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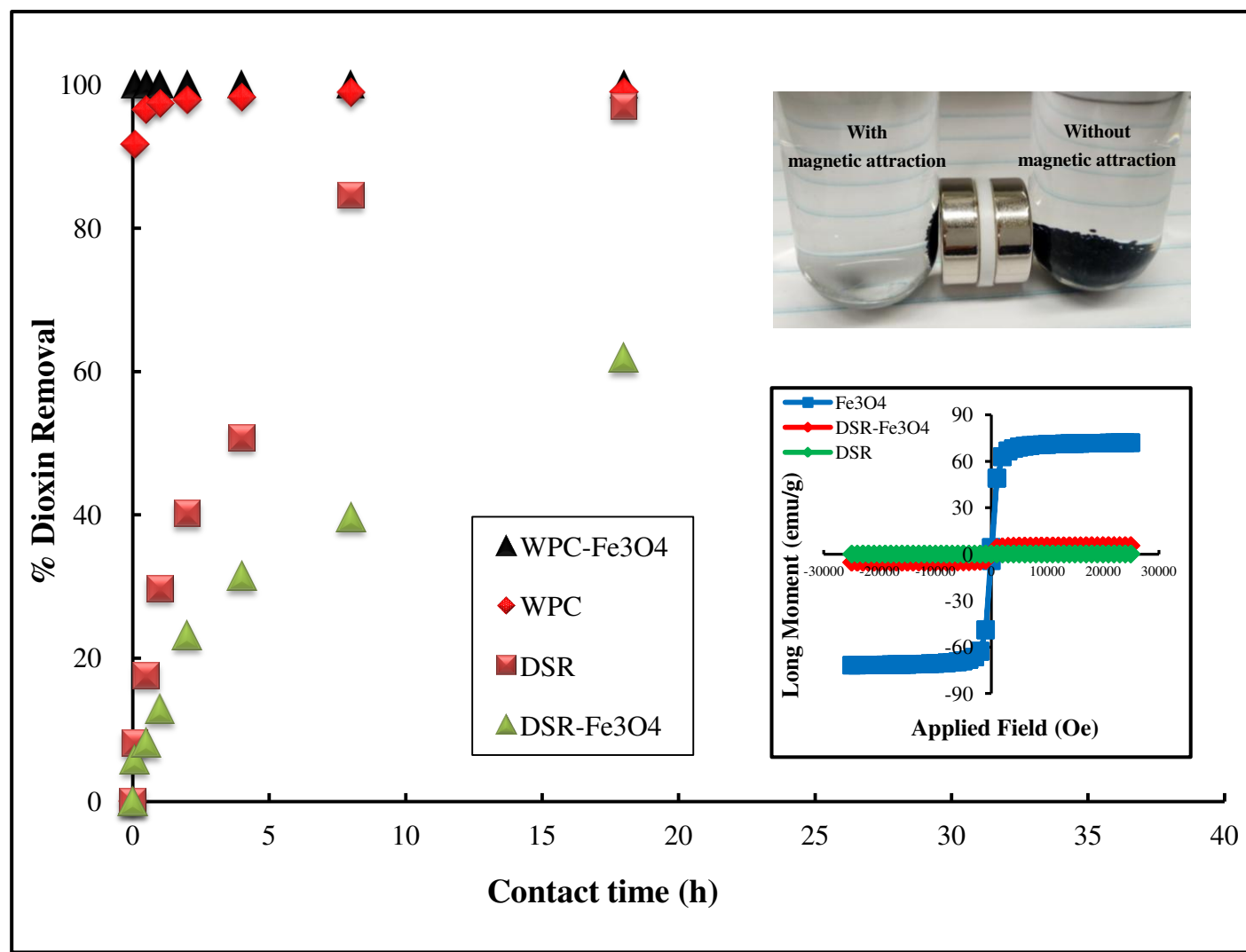
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# **Synthesis and evaluation of Fe<sub>3</sub>O<sub>4</sub>-impregnated activated carbon for dioxin removal**

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20 **Abstract**

21 Polychlorinated dibenzo-p-dioxins and -furans (PCDD/PCDFs) are highly toxic organic  
22 pollutants in soils and sediments which persist over timescales that extend from decades to  
23 centuries. There is a growing need to develop effective technologies for remediating  
24 PCDD/Fs-contaminated soils and sediments to protect human and ecosystem health. The use  
25 of sorbent amendments to sequester PCDD/Fs has emerged as one promising technology. A  
26 synthesis method is described here to create a magnetic activated carbon composite  
27 (AC-Fe<sub>3</sub>O<sub>4</sub>) for dioxin removal and sampling that could be recovered from soils using  
28 magnetic separation. Six AC-Fe<sub>3</sub>O<sub>4</sub> composites were evaluated (five granular ACs (GACs)  
29 and one fine-textured powder AC(PAC)) for their magnetization and ability to sequester  
30 dibenzo-p-dioxin (DD). Both GAC/PAC and GAC/PAC-Fe<sub>3</sub>O<sub>4</sub> composites effectively  
31 removed DD from aqueous solution. The sorption affinity of DD for GAC-Fe<sub>3</sub>O<sub>4</sub> was slightly  
32 reduced compared to GAC alone, which is attributed to the blocking of sorption sites. The  
33 magnetization of a GAC-Fe<sub>3</sub>O<sub>4</sub> composite reached 5.38 emu/g based on SQUID results,  
34 allowing the adsorbent to be easily separated from aqueous solution using an external  
35 magnetic field. Similarly, a fine-textured PAC-Fe<sub>3</sub>O<sub>4</sub> composite was synthesized with a  
36 magnetization of 9.3 emu/g.

37

38 Keywords: dibenzo-p-dioxin, granular activated carbon, Fe<sub>3</sub>O<sub>4</sub>, magnetic separation,

39 activated carbon-Fe<sub>3</sub>O<sub>4</sub> composite

## 40 **1. Introduction**

41 Polychlorinated dibenzo-p-dioxins (PCDDs) are prototypical persistent organic pollutants  
42 (POPs) that are commonly found in soils and sediments. Due to their exceptionally low water  
43 solubilities, these highly toxic lipophilic substances are highly bio-accumulative (Guruge et  
44 al., 2005; Maier et al., 2016; Champoux et al., 2017). Exposure to PCDDs, even at trace  
45 concentrations (Denison et al., 1989; Eljarrat et al., 2002), can result in measurable toxic and  
46 carcinogenic effects in mammals (Huwe, 2002; McKay, 2002; Charnley and Doull, 2005).  
47 PCDDs occur both naturally and from anthropogenic activities which include forest fires,  
48 coal combustion, iron ore sintering, chlorine bleaching of pulp and paper, waste incineration,  
49 and as by-products of pesticide manufacturing and the chlor-alkali process (Fiedler, 1996;  
50 Everaert and Baeyens, 2002; Kulkarni et al., 2008; Zheng et al., 2008; Zhou et al., 2016;  
51 Prisciandaroa et al., 2017; Zhao et al., 2017). Owing to their lipophilicity, PCDD/Fs  
52 accumulate in surface soils, sediments and biota, including the fatty tissues of fish (WHO,  
53 2010). In natural environments, they occur predominantly in the sorbed state associated with  
54 pyrogenic carbonaceous matter (PCM), amorphous organic matter, and clays (Ferrario et al.,  
55 2000; Fabietti et al., 2010). In fact, the significant role of PCM as a sorption domain has been  
56 well established (Cornelissen et al., 2005). As a group, PCDD/Fs are characterized by low  
57 aqueous solubilities and high octanol-water coefficients K<sub>ow</sub> (Shiu et al., 1988; Kim et al.,

58 [2002; Li et al., 2009](#)). Consequently, their concentrations in natural waters are extremely low  
59 with concentration ranges of pg/L to fg/L ([Charlestra et al., 2008; Cornelissen et al., 2008b;](#)  
60 [Louchouarn et al., 2018](#)). PCDD/F-contaminated soils are found in ecosystems worldwide  
61 ([Masunaga et al., 2001; Moon et al., 2008; Zheng et al., 2008](#)), and have proven difficult and  
62 expensive to remediate. For example, the estimated cleanup cost of a *single* Superfund site  
63 along the Passaic River which is contaminated by PCDD/Fs has exceeded one billion US  
64 dollars.

