-197 mg  $I_2$  g<sup>-1</sup>), while the lowest ones (77-88 mg  
342  $I_2$  g<sup>-1</sup>) were exhibited by materials produced under pyrolytic conditions (i.e. BC3 and BC7). The range  
343 determined for  $I_2In$  in BCs was about five times lower than that determined in virgin and regenerated  
344 ACs (i.e. AC1 and AC2), ranging between 540 and 1010 mg  $I_2$  g<sup>-1</sup>. As expected, the  $I_2In$  of AC3 (i.e.  
345 AC2 after some use in the potabilization plant) was lower than that of AC2 (438 mg  $I_2$  g<sup>-1</sup>), in  
346 agreement with the progressive pore saturation phenomena occurring during operation.

347 The  $MBIn$  showed a trend within the BCs and ACs clusters, and among them, similar to that described  
348 for  $I_2In$  (e.g. higher values for ACs than BCs and for BC5 and BC6 compared to the other BCs).  
349 Accordingly, as illustrated by **Figure S1** of the *Supplementary material*, the two indexes showed a  
350 very good linear correlation ( $R^2 = 0.945$ ,  $P \ll 0.05$ ), in agreement with findings observed elsewhere  
351 for different types of ACs and BCs (Del Bubba et al., 2020), even though a lower determination  
352 coefficient was observed by excluding ACs from the correlation ( $R^2 = 0.534$ ,  $P = 0.062$ ). The much  
353 lower correlation was mainly ascribable to the opposite trend observed for some pairs of materials,  
354 such as BC5 and BC6, the latter exhibiting lower  $MBIn$  but higher  $I_2In$  than the former. These findings

355 can be explained by the general differences in micro, meso, and macroporosity distributions of BCs  
356 due to the different experimental conditions adopted for their production.

### 357 3.1.5 Physisorption analysis

358 **Table 2** illustrated the results obtained for the porosimetry analyses (i.e. BET SSA, t-plot MiSSA and  
359 BJH desorption cumulative MeSSA) of the investigated BCs and ACs.

360 As expected, ACs exhibited much higher values of the SSA ( $561\text{-}1053\text{ m}^2\text{ g}^{-1}$ ) than BCs ( $136\text{-}309\text{ m}^2$   
361  $\text{g}^{-1}$ ), being the latter group characterized by a data trend similar to those observed for adsorption  
362 indexes. To elaborate, BC5 and BC6, in addition to showing greater values of the adsorption indexes,  
363 also exhibited the highest SSA, whereas BC7 had the lowest values of the aforementioned parameters.  
364 These findings can be explained by the well-recognized role of temperature in increasing the surface  
365 area (Liu et al., 2010; Ahmad et al., 2014), since BC5/BC6 and BC7 were obtained by the highest  
366 and lowest conversion temperature, respectively (**Table 1**). Indeed, very good linear correlations ( $R^2$   
367  $= 0.981\text{-}0.984$ ,  $P \ll 0.05$ ) were observed between SSA and adsorption indexes (see **Figures S2-A** and  
368 **S2-B** of the *Supplementary material*), even excluding ACs from the regression ( $R^2 = 0.781\text{-}0.904$ ,  
369 data not shown).

370 In general, BCs showed a higher percentage of macroporosity than ACs. However, it should be noted  
371 that BC3 and BC4, produced with the same patented PYREG® pyrolysis system, had a very small  
372 microporosity, comparable to that observed in ACs. The relative percentages of microporosity and  
373 mesoporosity were comparable in all materials, with the exception of AC1, which was strongly  
374 characterized by microporosity. Moreover, high correlations were found between MiSSA and  $I_2/\text{In}$   
375 ( $R^2 = 0.972$ ,  $P \ll 0.05$ ), as well as between the MeSSA and  $\text{MBIn}$  ( $R^2 = 0.922$ ,  $P \ll 0.05$ ) (**Figs. S2-C**  
376 **and S2-D**). However, similarly to findings observed for the correlation between adsorption indexes,  
377 also these relationships were mainly driven by the presence of ACs, since their exclusion strongly  
378 lowered the determination coefficients, making null the significance of the correlation (data not  
379 shown).



380 Differently from adsorption indexes, SSA is often reported as fundamental parameter for the  
381 characterization of sorption properties of BCs. In order to understand the overall significance of SSA  
382 data obtained here, it is therefore interesting to compare them with the values reported in literature  
383 for the numerous biochars obtained from vegetal feedstocks. However, the kind of conversion  
384 process, its temperature and time, as well as the type of biomass used, strongly affects the SSA of  
385 BCs. Accordingly, this comparison was restricted to biochars obtained from woody vegetal  
386 feedstocks, which represent the main type of biomass used for the production of BCs here  
387 investigated, obtaining SSA values in the range 2-637 m<sup>2</sup> g<sup>-1</sup> (Chen et al., 2016; Hansen et al., 2016;  
388 Chen et al., 2019; Grojzdek et al., 2021). Therefore, the SSA values between 136 and 309 m<sup>2</sup> g<sup>-1</sup>  
389 measured in this study are fully in the range reported in the literature.

### 390 3.2 Principal component analysis of the char characterization parameters

391 To summarize the wide group of information discussed above, deriving from the determination of the  
392 several characterization parameters in the ten char samples, a multivariate elaboration of the  
393 autoscaled original data was performed by means of PCA. In more detail, PCA elaboration included  
394 the following eleven parameters: conversion temperature (T), the seven parameters reported in **Table**  
395 **2** (i.e. ash, pH<sub>pzc</sub>, SSA, MiSSA, MeSSA, I<sub>2</sub>In, and MBIn), total PAHs (expressed as BaPy TEF  
396 concentrations), total PCBs, and total metals. Three principal components (PCs), characterized by  
397 eigenvalues > 1 and accounting for percentages of explained variances (E.V.) of 54.9%, 26.4%, and  
398 11.0%, were obtained (total E.V. = 92.3%). **Figure 1** illustrates the plots of scores (**Fig. 1A-B**) and  
399 loadings (**Fig. 1C-D**) of PC1 versus PC2 and PC1 versus PC3, which represent E.V. of 81.3% and  
400 65.9%, respectively. The contributions of each variable to the three significant PCs were not always  
401 well differentiated, even though most original variables showed remarkably different absolute values  
402 of loadings among the three components. In more detail, SSA, MiSSA, MeSSA, I<sub>2</sub>In, and MBIn  
403 contributed mainly in PC1, pH<sub>pzc</sub> and above all ash were mainly represented on PC2, whilst total  
404 PCBs exhibited by far the highest loading on PC3. Conversely, T was represented in PC1 and PC3 to  
405 the same extent, whereas total PAHs and metals contributed almost equally to PC2 and PC3. Among

406 the ten investigated chars, ACs clustered in both score plots, mainly due to their peculiar  
407 characteristics in terms of adsorption indexes and physisorption data. BC5 and BC6, which derived  
408 from the same gasification process, also clustered in both score plots mostly because of the  
409 particularly high values of ash and very low leachable concentrations of total PAHs. Actually, BC6  
410 was the closest char to the AC cluster, suggesting interesting adsorption properties. This consideration  
411 points out that PCA is a valuable tool to select the best sorbents for adsorption measurements, when  
412 one or more reference materials are included in the unsupervised multivariate analysis as  
413 comparators. BC2, BC3, and BC4 identified a further cluster in both score plots. Conversely, BC1  
414 and BC7 behaved as outliers, being they quite distant from the other BCs and the farthest from ACs,  
415 due to their peculiar values of the coordinates on PC2 and PC3. In fact, their scores, were mainly  
416 governed by the concentration values of ash (the highest in BC1 and the lowest in BC7), total PAHs  
417 (intermediate value for BC1 and the highest one for BC7), and total metals (the highest in BC1 and  
418 the lowest in BC7), which strongly contributed to these PCs.