65 Traditional site remediation has relied on removal of the contaminants via excavation or  
66 dredging followed by disposal in a hazardous waste landfill. Recently, sorbent amendments  
67 have gained attention as a means to lower or even eliminate bioavailability of soil/sediment-  
68 bound contaminants ([Ghosh et al., 2011; Cornelissen et al., 2012; Hale et al., 2012;](#)  
69 [Cornelissen et al., 2016; Cho et al., 2017](#)), and this has formed the basis of a new direction in  
70 management of sites contaminated with PCDD/Fs ([Ghosh et al., 2011](#)). Activated carbon  
71 (AC) materials (including granular activated carbon (GAC) and powdered activated carbon  
72 (PAC), has emerged as an effective sorbent amendment for this purpose ([Cornelissen et al.,](#)  
73 [2012; Denyes et al., 2013; Gomez-Eyles et al., 2013; Balasubramani and Rifai, 2018](#)).

74 The retrieval of the amendment with its sorbed contaminants after deployment has  
75 become a priority for a number of reasons. First, complete removal of contaminants, rather  
76 than just immobilization, is preferred by many environmental regulatory agencies ([e.g.,](#)

77 [USEPA, 1997](#)). Second, recovery of the sorbent amendment after its use as a passive sampler  
78 can help determine mass transfer kinetics ([Cornelissen et al., 2008a](#); [Oen et al., 2011](#)).  
79 Adsorbent magnetization is an emerging remediation area where magnetic separation  
80 simplifies isolation and regeneration ([Mohan et al., 2014](#)). Numerous studies have  
81 demonstrated that activated carbon/Fe<sub>3</sub>O<sub>4</sub> composites can be synthesized that maintain high  
82 surface area and high sorption affinities for a growing list of contaminants that includes  
83 organic dyes ([Do et al., 2011](#)), arsenic ([Zhang et al., 2007](#), [Zhang et al., 2010](#)), heavy metals  
84 ([Han et al., 2015](#)), pesticides and PAHs ([Mohan et al., 2014](#)). Our previous work showed  
85 that magnetic Fe<sub>3</sub>O<sub>4</sub> can be easily fabricated from the hydrothermal ferrite process and has  
86 the potential to remove/recover toxic/precious elements from aqueous solutions ([Tu et al.,](#)  
87 [2012](#); [Tu et al., 2013](#); [Tu et al., 2015](#)). To date, adsorbent magnetization has not been applied  
88 to applications involving PCDD/Fs.

89 In our prior work, we provided the first evidence that bioavailability of TCDD sorbed to  
90 two contrasting GACs and one PAC was eliminated in the mammalian (mouse) model. This  
91 conclusion was based upon the use of two highly sensitive bioassays, hepatic induction of  
92 cyp1A1 mRNA, an indirect measure of TCDD exposure, and immunoglobulin M antibody-  
93 forming cell response, a direct measure of immune response ([Boyd et al., 2017](#); [Sallach et al.,](#)  
94 [2019](#)). Prior to this, reductions in bioavailability had only been established based on simpler  
95 model organisms (e.g., worms) or passive samplers ([Fagervold et al., 2010](#); [Chai et al., 2011](#);

96 [Chai et al., 2012](#)). Although the ACs represented a wide range of particle size and pore size  
97 distributions, they were equally effective in eliminating the bioavailability of TCDD, making  
98 them viable candidates for remediation. In this study, we pursued an additional line of  
99 investigation to determine if these same ACs could be functionalized using *in situ* synthesis  
100 of Fe<sub>3</sub>O<sub>4</sub> for subsequent magnetic retrieval ([Indhu et al., 2015](#); [Choi et al., 2016](#)) without  
101 compromising their affinity for dioxins.

102 AC-Fe<sub>3</sub>O<sub>4</sub> composites were synthesized using the same GACs/PAC used in prior  
103 bioavailability studies ([Boyd et al., 2017](#); [Sallach et al., 2019](#)). The specific goals of the  
104 current work were to (1) synthesize GAC/PAC-Fe<sub>3</sub>O<sub>4</sub> composites using ACs shown to be  
105 effective in eliminating TCDD bioavailability in mammals, with emphasis on gaining new  
106 physicochemical insight into the interaction between GAC/PAC and Fe<sub>3</sub>O<sub>4</sub>, (2) characterize  
107 the composites using a combination of X-ray diffraction (XRD), scanning electron  
108 microscopy (SEM), N<sub>2</sub>-BET and micropore analysis, and superconducting quantum  
109 interference device (SQUID); and (3) evaluate sorption characteristics (kinetics and  
110 equilibration) of GAC/PAC, Fe<sub>3</sub>O<sub>4</sub>, and the GAC/PAC-Fe<sub>3</sub>O<sub>4</sub> composite for aqueous phase  
111 dibenzo-p-dioxin (DD). The compound DD served as an isostructural conservative surrogate  
112 for PCDD/Fs, which are important targets for sequestration using environmental geosorbents  
113 due to their extreme recalcitrance in natural environments ([Van Den Berg et al., 1998](#);  
114 [Sallach et al., 2019](#); [Johnston et al., 2012](#)).

## 115 2. Materials and methods

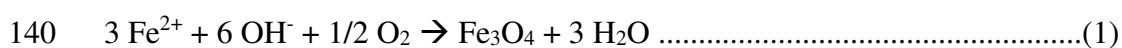
### 116 2.1. Chemicals and materials

117 All solutions were prepared with deionized water.  $\text{Fe}_3\text{O}_4$  was synthesized from ferrous  
118 sulfate  $\text{FeSO}_4$  (> 99.9%, Fisher Scientific, USA) and sodium hydroxide NaOH (99.5%,  
119 Fisher Scientific, USA). Dibenzo-p-dioxin (99% pure) was purchased from ChemService  
120 (West Chester, PA, USA) and was used as received. All the reagents are of analytical grade  
121 and used without further purification. Five GACs (DSRA, G60, FM-1, TOG-LF, and, F400)  
122 and one fine textured PAC (WPC) were purchased/obtained from USEPA, Sigma Aldrich,  
123 Cabot, Calgon Carbon Corp.,. Selected physical properties of the six GACs/PAC are given in  
124 Table 1.

### 125 2.2. Synthesis procedure for magnetic AC- $\text{Fe}_3\text{O}_4$ composites

126 The magnetic GAC/PAC- $\text{Fe}_3\text{O}_4$  composite synthesis followed two synthesis procedures  
127 modified from our previously published method (Tu et al., 2013). Method A was performed  
128 using 0.01 M  $\text{FeSO}_4$  at  $T=338$  K, pH 10, and a reaction time of 2 h. Method B was carried  
129 out using 0.1 M  $\text{FeSO}_4$  at  $T=298$  K, pH 10, and the same 2 h reaction time. Briefly, one gram  
130 of GAC/PAC material was dried under vacuum at room temperature for 20 h using a vacuum  
131 oven (Model 280 A, Fisher Scientific, USA). Dried GAC/PAC was then immersed in 0.01 M  
132 (Method A) or 0.1 M (Method B)  $\text{FeSO}_4$  solution (0.5 L) and mixed continuously at room  
133 temperature for 20 h. After mixing, the pH was adjusted to 10 by dropwise addition of 0.1 M

134 NaOH and then air was bubbled to the solution to initiate the reaction. The synthesis  
135 reactions were carried out at 338 K (Method A) or 298 K (Method B) for 2 h while  
136 maintaining the pH at 10. Finally, the resultant AC-Fe<sub>3</sub>O<sub>4</sub> material was rinsed 5x with DI  
137 water to remove free Fe<sub>3</sub>O<sub>4</sub>, and successfully magnetized GAC/PAC-Fe<sub>3</sub>O<sub>4</sub> composites was  
138 separated from the solution via magnet. The corresponding synthetic reaction of Fe<sub>3</sub>O<sub>4</sub> can be  
139 described as Eq. 1 (Tu et al., 2013).