419 In order to have a quantitative confirmation of the findings of PCA, CA was carried out, by using the  
420 complete linkage method and the Euclidean distances on the autoscaled values of the aforementioned  
421 eleven variables (**Figure 2**). The dendrogram confirmed the results of PCA, especially for BC2, BC3,  
422 and BC4, for BC5 and BC6, and for AC1 and AC2, which were grouped in three clusters at similarity  
423 percentages higher than 75%. It is also worth noting that BC5 and BC6 clustered with AC2 and AC3  
424 with a similarity of about 50%. CA also highlighted the high distance between the virgin activated  
425 carbon (AC1) and the regenerated ones (AC1 and AC2), which exhibited a very low degree of  
426 similarity (about 20%).

427 Based on the results of the multivariate characterization of BCs and ACs and their summarising  
428 picture obtained by PCA and CA, BC6 and BC7 were selected for the successive adsorption studies,  
429 as the closest and the farthest materials to the ACs cluster, respectively. Within this latter group, the  
430 virgin (AC1) and the regenerated (AC2) activated carbon were chosen as comparators.

431 3.3 Adsorption studies on DIAA and VOCs

432 Adsorption isotherm experiments were performed on BC6, BC7, AC1, and AC2, using the anion  
433 DIAA and the neutral VOCs benzene and 1,2-dichlorobenzene in order to (i) hypothesize possible  
434 retention mechanisms of BCs (Inyang and Dickenson, 2015) and (ii) estimate sorption capacity of  
435 chars towards these pollutants, which are of environmental concern, *per se*.

### 436 3.3.1 DIAA

437 As regards DIAA, at 5 µg/L, both ACs exhibited a quantitative removal for all the char concentrations  
438 tested, since target analyte was not detected in water solutions after 24 h of contact. Conversely,  
439 removal in the ranges of 18.8-70.8 % and 5.1-28.4% were observed for BC6 and BC7, respectively  
440 (**Table 3**). The different adsorption performances exhibited by BC6 and BC7 should be ascribed not  
441 only to the different surface area, but also to the surface charge, as derived by  $pH_{pzc}$  measures. In fact,  
442  $pH_{pzc}$  tests indicated a significantly higher positive surface charge for BC6 ( $pH_{pzc}=11.0$ ) than BC7  
443 ( $pH_{pzc}=7.0$ ) at the working pH value ( $pH=6.5$ ), which is responsible for electrostatic interactions  
444 between biochar and DIAA (Inyang and Dickenson, 2015).

445 As no detectable DIAA concentrations were found for ACs at 5 µg L<sup>-1</sup>, adsorption isotherm  
446 experiments were repeated using an initial concentration of 20 µg L<sup>-1</sup>. With this concentration, the  
447 removal percentage of the two ACs remained quite similar, ranging approximately from about 89 %  
448 to 100 % in both cases (**Table 3**). As illustrated by **Figure 3**, adsorption data were fitted by the  
449 linearized Freundlich equation, observing in all cases determination coefficients  $\geq 0.925$  and  
450 statistically significant models based on ANOVA ( $P$ -values  $\ll 0.05$ ). Values of  $K_F$  (**Table 3**) for BCs  
451 were about three orders of magnitude lower than those for ACs. Hence, BCs provided a poor  
452 adsorption ability compared to those determined for materials routinely used in water treatment  
453 plants. In more detail, based on the  $K_F$  values, BC6 exhibited sorption ability about 3 times higher  
454 than BC7. A similar efficiency ratio was observed for AC1 vs. AC2, in accordance with the fact that  
455 the former is a virgin material, while the latter is a regenerated char. Slope ( $1/n$ ) values of the  
456 regression lines (**Fig. 3**) were in all cases  $< 1$  (from 0.432 for BC7 to 0.736 for AC1), following the  
457 order  $BC7 \approx BC6 \ll AC2 < AC1$ . The values determined for slopes suggest an L-type isotherm

458 behaviour (European Centre for Ecotoxicology and Toxicology of Chemicals (ECETOC), 2013) for  
459 the adsorption of DIAA on the investigated chars, notwithstanding the pseudo-linear aspect of the  
460 experimental equilibrium concentration data (i.e. X/M vs. C<sub>e</sub>), which is probably related to the quite  
461 high concentrations of material and their narrow range tested here (i.e. about one order of magnitude).  
462 This means that when DIAA concentration increases, the relative adsorption decreases due to the  
463 saturation of adsorption sites available to DIAA, resulting in relatively less intense adsorption, with  
464 increasing the amount of chemical adsorbed onto the material, as commonly observed for the sorption  
465 of organic compounds on chars.

### 466 3.3.2 VOCs

467 Adsorption isotherm experiments on VOCs showed a not negligible variability of data, probably due  
468 to the high vapour pressure and low water solubility of these analytes (Henry constants of  $5.5 \cdot 10^{-4}$   
469 and  $2.3 \cdot 10^{-3} \text{ atm} \cdot \text{m}^3 \text{ mol}^{-1}$ , for benzene and 1,2-dichlorobenzene, respectively). In this regard, it  
470 should be noted that control experiments evidenced losses of both the investigated VOCs (about 15-  
471 25%). Accordingly, adsorption isotherms were not calculated for VOCs. However, it is possible to  
472 state that both BCs showed good adsorption properties, since, at both 20 and 5 µg/L the removal  
473 percentage was almost quantitative for ACs and BC6, and approximately equal to 60-70% and 70-  
474 80%, for BC7 towards benzene and 1,2-dichlorobenzene, respectively.

### 475 3.4 Removal tests in water samples collected in drinking water plants

476 The removal capabilities of BC6 and BC7 were additionally tested in two water samples (i.e. DSB  
477 and CB) collected from the drinking water plant treatment train (before entering the final refinement  
478 stage with activated carbon beds) and compared with those of commercial AC1. These tests were  
479 performed by putting in contact for 24 h 0.4 g of chars with 100 mL of DSB and CB spiked with 20  
480 µg L<sup>-1</sup> of DIAA or VOCs. In such a way, the possible competitive effects exhibited by the matrix can  
481 be assessed and results obtained from adsorption experiments in ultrapure water eventually  
482 confirmed. To better explain possible competitions mechanisms, Total Organic Carbon (TOC) was  
483 initially measured in the two samples, obtaining TOC values of 4.3 mg L<sup>-1</sup> and 2.4 mg L<sup>-1</sup> for CB and

484 DSB, respectively. The lower TOC value observed for DSB should be ascribed to the disinfection  
485 stage operated in this treatment train. The results obtained in these removal tests are summarized in  
486 **Table 4**. As a general consideration, the use of water samples collected within the treatment train of  
487 the potabilization plants did not alter the performance of the BCs, even though the lower  
488 performances of biochars compared to the activated carbon were confirmed. Results obtained in real  
489 water samples fully support the design of column experiments to assess accurately the removal  
490 capacity and exhausting time of BCs, with particular reference to BC6.

#### 491 **4 Conclusions**

492 Within the actions pursued in a circular economy approach fostered by European Union for waste  
493 management, the reuse of waste is promoted for the reduction of resources consumption. Biochar is  
494 one successful example of valorisation of wastes.