141 2.3. Characterization of AC-Fe<sub>3</sub>O<sub>4</sub> composite

142 The crystal phases were determined by X-ray diffraction (XRD; X'Pert Pro, Philips,  
143 Netherlands) using graphite monochromatic cobalt radiation over the 2θ range 10-80°. The  
144 surface morphology and particle size were examined by scanning electron microscopy (SEM;  
145 Nova NanoSEM, Oxford instruments, UK). The saturation magnetization of the adsorbent  
146 was measured using a Superconducting Quantum Interference Device (SQUID  
147 magnetometer; MPMS-3, Quantum Design, USA) at 300 ±1 K. N<sub>2</sub> BET and micropore  
148 analysis was conducted using a Micromeritics 3Flex Multiport Chemi/Physi/Micropore  
149 Analyzer.

150 2.4. Measurement of adsorption isotherm

151 Batch adsorption experiments were conducted in duplicate using five initial aqueous DD  
152 concentrations (0.18, 0.3, 0.4, 0.6 and 0.8 mg/L) prepared by a serial dilution of 800 mg/L of

153 DD methanol stock solution. The amount of methanol in the aqueous solutions was 0.1%  
154 which is considered to have minimal cosolvent effects. Aqueous solutions of DD were  
155 sonicated for 60 min at room temperature in a water bath sonicator prior to mixing with the  
156 GAC/PAC (Branson 120, Branson Ultrasonics, Danbury, CT, USA).

157 Two and half (2.5) mg of the adsorbent (GAC, PAC, GAC-Fe<sub>3</sub>O<sub>4</sub>, PAC-Fe<sub>3</sub>O<sub>4</sub>) was  
158 placed in 30 mL Corex glass tubes (Kimble, Vineland, NJ, USA) with  
159 polytetrafluoroethylene (PTFE)-lined screw caps, and mixed with a 30 mL aliquot of DD in  
160 aqueous solutions (methanol 0.1 %). Control samples containing the initial aqueous DD  
161 solutions 0.6 and 0.8 mg/L of DD solution without AC were prepared for calibration to  
162 determine the losses of DD in the batch reactor. Measured values of DD in the control  
163 samples ranged from 0.58 to 0.62 mg/L and 0.78 to 0.82 mg/L for the 0.6 and 0.8 mg/L DD  
164 solutions, respectively. These results indicated that loss of DD to glassware can be ignored.  
165 The suspensions were sonicated for 30 sec prior to shaking at a speed of 60 rpm in a rotary  
166 shaker (Glas-Col, Terre Haute, IN, USA) at room temperature for 10-48 h to achieve the  
167 apparent sorption equilibrium. The supernatant and adsorbent were separated by  
168 centrifugation for GAC/PAC and by external magnetic field for GAC/PAC-Fe<sub>3</sub>O<sub>4</sub>  
169 composites. An aliquot of 1.0 mL of supernatant and DD standards (0.0, 0.18, 0.3, 0.4, 0.6  
170 and 0.8 mg/L) were transferred to HPLC vials. In order to prevent any sorption of DD by  
171 HPLC vials, 0.5 mL of methanol (99.9%) was added to each vial prior to the addition of the

172 supernatant. HPLC vials containing supernatant and methanol were vortexed for 30 sec using  
173 a digital mini-vortexer (VWR, Radnor, PA, USA). Samples were then analyzed for DD  
174 concentrations by direct injection of 50  $\mu$ L into a Thermo Scientific high-performance liquid  
175 chromatography (HPLC) system (Ultimate 3000) equipped with a UV detector and a 150 $\times$   
176 4.60 mm 5 micron Luna 5  $\mu$ m C8(2) 100 Å column (S/N 514816-4). Isocratic elution was  
177 performed using a mobile phase of 80% methanol: 20% water (v/v) with a flow rate of 1.0  
178 mL/min and wavelength of 223 nm for detection.

179 The amount of DD sorbed ( $q_e$ , mg/kg) was calculated as the difference between the  
180 amount initially added and that remaining in the solution after equilibration (Eq. 2):

181 
$$q_e = \frac{(C_o - C_e) \times V}{m_{ads}} \dots \dots \dots (2)$$

182 where  $C_o$  and  $C_e$  are DD concentration in liquid phase at time zero and after equilibration  
183 (mg/L), respectively;  $V$  is the solution volume used in DD adsorption (L);  $m_{ads}$  is GAC/PAC  
184 mass (kg).

185 2.5. Desorption

186 Following collection of the supernatant after equilibrium had been reached, the remaining  
187 supernatant was carefully decanted and the solid phase was re-suspended in a 30 mL solution  
188 of 25% and 99.9% methanol and water (v/v). Tubes were sonicated for 30 sec prior to  
189 shaking 24 h at 60 rpm to ensure equilibrium. Then, the supernatant and adsorbent were  
190 separated by either centrifugation (GAC/PAC) or external magnetic field (GAC/PAC- $Fe_3O_4$

191 composite). Approximately 1.5 mL aliquots of supernatant and standards of 0.18, 0.3, 0.4, 0.6  
192 and 0.8 mg/L were transferred to HPLC vials for HPLC analysis. The amount of DD  
193 desorbed was calculated directly from the concentration of DD present in the supernatant  
194 following equation Eq. 3:

195 Desorption efficiency =  $\frac{C \times V}{X} \times 100\%$ .....(3)

196 where C (mg/L) is the concentration of DD in the desorption solution, V (L) is the volume of  
197 the desorption solution, and X (g) is the amount of DD adsorbed.

198

### 199 **3. Results and discussion**

#### 200 **3.1. Adsorbent characterization in GAC**

201 Six activated carbon materials consisting of five granular activated carbon (GAC)  
202 materials (F400, FM-1, G60, TOG-LF, and DSRA) and one fine textured powdered activated  
203 carbon powder (PAC) (WPC) were functionalized with Fe<sub>3</sub>O<sub>4</sub>. Selected physical properties of  
204 the six materials are given in Table 1. All but one of these GAC/PAC materials (DSRA)  
205 were used in recent TCDD bioavailability studies and were found to be highly effective in  
206 eliminating mammalian bioavailability of TCDD (Boyd et al., 2017; Sallach et al., 2019).

207 The activated carbon materials (Table 1) were functionalized using two different  
208 magnetite synthesis methods. Given their large specific surface areas and micropore (0-2 nm  
209 pores) volumes (Table 1), some magnetite synthesis was expected to occur in the micro- and

210 mesopores (2-50 nm pores) of the GAC/PAC, along with surface decoration of exterior  
211 surfaces of the GAC/PAC particles rendering the GAC/PAC-Fe<sub>3</sub>O<sub>4</sub> composites magnetic.  
212 Fe<sub>3</sub>O<sub>4</sub> synthesis Method A was performed at elevated temperature (338 K) using 0.01 M  
213 FeSO<sub>4</sub>. The resulting magnetization of the PAC-Fe<sub>3</sub>O<sub>4</sub> composite was successful with a value  
214 of 9.7 emu/g (Table 1). In contract, observed magnetization values measured at 300 K for the  
215 five GACs were weak with values of 0.61 (DSRA), 0.54 (FM-1), 0.49 (TOG-LF), 0.46 (G60)  
216 and 0.35 (F-400) emu/g (Table 1). For comparison, the magnetization of Fe<sub>3</sub>O<sub>4</sub> synthesized  
217 using Synthesis Method A without GAC/PAC was 71.9 emu/g. Weak magnetization values  
218 in the ranged of 0.35-0.61 emu/g were not sufficient to allow rapid magnetic separation.