495 In this paper, seven BCs obtained from gasification or pyrolysis processes of waste vegetal biomass,  
496 were characterized in depth for numerous parameters, in comparison with a virgin commercial AC, a  
497 freshly regenerated AC, and a regenerated AC in use at a potabilization facility. The characterization  
498 included the evaluation of “environmental concern” parameters (e.g. PAHs and metals release), for  
499 which mandatory limits are provided at European level for materials intended as sorbents for drinking  
500 water filtration, but seldom evaluated elsewhere. Most BCs met these limits, whilst the “sorption  
501 performance parameters” regulated in the European standard (i.e. I<sub>2</sub>In and ash in the UNI EN 12915-  
502 1) were in almost all cases outside the acceptance thresholds, suggesting lower efficiencies compared  
503 to ACs. However, the sorption ability of a given material towards a specific molecule is the result of  
504 a set of characteristics, which all contribute together to the overall removal efficiency, thus suggesting  
505 the importance of following a multivariate approach. Indeed, multivariate analyses performed in this  
506 work (i.e. PCA and CA) allowed for easily identifying the materials with the closest (BC5 and BC6)  
507 or the farthest (BC1 and BC7) characteristics to those of ACs. Accordingly, the multivariate approach  
508 should be promoted in the exploration of data deriving from material characterization, rather than the  
509 evaluation of individual characteristics, even if they are regulated by legislation.

510 Adsorption tests towards DIAA and VOCs carried out in ultrapure water highlighted the much lower  
511 sorption ability of BC7 compared to BC6. These results were in agreement with findings of  
512 multivariate analyses, therefore suggesting intermediate sorption performances for BC1-BC5.  
513 Interestingly, removal tests in waters withdrawn from potabilization plants did not evidence any  
514 significant decrease of the sorption ability of BC6 and BC7 towards the investigated contaminants  
515 compared to tests in ultrapure water, thus supporting the implementation of column experiments for  
516 establishing the maximum loading capacity of the materials in experimental conditions more similar  
517 to the real scale.

518 Even though the sorption performances of BCs are much lower than those of ACs, it should be noted  
519 that BCs did not undergo any physical or chemical activation process, which can surely improve their  
520 removal capacity. Moreover, the management of waste biomass to produce biochar as adsorbent for  
521 water treatment may be regarded as a “win-win” solution for pursuing circular economy principles  
522 and protecting the environment.

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**Table 1** – Status of operation and production conditions of biochars (BC) and commercial activated carbons (AC); n.a. = not available

Sample	Status	Feedstock	Thermal treatment	Temperature (°C)	Contact time (min)
BC1	Virgin	Wood waste mixture <sup>a</sup>	Pyrolysis	550	10
BC2	Virgin	Wood waste mixture <sup>b</sup>	Pyrolysis	550	10
BC3	Virgin	Wood waste mixture <sup>b</sup>	Pyrolysis	550-600	15
BC4	Virgin	Herbal pomace	Pyrolysis	550-600	15
BC5	Virgin	Wood waste mixture <sup>c</sup>	Gasification	800-900	10
BC6	Virgin	Wood waste mixture <sup>d</sup>	Gasification	800-900	10
BC7	Virgin	Corn cob	Pyrolysis	450	30
AC1	Virgin	Coconut	Pyrolysis+physical activation	800-950	n.a
AC2	Regenerated	Coconut	Pyrolysis+physical activation	800-950	n.a
AC3	In use	Coconut	Pyrolysis+physical activation	800-950	n.a

<sup>a</sup> Composition: 100% Poplar; <sup>b</sup> Unknown composition; <sup>c</sup> Approximate composition: Pine 60%, Beech 25%, Hazel 15%; <sup>d</sup> Approximate composition: Pine 40%, Beech 30%, Hazel 20%, Spruce 10%

**Table 2** – Ash (%), pH of the point of zero charge ( $\text{pH}_{\text{pzc}}$ ), specific surface area (SSA, BET method,  $\text{m}^2 \text{g}^{-1}$ ), surface area of micropores (MiSSA, t-plot method,  $\text{m}^2 \text{g}^{-1}$ ), surface area of mesopores (MeSSA, BJH model – desorption cumulative surface area,  $\text{m}^2 \text{g}^{-1}$ ), iodine index ( $\text{I}_2\text{In}$ ,  $\text{mg g}^{-1}$ ), and methylene blue index (MBIn,  $\text{mg g}^{-1}$ ), determined in biochars (BCs) and activated carbons (ACs). For the parameters tested by replicated analyses ( $n=3$ ), mean and standard deviations (in bracket) are reported. Available limits set by European regulation EN 12915-1 are also reported; n.a. = not available.

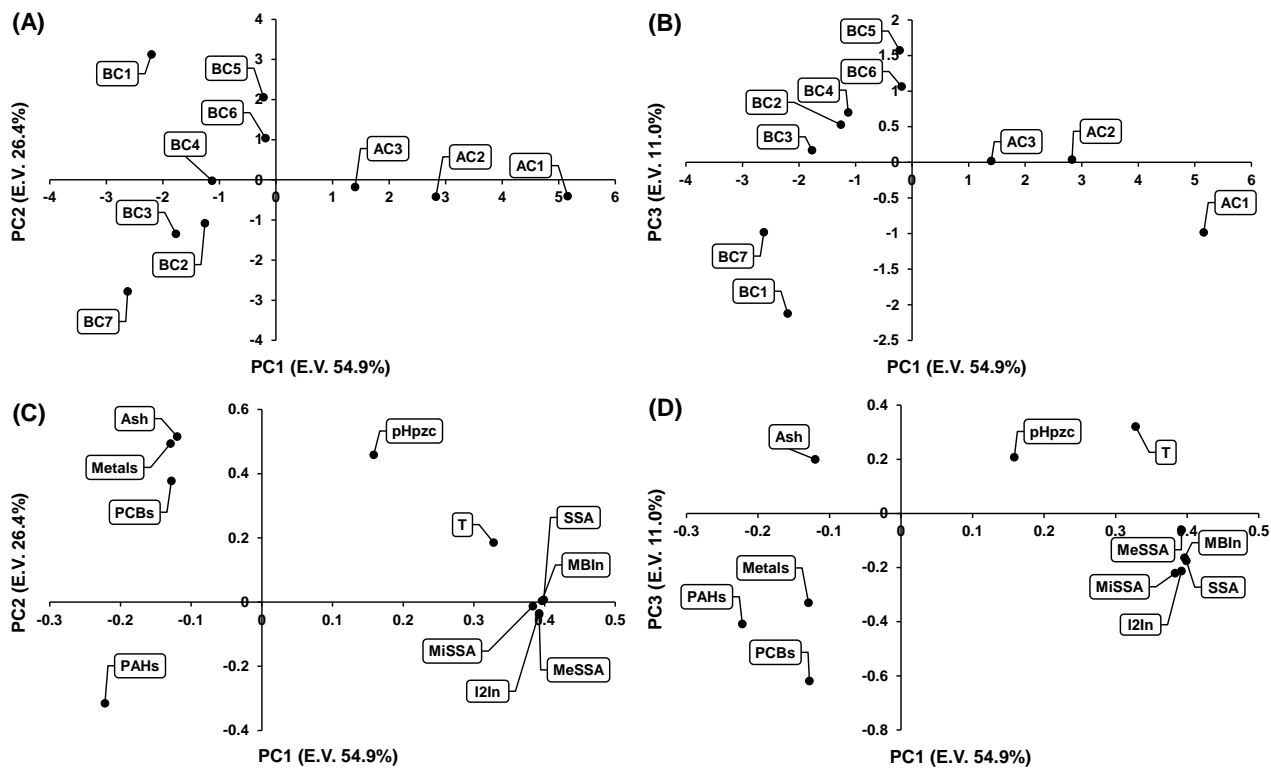
Sample	Ash	$\text{pH}_{\text{pzc}}$	SSA	MiSSA	MeSSA	$\text{I}_2\text{In}$	MBIn
EN 12915-1	15	n.a.	n.a.	n.a.	n.a.	600	n.a.
BC1	42 (8)	10.5	253 (19)	81 (15)	53 (12)	129 (1)	4 (1)
BC2	12 (2)	8.6	243 (22)	43 (11)	63 (14)	144 (1)	4 (2)
BC3	14.1 (0.5)	8.4	153 (20)	65 (14)	76 (17)	88 (1)	1.4 (0.8)
BC4	29.2 (0.2)	9.1	222 (18)	92 (22)	117 (25)	124 (1)	4 (2)
BC5	49 (4)	12.0	302 (23)	121 (26)	97 (20)	156 (1)	6.0 (0.7)
BC6	25 (2)	11.0	309 (21)	80 (19)	136 (30)	197 (1)	4.1 (0.4)
BC7	6.2 (0.1)	7.0	136 (12)	33 (10)	40 (11)	77 (1)	2.2 (0.4)
AC1	7 (2)	10.9	1053 (88)	634 (118)	384 (81)	1010 (1)	20 (2)
AC2	13 (4)	9.9	714 (65)	300 (72)	359 (78)	540 (1)	15 (1)
AC3	21 (8)	8.1	561 (38)	205 (51)	273 (59)	438 (1)	11 (2)