219 Because the magnetization values resulting from synthesis Method A for the granular  
220 activated carbon samples were weak (0.35-0.61 emu/g), the Fe<sub>3</sub>O<sub>4</sub> synthesis procedure was  
221 modified using synthesis Method B, which utilized a higher concentration of FeSO<sub>4</sub> (0.1 M)  
222 and a lower temperature of 298 K. The granular activated carbon sample DSRA was selected  
223 because it showed the highest level of magnetization among the five GAC materials  
224 evaluated using Method A (Table 1). Synthesis Method B resulted in a DSRA-Fe<sub>3</sub>O<sub>4</sub>  
225 composite with a significantly improved magnetization value of 5.38 emu/g (Table 1). No  
226 residual magnetism was detected in either of the GAC(DSRA)-Fe<sub>3</sub>O<sub>4</sub> (Synthesis Method B)  
227 or PAC(WPC)/Fe<sub>3</sub>O<sub>4</sub> (Synthesis Method A) composites indicating that these two materials  
228 are superparamagnetic (Table 1). For simplicity, the GAC(DSRA)-Fe<sub>3</sub>O<sub>4</sub> complex (using

229 Synthesis Method B) will be referred to as GAC-Fe<sub>3</sub>O<sub>4</sub>(B) and the PAC(WPC)-Fe<sub>3</sub>O<sub>4</sub>  
230 complex (using Synthesis Method A) will be referred to as PAC-Fe<sub>3</sub>O<sub>4</sub>(A), where (A) and  
231 (B) refer to Synthesis Methods A and B, respectively.

232 The GAC-Fe<sub>3</sub>O<sub>4</sub>(B) composite in aqueous suspension was efficiently recovered by  
233 applying an external magnetic field. The complete (~100%) separation of the GAC-Fe<sub>3</sub>O<sub>4</sub>(B)  
234 composite from solution using a magnet was achieved within only 20 seconds (Supplemental  
235 Information Fig. S1). When the external magnetic field was removed, the GAC-Fe<sub>3</sub>O<sub>4</sub>(B)  
236 composite could be readily dispersed again in water by physical shaking.

237 The XRD patterns of the GAC-Fe<sub>3</sub>O<sub>4</sub>(B) and PAC-Fe<sub>3</sub>O<sub>4</sub>(A) composites are shown in  
238 Fig. 2. The observed diffraction peaks at d-spacings of 4.846, 2.968, 2.531, 2.423, 2.099,  
239 1.713, 1.615, and 1.484 Å matched the XRD reflections of Fe<sub>3</sub>O<sub>4</sub> (JCPDS file number 04-  
240 007-9093). No other peaks were detected in the XRD pattern of the GAC-Fe<sub>3</sub>O<sub>4</sub>(B)  
241 confirming that the only crystalline phase present is Fe<sub>3</sub>O<sub>4</sub> nanoparticles in the GAC-  
242 Fe<sub>3</sub>O<sub>4</sub>(B) composite. In addition to the Fe<sub>3</sub>O<sub>4</sub> peaks, the PAC-Fe<sub>3</sub>O<sub>4</sub>(A) composite had small  
243 peaks at 24.2, 31.0 and 58.5 °2θ.

244 Further characterization of the GAC and the GAC-Fe<sub>3</sub>O<sub>4</sub>(B) composite was provided by  
245 SEM imaging of the two materials at different levels of magnification (Figs. 3a-3f). The  
246 average bulk size of the GAC was ~1 mm with <5% passing through a 40 US Mesh sieve  
247 (420 μm) (Fig. 3a); 'large' pores were observed ranging in size from several μm to >10 μm

248 (Figs. 3b and 3c). The synthesized  $\text{Fe}_3\text{O}_4$  particles were observed to be spherical, and their  
249 primary particle size ranged between 20 and 120 nm (Fig. 3d). From the SEM images it is  
250 evident that the synthesized  $\text{Fe}_3\text{O}_4$  nanoparticles were randomly distributed on the surfaces  
251 and pores of the GAC particles (Fig. 3f).

### 252 3.2 $\text{N}_2$ BET and Textural Analysis

253  $\text{N}_2$  BET and micro-textural analysis of the activated carbon materials prior to magnetite  
254 synthesis are presented in Table 1. The supplier of the activated carbon, feedstock, along with  
255  $\text{N}_2$  BET surface area and micropore analysis are presented in Table 1. The five granular  
256 activated carbon samples ranged in percentage micropore (0-2 nm) volume from 29 to 82%  
257 of the total micropore and mesopore (2-50 nm) volume. The fine textured WPC powder had  
258 very little mesoporosity with 91% of its pore volume in the micropore range. Of the six  
259 samples, five were used in prior toxicology studies to assess TCDD bioavailability (Boyd et  
260 al., 2017; Sallach et al., 2019). Although these ACs represented a wide range of particle size  
261 and pore size distributions, they were equally effective in eliminating the bioavailability of  
262 TCDD, making them viable candidates for remediation. After  $\text{Fe}_3\text{O}_4$  synthesis, the specific  
263 surface area of the GAC(DSRA)- $\text{Fe}_3\text{O}_4$  composite decreased from 822 to 633  $\text{m}^2/\text{g}$  along  
264 with a modest reduction in micropore volume (0.388 to 0.262  $\text{g}/\text{cc}$ ) indicative of some partial  
265 pore blocking by the magnetite particles (Table 1).

### 266 3.3. Batch equilibrium sorption and kinetics

267 Batch sorption isotherms of dibenzo-p-dioxin (DD) to GAC and GAC-Fe<sub>3</sub>O<sub>4</sub>(B) are  
268 shown in Fig. 4. Both GAC and GAC-Fe<sub>3</sub>O<sub>4</sub> composite showed a high affinity for aqueous  
269 phase DD at low equilibrium concentrations (<0.005 mg/L) up to a sorbed concentration  
270 about of about 4000 mg/kg. At higher equilibrium concentrations (0.005-0.12 mg/L), sorption  
271 isotherms showed some nonlinear behavior exhibiting high affinity at low equilibrium  
272 concentration followed by an “S-shaped” response (Giles and Smith, 1974). The GAC-  
273 Fe<sub>3</sub>O<sub>4</sub>(B) isotherm is shifted to higher equilibrium concentrations (i.e., lower affinity)  
274 compared the GAC, however, both materials sorbed > 8000 mg/kg. These results could be  
275 explained by sorption processes involving easily accessible external sites and less accessible  
276 pores. Since Fe<sub>3</sub>O<sub>4</sub> demonstrated no sorption affinity for DD, its presence within the GAC  
277 composite likely blocked some sorption sites or access to certain pores manifesting a slight  
278 decrease in DD affinity.

279 The kinetics of DD sorption by GAC and GAC-Fe<sub>3</sub>O<sub>4</sub> composite were evaluated over a  
280 period of 40 h using the batch equilibration method (described above) with initial aqueous  
281 phase DD concentrations of 0.18, 0.4, and 0.8 mg/L (Fig. 5). Sorption kinetics of DD can be  
282 separated into two phases. Initially, within the first 10 h, both GAC and GAC-Fe<sub>3</sub>O<sub>4</sub>  
283 demonstrated comparatively rapid uptake of DD for all three initial concentrations. This was  
284 followed by a slower phase (>50 h) to reach apparent equilibrium.