**Table 3** – Sorbent masses of chars (M, g) used in adsorption experiments of diiodoacetic acetic (DIAA), DIAA equilibrium concentrations (Ce, mg L<sup>-1</sup>), DIAA removal (R, %), ratio of the amount of DIAA adsorbed per mass unit of sorbent (X/M, mg g<sup>-1</sup>), and values of the Freundlich constant (K<sub>F</sub>, mg<sup>1-1/n</sup> L<sup>1/n</sup> g<sup>-1</sup>). Initial concentrations of DIAA tested for each char are reported in bracket.

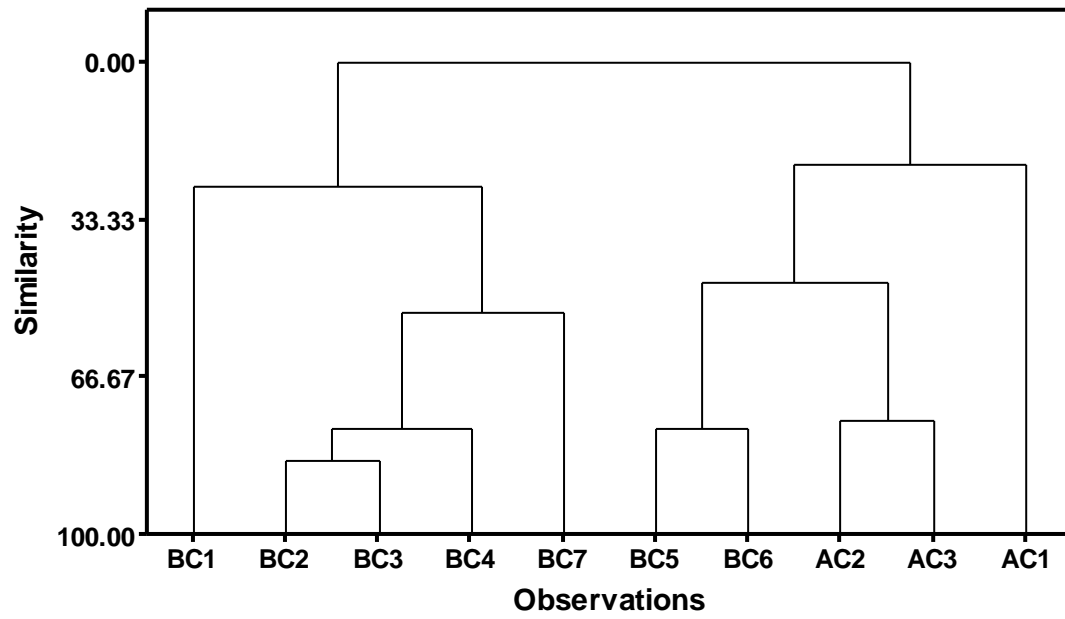
M	Ce	R	X/M	K <sub>F</sub>
<u>BC6 (5 µg L<sup>-1</sup>)</u>				
0.490	0.00146	70.8	0.00029	0.00484
0.324	0.00226	54.8	0.00034	
0.243	0.00272	45.6	0.00038	
0.163	0.00335	33.0	0.00040	
0.113	0.00377	24.6	0.00044	
0.084	0.00406	18.8	0.00045	
<u>BC7 (5 µg L<sup>-1</sup>)</u>				
0.459	0.00358	28.4	0.00012	0.00141
0.304	0.00402	19.6	0.00013	
0.218	0.00426	14.8	0.00014	
0.133	0.00454	9.2	0.00014	
0.103	0.00465	7.0	0.00014	
0.072	0.00475	5.1	0.00014	
<u>AC1 (20 µg L<sup>-1</sup>)</u>				
0.486	0.000032	99.8	0.00164	3.18
0.323	0.000051	99.7	0.00247	
0.244	0.000102	99.5	0.00326	
0.163	0.000147	99.3	0.00487	
0.122	0.000278	98.6	0.00647	
0.033	0.001510	92.5	0.02241	
0.017	0.001970	90.2	0.04242	
<u>AC2 (20 µg L<sup>-1</sup>)</u>				
0.244	0.000028	99.9	0.00327	0.922
0.163	0.000048	99.8	0.00489	
0.122	0.000103	99.5	0.00652	
0.083	0.000208	99.0	0.00954	
0.033	0.001630	91.9	0.02227	
0.016	0.002100	89.5	0.04475	

**Table 4** – Mean values (n=3) and standard deviation (in bracket) of the removal performances (%) of 0.4 g of BC6, BC7, and AC1 towards 20 µg/L of DIAA, benzene, and 1,2-dichlorobenzene (contact time 24 h) in 100 mL-aliquots of two real water samples (DSB and CB) from a potabilization plant, in comparison with ultrapure water (UP). Tests were performed in triplicate. “Q” means quantitative removal, i.e. concentration of the contaminant at the end of the experiment below the detection limit.

	DIAA			Benzene			1,2-Dichlorobenzene		
	UP	DSB	CB	UP	DSB	CB	UP	DSB	CB
<b>BC6</b>	47 (1)	41 (3)	37 (2)	Q	Q	Q	Q	Q	Q
<b>BC7</b>	14 (2)	12 (3)	15 (3)	74 (10)	60 (12)	51 (9)	76 (9)	78 (11)	74 (10)
<b>AC1</b>	99 (1)	99 (4)	99 (3)	Q	Q	Q	Q	Q	Q

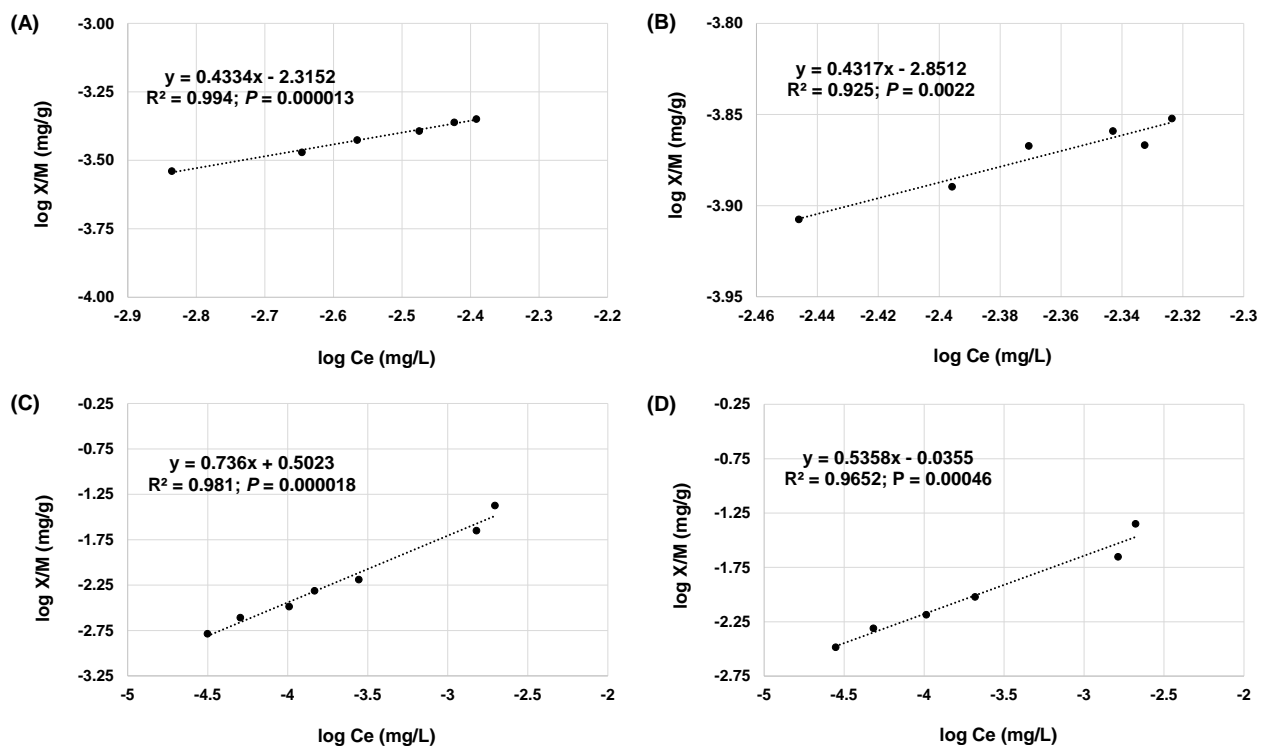


**Figure 1** – Score (A-B) and loading (C-D) plots of PC1 versus PC2 and PC1 versus PC3, representing a percentage of explained variance (E.V.) of 81.3% and 65.9%, respectively. PCA values were calculated using the autoscaled values determined for the eleven original variables in the ten char samples. Note that the terms PAHs, PCBs, and Metals refer to their total leachable concentrations.



**Figure 2** – Dendrogram of similarity of the ten investigated char samples, calculated through the complete linkage method on the basis of Euclidean distances of the autoscaled values of the eleven original variables.





**Figure 3** – Plots of linearized Freundlich isotherms obtained for BC6 (A), BC7 (B), AC1 (C), and AC2 (D).

## SUPPLEMENTARY MATERIAL

### **Biochars intended for water filtration: a comparative study with activated carbons of their physicochemical properties and removal efficiency towards neutral and anionic organic pollutants**

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## S1. Biochar and activated carbon characterization

The **physisorption analysis** of chars were performed by nitrogen adsorption and desorption experiments at  $-196^{\circ}\text{C}$  using a Porosity Analyser Thermo Fisher (Milan, Italy) model SORPTOMATIC 1990. The surface area was calculated by using the Brunauer-Emmet-Teller (BET) and the Langmuir methods applied to nitrogen adsorption data in the relative pressure ( $P/P^{\circ}$ ) range of 0.05-0.25. The total pore volume was determined from the amount of nitrogen adsorbed at  $P/P^{\circ}=0.98$ . The mesopore size distribution was determined by the Barret-Joyner-Halenda (BJH) model applied to desorption data and the assessment of microporosity was carried out by the t-plot method. For the porosimetry of biochars, a minimum equilibrium interval of 30 s with a maximum relative tolerance of 5% of the targeted pressure and an absolute tolerance of 5 mmHg were used. Before analysis, materials were degassed in situ, by heating at  $300^{\circ}\text{C}$ , at a rate of  $5^{\circ}\text{C min}^{-1}$ , under a high vacuum ( $<10^{-8}$  mbar) for 12 h, provided by an oil sealed mechanical vacuum pump coupled with a high vacuum system. For the porosimetry a minimum equilibrium interval of 10 s with a maximum relative tolerance of 2% of the targeted pressure and an absolute tolerance of 2 mmHg were adopted.

The **ash content procedure** is provided by the ASTM International D 2866-11: a known amount of sorbent is placed in a muffle at  $650^{\circ}\text{C}$  for 1 hour and then weighted. According to the UNI EN 12902 standard, ashes should be below 15%.

The **pH at the zero-charge point ( $\text{pH}_{\text{pzc}}$ )** was determined according to the pH drift method: 10 aliquots of 20 mL of 0.01 mol/L  $\text{KNO}_3$  solution, whose pH was corrected in a range from 2 to 12, were prepared. 0.12 g of sorbent was added to each solution and, after stirring for 24 hours, the pH value was measured. To calculate the  $\text{pH}_{\text{pzc}}$ , the initial pH value of each solution was plotted against the final pH value: the  $\text{pH}_{\text{pzc}}$  corresponds to the intersection of the curve so obtained with the x axis.

The **extractable substances**, i.e. metals, Polycyclic Aromatic Hydrocarbons (PAHs) and Polychlorinated Biphenyls (PCBs), were determined for each sorbent following the UNI EN 12902 standard: 1 L of a solution containing  $\text{NaHCO}_3$  (0.5 mmol/L),  $\text{CaCl}_2$  (0.3 mmol/L), and  $\text{MgSO}_4$  (0.2 mmol/L), at pH 7, was put in contact for 24 hours with 10 g of substrate. The content of Sb, As, Cd, Cr, Pb, Hg, Ni and Se was measured by ICP-MS, whereas PAHs and PCBs were determined by GC-MS. The analysis of a blank solution, without any material, was simultaneously performed.

For metals, a calibration curve was built from 0.25 to 20  $\mu\text{g/L}$ ; all the samples were acidified by 2% (v/v)  $\text{HNO}_3$  before the ICP-MS analysis.

For the determination of PAHs and PCBs, the extracted solution was spiked with 23 ppt of labelled PAHs and PCBs (for apparent recovery calculation) and then pre-concentrated by solid phase extraction (SPE), using reversed phase cartridges according to *Environmental Engineering and Management Journal*, 18(10), 2019, Pages 2133-2143, and then analyzed by GC-MS.

The following operating conditions were adopted for GC-MS analysis. Column: HP 5MS UI (5% - phenyl)-methylpolysiloxane; Gas Carrier: He (1 mL/ min); Injection type: Pulsed Splitless; Injection pressure: 40 psi for 2.5 minutes; Injection volume 2 µl. The oven ramp was set as follows: starting temperature: 40°C, hold for 2 min; ramp to 176 °C, 12 °C/min rate; ramp to 196°C, 5 °C/min rate, hold for 3 mins; ramp to 224°C, 12 °C/min rate; ramp to 244 °C, 12°C/min rate, hold for 3 min; ramp to 270 °C, 7°C/min rate, hold for 3 min; final ramp to 300 °C, 5°C/min, hold for 10 min to completely clean and restore the GC column. The total run time for the complete separation of PAHs and PCBs is 52 min.