285 For the lower and intermediate initial DD concentrations of 0.18 and 0.4 mg/L, DD  
286 uptake from aqueous solution by GAC was rapid and nearly complete within the first 10 h;  
287 the percent DD removal approached 100 percent (Fig 5c-f). However, at the higher initial DD  
288 concentration of 0.8 mg/L, there is a more gradual approach to apparent equilibrium over  
289 time (Fig. 5a-b). For the initial concentrations of 0.18 and 0.4 mg/L, DD adsorption by GAC  
290 was essentially complete by 10 h, and the total uptake of DD from aqueous solution  
291 approached 100 percent. For the initial concentration of 0.8 mg/L the data indicate that  
292 equilibrium had not been achieved after 50 h for GAC-Fe<sub>3</sub>O<sub>4</sub>.

293 For this system, it is assumed that sorption kinetics are controlled surface adsorption with  
294 associated resistance to film diffusion followed by an emerging contribution to DD sorption  
295 via intraparticle diffusion. Pseudo-second order kinetic models are commonly used to  
296 describe sorption kinetics for these types of interactions (Ho and, McKay, 1999;  
297 Amarasinghe and Williams, 2007). The pseudo-second order kinetic model was able to fit  
298 the experimental data well (see Table 2). The pseudo-second order model results are plotted  
299 on the kinetics data shown in Fig. 5.

300 The correlation coefficients ( $R^2$ ) and kinetic parameters derived from the pseudo-second  
301 order models are summarized in Table 2. These results suggest that the rate-limiting step may  
302 be some type of site-specific mechanism involving direct interaction between the sorbent and  
303 sorbate (Amarasinghe and Williams, 2007). The kinetic rate constant ( $k_2$ ) from the pseudo-

304 second order model decreased with increasing initial DD concentrations (Table 1), indicating  
305 that the DD adsorption rates are faster at lower concentrations. In other words, the time  
306 required to reach equilibrium increased as the initial DD concentration increased. This is  
307 likely due to competition for active surface sites and pores of the sorbent which is greater at a  
308 higher DD concentration.

### 309 3.4 Desorption of DD

310 Desorption of DD into solutions of either 25% or 99.9% methanol and water (v/v)  
311 desorption solutions were used to evaluate the reversibility of DD sequestration by GAC and  
312 the GAC-Fe<sub>3</sub>O<sub>4</sub> composite. Desorption of DD from GAC and GAC-Fe<sub>3</sub>O<sub>4</sub> in 99.9% methanol  
313 was 19% and 14.3% (after 20 h), respectively, and 11.2% and 12.4% for 25%  
314 methanol/water, respectively. These results agree with our prior study that showed 22-27%  
315 of TCDD bound to three of the activated carbons samples used in the present study could be  
316 desorbed after 64 hours of Soxhlet extraction using toluene (Sallach et al., 2019).  
317 Magnetizing GAC with Fe<sub>3</sub>O<sub>4</sub> had little to no effect on the propensity of DD to desorb. That  
318 the fraction of released DD was less than 20% even for 99.9% methanol clearly indicated the  
319 strong affinity between DD and GAC; once sorbed DD appears to be largely irreversibly-  
320 sequestered within the pore structure of GAC. The resistance to desorption, even into  
321 methanol, is consistent with our prior observation that sequestration of 2,3,7,8-TCDD by AC  
322 eliminated its mammalian bioavailability (Boyd et al., 2017; Stedtfeld et al., 2017).

### 323 3.5 Sorption removal comparison of fine-textured AC-Fe<sub>3</sub>O<sub>4</sub> to GAC-Fe<sub>3</sub>O<sub>4</sub>

324 The uptake of DD by GAC, GAC-DSR-Fe<sub>3</sub>O<sub>4</sub>(B), PAC and PAC-Fe<sub>3</sub>O<sub>4</sub>(A) as a function  
325 of time are shown in Fig. 6. As expected, uptake of DD by the fine-textured AC (WPC) was  
326 rapid and nearly stoichiometric. More than 90% of the WPC has a particle size of < 45 μm  
327 (Sallach et al., 2019). In contrast, sorption kinetics for DD uptake by the coarse-textured  
328 GAC and GAC-Fe<sub>3</sub>O<sub>4</sub>(B) were considerably slower. Although the surface area of the PAC,  
329 GAC and their Fe<sub>3</sub>O<sub>4</sub> derivatives are comparable, most of the surface area in GAC can only  
330 be accessed through intraparticle diffusion, resulting in slower sorption kinetics (Figs. 5-6).

## 331 4. Discussion

332 The activated carbon materials, including GAC and PAC, used here to form the magnetic  
333 variants were also used in our prior bioavailability studies (Boyd et al., 2017; Stedtfeld et al.,  
334 2017; Sallach et al., 2019), along with natural geosorbents including clay minerals and silica  
335 (Boyd et al., 2011; Kaplan et al., 2011; Chai et al., 2016). Among these, only GAC and PAC  
336 eliminated the bioavailability of sorbed TCDD to a mammalian (mouse) model.  
337 Mammalian bioavailability was evaluated in our prior work using two highly sensitive  
338 bioassays, hepatic induction of cyp1A1 mRNA, an indirect measure of TCDD exposure, and  
339 immunoglobulin M antibody-forming cell response, a direct measure of immune response  
340 (Boyd et al., 2017; Sallach et al., 2019). In contrast to the complete elimination of TCDD  
341 bioavailability achieved via sequestration by GAC/PAC, TCDD bound to clay minerals and

342 silica was found to be 100% bioavailable. The lack of mammalian bioavailability of TCDD  
343 bound to GAC/PACs was consistent with a related study showing contaminant bioavailability  
344 to lower organisms was significantly decreased in the presence of AC (Chai et al., 2012). In  
345 addition, attempts to extract sorbed TCDD from ACs using Soxhlet extraction revealed that  
346 only a minor fraction of the total TCDD present could be recovered (Sallach et al., 2019).  
347 From an applied perspective, these laboratory results are now leading to the use of GAC/PAC  
348 sorbent amendments in large-scale remediation efforts for impacted soils, sediments and  
349 water bodies (Samuelsson et al., 2017; Payne et al., 2019; Cornelissen et al., 2016;  
350 Beckingham et al., 2011). Given our earlier results showing elimination of mammalian  
351 TCDD bioavailability, creating magnetic GAC/PAC composites that could be used as a  
352 retrievable form of GAC/PAC sorbent amendments was attempted. The ability to retrieve  
353 magnetized GAC/PAC would enhance their utility as passive samplers in field settings and in  
354 ongoing laboratory studies. For example, our earlier mammalian studies would have  
355 benefited from using a magnetic AC to determine the amount TCDD in fecal pellets from  
356 mice that were dosed with TCDD–AC (Boyd et al., 2017; Stedfeldt et al., 2017; Sallach et al.,  
357 2018). Likewise, the ability to ultimately retrieve (magnetized) GAC/PAC sorbent  
358 amendments used to remediate areas with especially high levels of contamination would  
359 provide both an immediate benefit, i.e., bioavailability reduction, and make contaminant  
360 removal possible in the longer term.

361 The synthesis procedure using a lower concentration of  $\text{FeSO}_4$  (0.01 M) at an elevated  
362 temperature (338 K), Synthesis Method A, was successful in synthesizing a magnetic PAC-  
363  $\text{Fe}_3\text{O}_4$  composite (PAC- $\text{Fe}_3\text{O}_4$ (A)). In the case of the PAC, magnetization most likely  
364 occurs on the external surfaces of fine-textured AC and this could be accomplished using the  
365 lower concentration of  $\text{FeSO}_4$ . However, Method A only produced weak magnetization  
366 values for the GAC- $\text{Fe}_3\text{O}_4$  composites. One could argue that this procedure was not able to  
367 synthesize  $\text{Fe}_3\text{O}_4$  particles within the coarse textured GACs. The procedure was modified  
368 using a higher concentration of  $\text{FeSO}_4$  (0.1 M) and lower temperature that resulted in a  
369 sufficiently magnetic GAC- $\text{Fe}_3\text{O}_4$ (B) composite.