The **iodine index** was determined according to CEFIC test method. In detail, an aliquot of substrate (previously dried at 150 ° C until a constant mass is obtained) was boiled for 30 seconds in a flask with 10 mL of 5% HCl. Subsequently, 100 mL of a standardized 0.1N iodine solution (titrated by 0.1 N sodium thiosulfate) were added and the flask was shaken vigorously for about 30 seconds. The solution was immediately filtered on a gravity paper filter (first 20 mL aliquot is discarded).

Aliquots of 50 mL of the filtered solution were titrated with 0.10 M sodium thiosulfate until the yellow color disappeared after addition of 2 mL of a starch solution titration was continued until the blue color disappeared. The final volume of the sodium thiosulphate solution used was noted to evaluate the mg of iodine adsorbed.

The iodine number was calculated using the following formula:

$$I_n = \frac{X}{M} A$$

- *X = mg of iodine adsorbed by the material*
  - $X = (12693 N_1) - (279.246 N_2 V)$  where  $N_1$ : normality of iodine solution,  $N_2$ : normality of thiosulphate solution;  $V$  is the volume of sodium thiosulphate solution in mL.
- $M$  is the mass of activated carbon in g;
- $A$  is the correction factor depending on the residual normality ( $N_r$ ) of the filtrate and that is reported in a conversion table of CEFIC test method. This factor  $A$  may be applied if  $N_r$  is between 0.008 and 0.0334, when  $N_r = N_2 (V/50)$ . If  $N_r$  is outside the range, the determination must be repeated.

**The methylene blue index** was determined according to CEFIC test method and represents the volume (mL) of a methylene blue solution (1.2 g/L) adsorbed by 0.1 g of sorbent placed in contact with it.

A methylene blue reference solution (C=1.2 g/L) was prepared, and its absorbance was measured diluting 5 mL of this solution with 0.25% (v/v) acetic acid to 1L. Absorbance measured at 620 nm should be 0.840±0.01; if higher, the reference solution should be diluted with ultrapure water to obtain the expected value.

Subsequently, 100 mg of dried sorbent were added to 25 mL of the 1.2g/L reference solution. The suspension is stirred for 30 minutes, filtered on a paper filter, discarding the first 5 mL of the filtrate and 1 mL of the filtered solution was diluted 1:100 in 0.25% acetic acid solution. Simultaneously, the reference solution was diluted 1:100 in 0.25% acetic acid. The absorbance of both solutions was measured at 620 nm.

The methylene blue index  $x$  is calculated according to:

$$x = \frac{ABS0 \times 25}{abs1} ;$$

- ABSO is the absorbance of methylene blue adsorbed by the sample, calculated as abs1 – abs2;
- abs 1 is the absorbance of the reference solution, diluted 1:100;
- abs 2 is the absorbance of the filtrate, diluted 1:100;

## **S2. Adsorption isotherms: determination of the residual concentrations of diiodoacetic acid by IC-MS/MS**

Diiodoacetic acid was determined by IC-MS/MS using a Thermo Fisher Scientific (Waltham, MA USA) ICS-5000 IC system. The system includes a DP dual pump module for analytical and capillary applications, a CD conductivity detector, an AS autosampler, and a Reagent-Free (RFIC) eluent generator EG-5000 with ECG III cartridges KOH to provide the gradient of KOH (mobile phase) using deionized water from an AXP-MS pump (Thermo Fisher Scientific). For sample injections (120 µL), two autosamplers without (AS-DV) and with sample tray temperature control (AS-AP) set at 9±1 °C were used; both were from Thermo Fisher Scientific. Separations were performed on an IonPac AS24 (250x2 mm i.d.) coupled with a guard column IonPac AG24 (50x2 mm i.d.) both from Thermo Fisher Scientific, thermostatted at 15 °C in order to minimize the analyte degradation at high pH values. Eluent gradient (0.3 mL/min) was set as follows, 7 mM KOH: t=0-15.1 min; 7-15.5 mM KOH: t=15.1-25.8 min; 60 mM KOH: t=25.9 min, keep until 46 min; 7 mM KOH; t=47-58 min.

To remove trace anion contaminants from hydroxide eluent and to minimize base line shifts during gradient operation, an electrolytically continuously regenerated trap column (CR-ATC, 8% DVB crosslinking, 55 µm particle size) was installed in the eluent line after the pump prior to the sample injection. After eluent generation and before the separation column. Electrolytic suppression was accomplished using an ASRS 500 (2-mm) from Thermo Fisher Scientific.

A TSQ Endura triple-stage quadrupole mass spectrometer with ESI interface (HESI-II) was employed for detection. A diverter valve was used to waste the anion interfering species from matrix, thus preventing inorganic anions to enter the MS equipment. After the IC suppressor and before the ESI inlet, acetonitrile (CH<sub>3</sub>CN) was added to the eluate at 0.3 mL/min through an additional AXP-MS pump. The m/z ratios of the precursor ion were 184.878 and 126.889, respectively. RF lens value was 51.236 V with CE 10.253 V.

### S3. Adsorption isotherms: determination of the residual concentrations of VOCs by SPE-GC/MS

Benzene and 1,2-dichlorobenzene were extracted by solid-phase extraction (SPE) using a SPE Vacuum manifold and a polymeric reversed-phase cartridge (STRATA XL-100  $\mu$ m, Phenomenex, Torrance, USA). The cartridge was firstly conditioned (20 psi) with 5 mL CH<sub>2</sub>Cl<sub>2</sub>, 5 mL 2-propanol, 5 mL H<sub>2</sub>O, and then loaded (50 psi) with 100 ml of water sample. The water sample container was subsequently rinsed with 20 mL of a 2-propanol-water (10:90, v/v) solution. After loading, the cartridge was washed (20 psi) with 5 ml H<sub>2</sub>O, and 5 mL of a 2-propanol-water (85:15, v/v) solution. The cartridge was dried for 10 min, capped, and analytes were finally eluted with two aliquots of 1.0 mL CH<sub>2</sub>Cl<sub>2</sub>. The eluted extract was finally diluted 50 times and injected for GC/MS analysis.

The following operating conditions were adopted for GC-MS analysis. Column: HP 5MS UI (5% - phenyl)-methylpolysiloxane; gas carrier: He (1.5 mL/min); Injection type: Pulsed Splitless; Injection pressure: 40 psi for 2.5 minutes: Injection volume 2  $\mu$ l.

The oven ramp was set as follows: starting temperature: 35°C, hold for 2 min; ramp to 50 °C, 4 °C/min rate; ramp to 120°C, 10 °C/min rate; ramp to 220°C, 30 °C/min rate, hold for 3 min. Retention times were 2.8 min for benzene and 10.8 min for 1,2-dichlorobenzene. Quantitative analysis were performed in SIM mode at the following m/z values: 78 (benzene) and 146 (1,2-dichlorobenzene).