370 Both the GAC- $\text{Fe}_3\text{O}_4$ (B) and PAC- $\text{Fe}_3\text{O}_4$ (A) composites revealed the presence of  $\text{Fe}_3\text{O}_4$   
371 based on X-ray diffraction analysis (Fig. 2) and it is possible that careful XRD studies could  
372 be used as a surrogate for the more difficult to obtain SQUID magnetization results.  $\text{N}_2$  BET  
373 surface area and textural analysis showed that both the GAC and PAC materials were  
374 characterized by high  $\text{N}_2$ -surface area (802-822  $\text{m}^2/\text{g}$ ). The specific surface area of the  
375 GAC- $\text{Fe}_3\text{O}_4$ (B) composite showed a modest reduction in both surface area (822 to 633  $\text{m}^2/\text{g}$ )  
376 and micropore volume (0.38 g/cc to 0.26 g/cc) compared to the starting GAC (Table 1).  
377 The batch sorption isotherms of dibenzo-p-dioxin (DD) on the GAC and GAC- $\text{Fe}_3\text{O}_4$ (B)  
378 composites showed that sorption affinity of DD was slightly reduced due to the presence of  
379 magnetite particles (GAC- $\text{Fe}_3\text{O}_4$ (B)) compared GAC. The decrease in surface area and DD

380 sorption is interpreted as some blocking of pore throats. Overall, the sorption isotherms for  
381 both GAC and GAC-Fe<sub>3</sub>O<sub>4</sub>(B) showed some sorption nonlinearity, consistent with a range of  
382 sorption sites of varying accessibility.

383 The sorption kinetics of DD uptake by GAC, GAC-Fe<sub>3</sub>O<sub>4</sub>(B), PAC and PAC-Fe<sub>3</sub>O<sub>4</sub>(A)  
384 were strongly dependent on particle size. Uptake of DD by the fine textured PAC and PAC-  
385 Fe<sub>3</sub>O<sub>4</sub>(A) composite was rapid and complete within 10 hours. In contrast, sorption uptake  
386 was much slower for GAC and the GAC-Fe<sub>3</sub>O<sub>4</sub>(B) composite (Fig. 4 and 5) owing to the  
387 larger particle size. Sorption equilibria had not been reached for the GAC-Fe<sub>3</sub>O<sub>4</sub>(B)  
388 composite after 40 h. These results are consistent with prior work showing the influence of  
389 particle size on sorption kinetics of hydrophobic organic solutes on activated carbon and  
390 biochars (Ahn et al., 2005; Kang et al., 2018). As shown, the rate of DD uptake by the fine  
391 textured AC and AC-Fe<sub>3</sub>O<sub>4</sub>(A) is rapid with showing 97% removal of DD from aqueous  
392 solution after one hour. The larger sized GAC, with particle diameters ~ 1 mm (5% < 420  
393 μm), showed much slower uptake (Fig. 6a) and is attributed to the longer diffusion pathways  
394 to binding sites and pore structures in GAC.

395 We demonstrated that both a GAC and a PAC could be magnetized and, more  
396 importantly, the GAC-Fe<sub>3</sub>O<sub>4</sub>(B) and PAC-Fe<sub>3</sub>O<sub>4</sub>(A) composites maintained high sorption  
397 affinity for dioxin. Particle size was a dominant factor in controlling sorption kinetics, with  
398 the fine-textured PAC showing nearly complete uptake of dioxin within 1 hour compared to

399 considerably slower uptake by the coarse texture GAC. These differences could be significant  
400 in animal dosing studies but less significant for materials deployed as passive samplers over  
401 periods of months to years. Finally, these results could prove useful in the design of large-  
402 scale recoverable geosorbents manufactured for contaminant removal.

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410

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614 Table titles:

615 Table 1. Selected physical properties of GAC, GAC-Fe<sub>3</sub>O<sub>4</sub> and PAC.

616 Table 2. Kinetic parameters at different concentration for adsorption of DD by using DSRA  
617 and GAC-Fe<sub>3</sub>O<sub>4</sub> composite.

618

619 Figure Captions

620 Fig. 1. Saturation magnetization of GACs and PAC-Fe<sub>3</sub>O<sub>4</sub> composites measured by SQUID.

621 Fig. 2. X-ray powder diffraction (XRD) patterns of the GAC-Fe<sub>3</sub>O<sub>4</sub>(B) and PAC-Fe<sub>3</sub>O<sub>4</sub>(A)  
622 composites along with reference reflections for magnetite Fe<sub>3</sub>O<sub>4</sub>.

623 Fig. 3. Scanning electron microscopy (SEM) images of (a) DSRA (120X magnitude); (b)

624 DSRA (500X magnitude); (c) DSRA (800X magnitude); (d) GAC-Fe<sub>3</sub>O<sub>4</sub> composite

625 (50000X magnitude); (e) GAC-Fe<sub>3</sub>O<sub>4</sub> composite (150000X magnitude); and (f) GAC-

626 Fe<sub>3</sub>O<sub>4</sub> composite (350000X magnitude).

627 Fig. 4. Batch equilibrium sorption isotherms of dibenzo-p-dioxin (DD) on GAC (black  
628 squares) and GAC-Fe<sub>3</sub>O<sub>4</sub>(B) (red circles). Conditions: T=298 K, solution volume=30  
629 mL, adsorbent =2.5 mg, contact time=10-48 h.

630 Fig. 5. Sorption kinetics of dibenzo-p-dioxin (DD) uptake by GAC and GAC-Fe<sub>3</sub>O<sub>4</sub>(B) over  
631 40 h of contact. Top figure shows results from initial concentration of 0.8 mg/L,  
632 middle figure shows results from initial concentration of 0.4 mg/L, and lower figure  
633 shows initial concentration of 0.18 mg/L. GAC is represented by black squares and  
634 GAC-Fe<sub>3</sub>O<sub>4</sub>(B) is represented by red circles. Conditions: T=298 K, solution  
635 volume=30 mL, adsorbent =2.5 mg.

636 Fig. 6. Sorption kinetics of dibenzo-p-dioxin (DD) uptake by GAC, GAC-Fe<sub>3</sub>O<sub>4</sub>(B), PAC and  
637 PAC-Fe<sub>3</sub>O<sub>4</sub>(A) over 18 h of contact. PAC is represented by black squares, PAC-  
638 Fe<sub>3</sub>O<sub>4</sub>(A) is represented by solid red circles, GAC is represented by open black  
639 squares, and GAC-Fe<sub>3</sub>O<sub>4</sub>(B) is represented by open red circles. Conditions: T=298 K,  
640 solution volume=30 mL, adsorbent =2.5 mg. Conditions: DD concentration=0.8  
641 mg/L, T=298 K, solution volume=30 mL, adsorbent=2.5 mg, contact time=1 h.