**Table S1** – Apparent recoveries of the SPE procedure for labelled PAHs and PCBs from the seven samples of biochars.

	BC 1	BC 2	BC 3	BC 4	BC 5	BC 6	BC 7
<sup>13</sup> C <sub>6</sub> -BaA	91.2 (0.1)	85.7 (1.6)	23.8 (4.3)	76.7 (3.6)	46.7 (2.1)	51.0 (3.8)	13.7 (1.1)
<sup>13</sup> C <sub>6</sub> -Chr	95.3 (0.5)	89.7 (1.1)	46.4 (2.9)	83.7 (7.0)	76.8 (1.4)	81.1 (2.2)	20.5 (2.2)
<sup>13</sup> C <sub>6</sub> -BbFl	76.0 (0.4)	83.3 (0.2)	19.4 (5.8)	73.5 (7.1)	61.2 (3.1)	76.4 (4.1)	15.4 (3.2)
<sup>13</sup> C <sub>6</sub> -BkFl	83.0 (3.5)	82.9 (2.1)	31.6 (1.9)	73.5 (4.3)	60.1 (2.5)	77.0 (3.6)	23.8 (3.3)
<sup>13</sup> C <sub>4</sub> -BaP	67.9 (4.3)	72.6 (1.5)	24.8 (7.3)	63.0 (4.5)	58.6 (3.0)	68.3 (2.8)	16.2 (0.4)
<sup>13</sup> C <sub>4</sub> -Ind	68.7 (6.8)	65.5 (1.6)	12.4 (3.1)	27.9 (0.1)	33.1 (2.6)	51.4 (3.1)	12.8 (1.1)
<sup>13</sup> C <sub>6</sub> -DBA	53.2 (2.1)	49.0 (1.3)	11.9 (2.4)	23.2 (1.7)	33.5 (2.9)	46.8 (1.1)	14.8 (4.3)
<sup>13</sup> C <sub>12</sub> -BP	57.8 (3.5)	48.5 (1.2)	14.7 (2.4)	27.4 (1.4)	36.7 (2.7)	45.8 (2.0)	14.2 (4.0)
<sup>13</sup> C <sub>12</sub> -PCB28	93.6 (2.8)	96.6 (4.0)	55.0 (9.5)	89.1 (3.8)	77.3 (3.1)	83.1 (4.1)	33.2 (4.5)

<sup>13</sup> C <sub>12</sub> -PCB52	90.6 (4.7)	97.3 (2.2)	60.6 (10.9)	92.4 (2.7)	70.8 (1.8)	76.7 (6.4)	37.7 (9.4)
<sup>13</sup> C <sub>12</sub> -PCB118	93.5 (0.7)	76.2 (2.1)	32.8 (2.3)	58.5 (5.2)	55.2 (4.1)	61.0 (3.7)	11.5 (5.7)
<sup>13</sup> C <sub>12</sub> -PCB153	86.9 (1.4)	79.4 (3.1)	35.8 (1.1)	59.8 (6.8)	54.7 (2.5)	58.8 (2.0)	13.3 (6.4)
<sup>13</sup> C <sub>12</sub> -PCB180	71.0 (3.5)	75.2 (1.8)	24.5 (8.2)	52.0 (13.0)	48.1 (2.8)	61.0 (4.4)	9.0 (3.6)

**Table S2** – Apparent recoveries of the SPE procedure for labelled PAHs and PCBs from the three samples of activated carbons.

	AC1	AC2	AC3
<sup>13</sup> C <sub>6</sub> -BaA	60.6 (14.4)	96.3 (5.0)	61.9 (14.6)
<sup>13</sup> C <sub>6</sub> -Chr	75.2 (20.6)	94.5 (6.4)	76.0 (20.6)
<sup>13</sup> C <sub>6</sub> -BbFl	49.8 (11.6)	51.4 (2.7)	55.1 (12.9)
<sup>13</sup> C <sub>6</sub> -BkFl	59.0 (13.3)	52.8 (2.1)	58.0 (13.6)
<sup>13</sup> C <sub>4</sub> -BaP	39.5 (9.2)	48.3 (2.2)	48.2 (11.3)
<sup>13</sup> C <sub>4</sub> -Ind	13.8 (2.5)	47.3 (0.2)	16.8 (3.7)
<sup>13</sup> C <sub>6</sub> -DBA	13.8 (2.0)	35.4 (3.2)	19.9 (4.6)
<sup>13</sup> C <sub>12</sub> -BP	12.7 (1.1)	34.4 (0.5)	18.9 (4.4)
<sup>13</sup> C <sub>12</sub> -PCB28	43.6 (6.4)	76.2 (1.0)	56.4 (12.5)
<sup>13</sup> C <sub>12</sub> -PCB52	30.5 (3.5)	72.7 (12.6)	55.0 (14.1)
<sup>13</sup> C <sub>12</sub> -PCB118	16.6 (3.3)	49.5 (12.4)	37.6 (11.2)
<sup>13</sup> C <sub>12</sub> -PCB153	13.7 (3.4)	55.2 (5.3)	28.2 (8.6)
<sup>13</sup> C <sub>12</sub> -PCB180	12.1 (2.7)	52.3 (5.9)	19.5 (5.5)

The evaluation of extracted PAHs and PCBs was preceded by the determination of SPE extraction recovery of labelled surrogates from each char. Apparent recoveries were calculated after spiking the extracted solutions with labelled surrogates and were expressed as average of two replicated SPE extractions on two spiked aliquots of char. Hence, averages are referred to n=14 and n= 6 data for the seven biochar, and for the three activated carbons, respectively.

For biochars, average recoveries were 50.7% for PAHs (relative standard deviation, RSD, below 7.0%) and 62.1% for PCBs (RSD below 7.0%). For activated carbons, average recoveries were 47.4% for PAHs (RSD below 21%) and 41.3% for PCBs (RSD below 13%).

It is interesting to note that for both PAHs and PCBs, the average of apparent extraction recoveries for ACs is lower than that observed for BCs. This may be ascribed to a higher matrix effect exhibited by ACs due to possible leaching from the material. This hypothesis is supported by the higher RSD values observed for ACs with respect to BCs and by the total organic carbon measurements (see *Adsorption tests in real water samples* section) performed on selected BC and AC samples.

Additionally, for both the classes of biochar and activated carbon, extraction recoveries increase with the decrease of log P of analytes (data not shown), in agreement with the lower competitive effect exhibited by the reversed-phase cartridge used during SPE experiments.

#### S4. Release of PAHs, PCBs, and metals from biochars and activated carbons

**Table S3** – Extractable PAHs from the chars studied. Concentrations are expressed in ng/L. The sum of concentrations of asterisked PAHs (regulated by the UNI EN 12915-1) is also reported. Regulated limit for these compounds is 20 ng/L.

	BC1	BC2	BC3	BC4	BC5	BC6	BC7	AC1	AC2	AC3
Naphthalene	0.55 (0.01)	< LOD	21.9 (0.97)	4.15 (0.07)	6.64 (20)	7.2 (0.13)	83.14 (3.69)	2.2 (0.09)	12.5 (0.45)	10.91 (0.24)
Acenaphthylene	< LOD	< LOD	0.99 (0.03)	1.18 (0.08)	0.56 (0.07)	0.94 (0.01)	6.22 (0.18)	< LOD	< LOD	< LOD
Acenaphthene	0.9 (0.5)	< LOD	3.77 (0.06)	0.94 (0.09)	2.04 (0.09)	2.2 (0.53)	33.34 (3.17)	< LOD	2.73 (0.07)	3.03 (0.15)
Fluorene	< LOD	3.56 (0.41)	83.26 (3.42)	8.34 (0.2)	< LOD	< LOD	261.55 (13.33)	1.04 (0.03)	2.34 (0.11)	3.29 (0.21)
Phenanthrene	15 (4)	34.47 (0.75)	164.25 (5.56)	8.00 (0.03)	< LOD	< LOD	329.54 (19.31)	7.04 (0.03)	4.92 (0.21)	7.52 (0.15)
Anthracene	1.7 (0.01)	4.36 (0.06)	36.77 (0.94)	0.88 (0.07)	< LOD	< LOD	72.03 (2.84)	1.99 (0.17)	2.02 (0.28)	2.69 (0.03)
Fluoranthene*	3.37 (0.01)	7.21 (0.07)	3.41 (0.20)	1.64 (0.03)	< LOD	< LOD	12.71 (0.3)	< LOD	< LOD	< LOD
Pyrene	3.09 (0.48)	6.36 (0.14)	5.56 (0.03)	0.98 (0.01)	2.05 (0.15)	1.45 (0.07)	26.64 (0.8)	< LOD	0.87 (0.06)	1.11 (0.07)
Benzo[a]pyrene	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD
Chrysene	0.96 (0.14)	< LOD	1.34 (0.09)	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD
Benzo[b]fluoranthene*	1.95 (0.42)	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD
Benzo[k]fluoranthene*	1.53 (0.30)	< LOD	< LOD	< LOD	< LOD	< LOD	0.55 (0.11)	< LOD	< LOD	< LOD
Benzo[g,h,i]perylene*	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD
Indeno[1,2,3-c,d]pyrene*	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD
Dibenzo[a,h]anthracene	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD
Benzo[g,h,i]perylene*	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD
<b>Σ* UNI EN 12915-1</b>	6.85	7.21	3.41	1.64	< LOD	< LOD	13.26	< LOD	< LOD	< LOD
<b>Σ</b>	29.39	55.96	321.25	26.11	11.29	11.79	825.72	12.27	25.38	28.55
<b>Σ (Benzo[a]pyrene eq. conc.)</b>	0.398	0.095	0.664	0.034	0.011	0.012	1.528	0.030	0.044	0.053