642

1 Figure Captions

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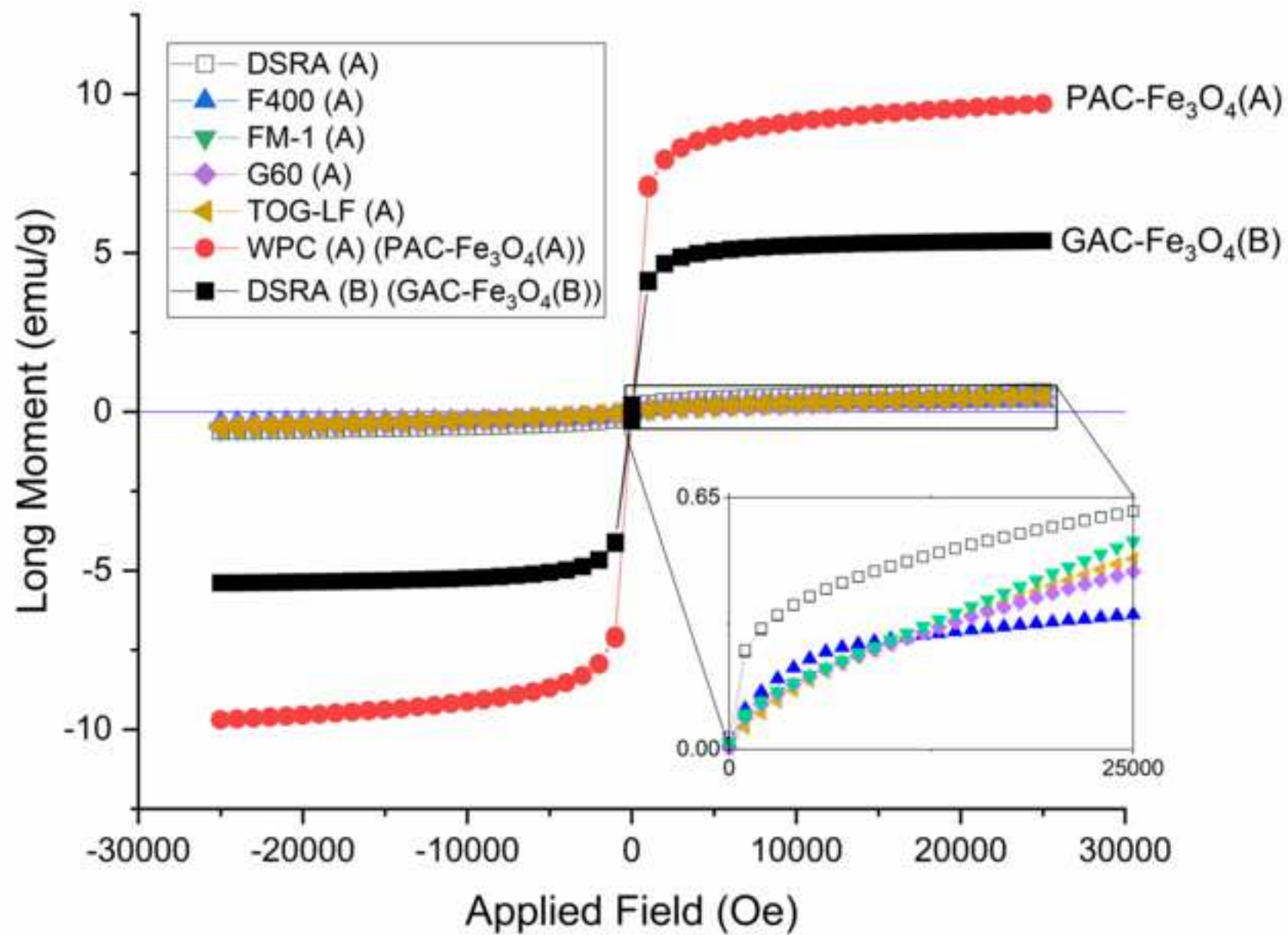
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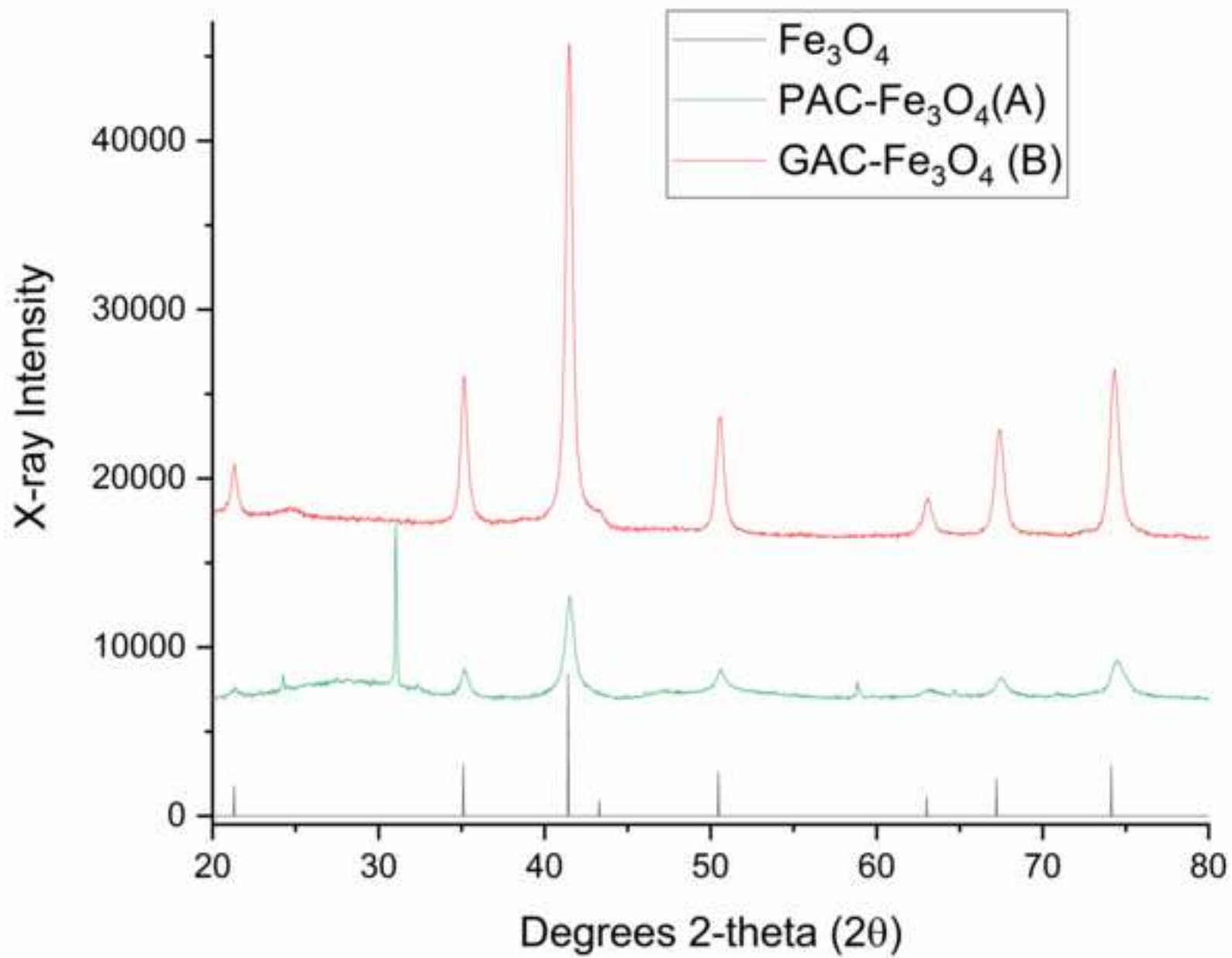
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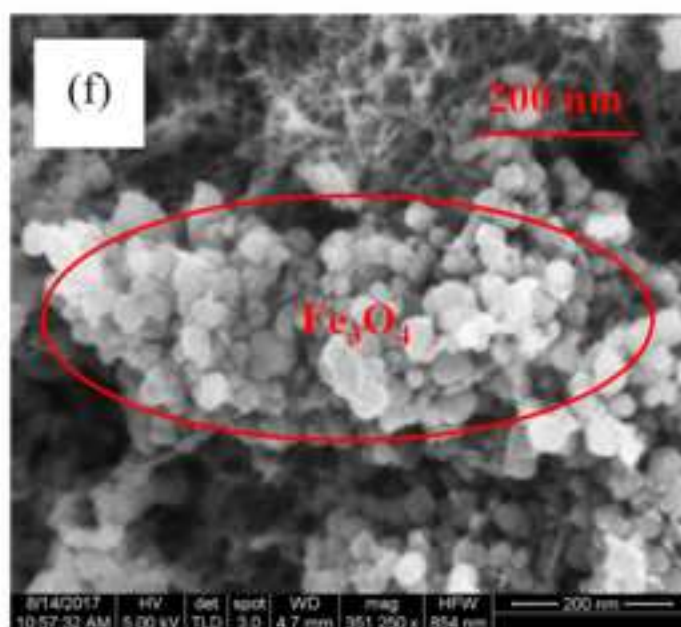
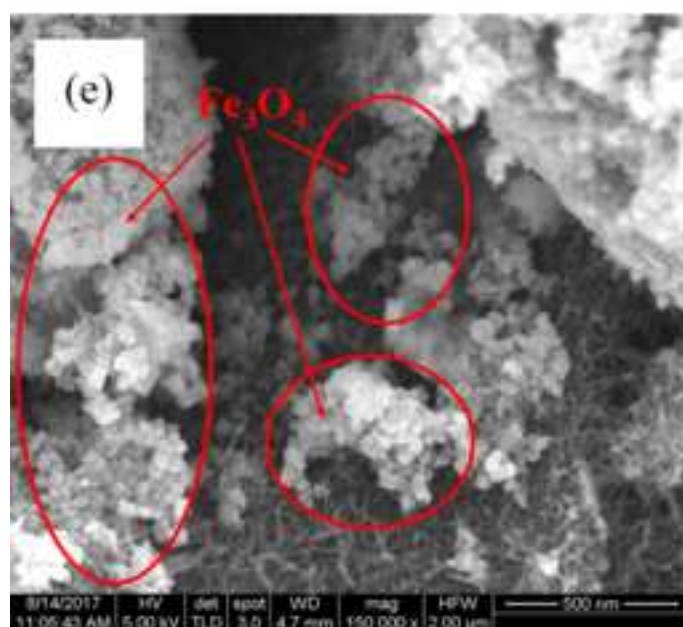
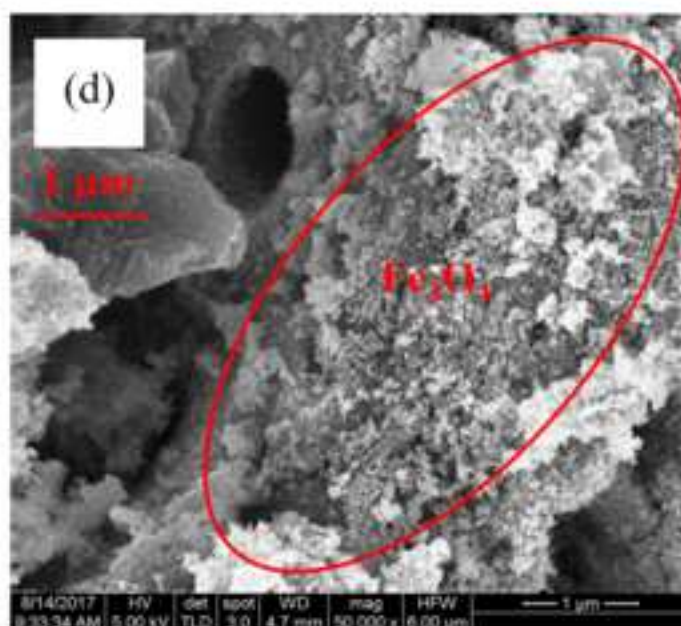
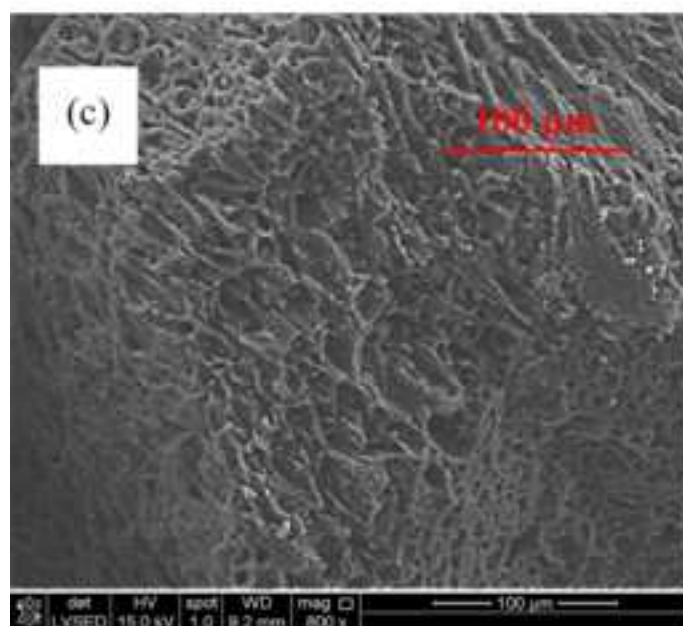
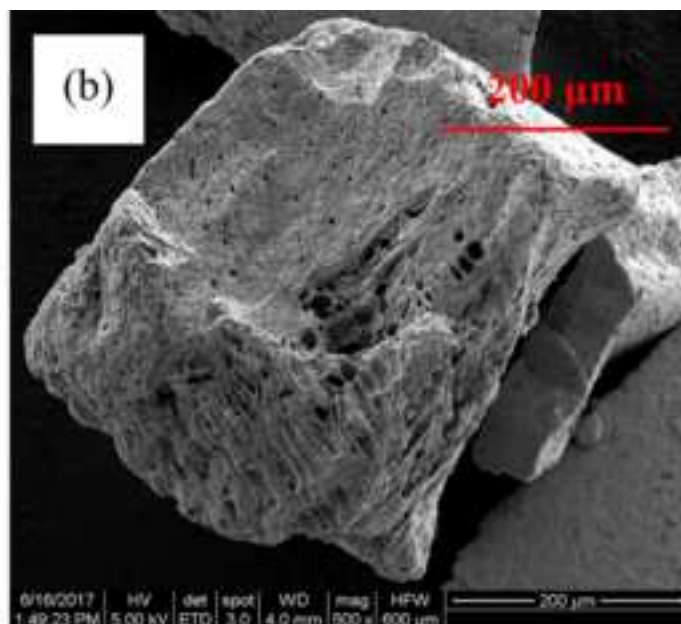