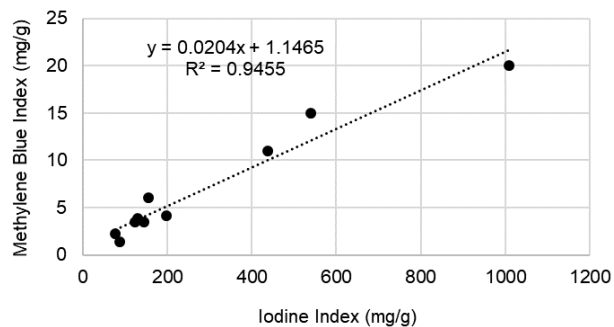


**Table S4** – Extractable PCBs from the chars studied. Concentrations are expressed in ng/L.

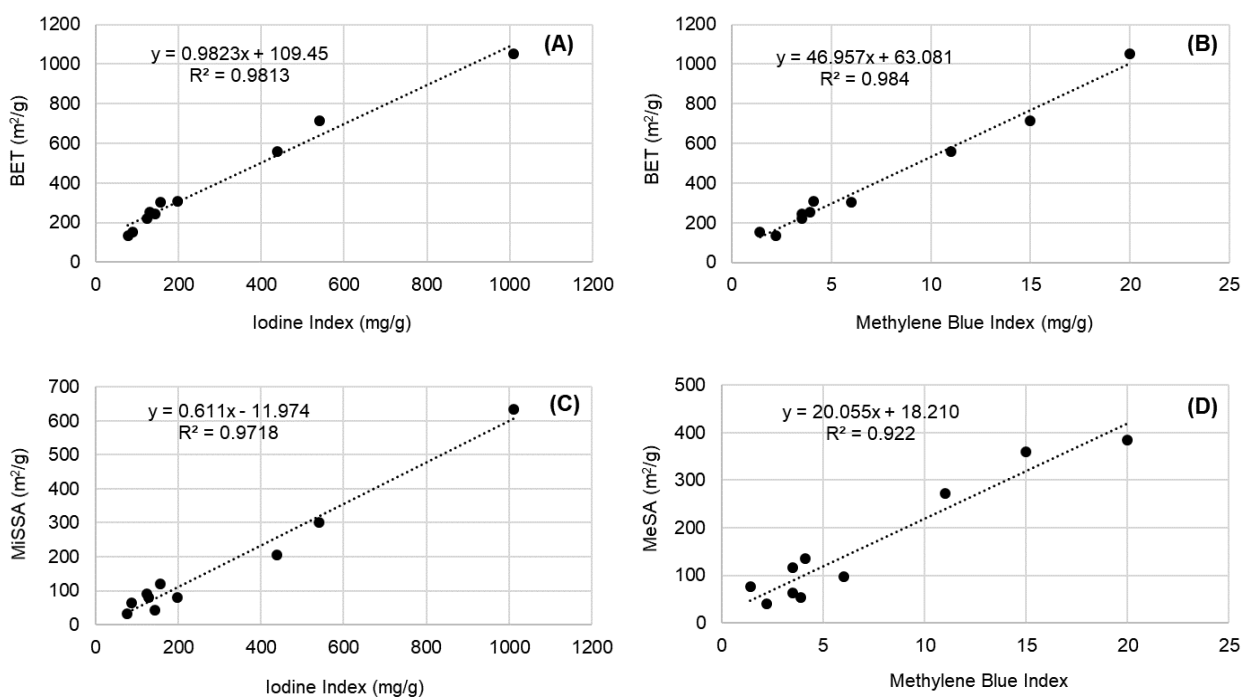
	<b>BC1</b>	<b>BC2</b>	<b>BC3</b>	<b>BC4</b>	<b>BC5</b>	<b>BC6</b>	<b>BC7</b>	<b>AC1</b>	<b>AC2</b>	<b>AC3</b>
<b>PCB15</b>	0.5 (0.1)	0.56 (0.05)	1.29 (0.02)	2.8 (0.1)	21.7 (0.1)	29 (1)	20 (2)	< LOD	4 (2)	1.1 (0.2)
<b>PCB101</b>	6.3 (0.2)	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD
<b>PCB81</b>	8 (2)	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD
<b>PCB118</b>	7 (2)	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD
<b>PCB123</b>	12 (3)	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD
<b>PCB153</b>	16 (4)	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD
<b>PCB167</b>	21 (5)	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD
<b>PCB180</b>	25 (7)	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD
<b>Σ</b>	95.8	0.56	1.29	2.8	21.7	29	20	< LOD	4	1.1

**Table S5** – Extractable metals from the chars studied. Concentrations are expressed in µg/L. The limits established by the UNI EN 12915-1 are also reported.

	<b>BC1</b>	<b>BC2</b>	<b>BC3</b>	<b>BC4</b>	<b>BC5</b>	<b>BC6</b>	<b>BC7</b>	<b>AC1</b>	<b>AC2</b>	<b>AC3</b>	<b>UNI EN limits</b>
<b>Cd</b>	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	0.5
<b>As</b>	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	10
<b>Cr</b>	<b>17.3 (0.1)</b>	< LOD	< LOD	2.8 (0.1)	3.1 (0.1)	3.45 (0.07)	< LOD	< LOD	< LOD	< LOD	5
<b>Ni</b>	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	15
<b>Pb</b>	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	5
<b>Sb</b>	< LOD	< LOD	0.73 (0.05)	< LOD	1.07 (0.01)	2.68 (0.05)	< LOD	< LOD	< LOD	< LOD	3
<b>Se</b>	< LOD	< LOD	< LOD	< LOD	1.01 (0.02)	0.55 (0.08)	< LOD	< LOD	< LOD	<b>5.0 (0.2)</b>	3
<b>Hg</b>	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	< LOD	0.3



**Figure S1** – Dispersion plot and least square regression line of I<sub>2</sub>In vs. MBIn.



**Figure S2** – Dispersion plots and least square regression lines of (A) BET specific surface area vs. I<sub>2</sub>In, (B) BET specific surface area vs. MBIn, (C) micropore specific surface area vs. I<sub>2</sub>In, and (D) mesopore specific surface area vs. MBIn.