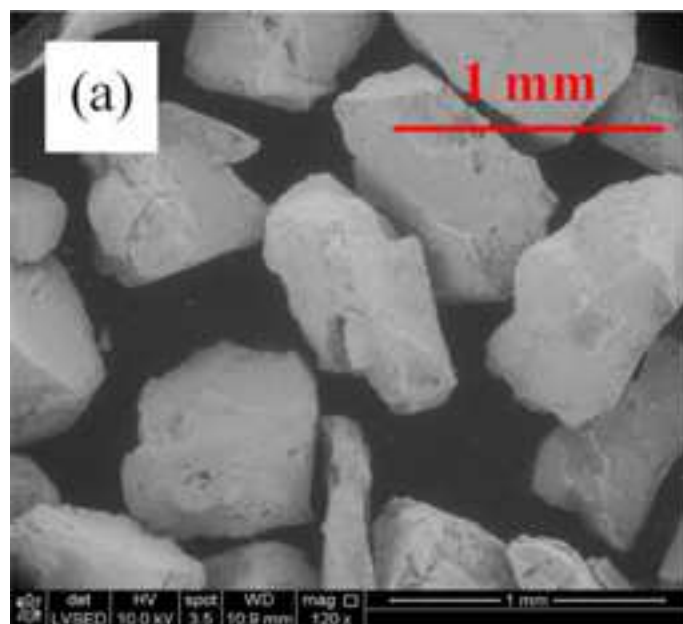
20  $\text{Fe}_3\text{O}_4(\text{A})$  is represented by solid red circles, GAC is represented by open black  
21 squares, and GAC- $\text{Fe}_3\text{O}_4(\text{B})$  is represented by open red circles. Conditions:  $T=298\text{ K}$ ,  
22 solution volume=30 mL, adsorbent =2.5 mg. Conditions: DD concentration=0.8  
23 mg/L,  $T=298\text{ K}$ , solution volume=30 mL, adsorbent=2.5 mg, contact time=1 h.

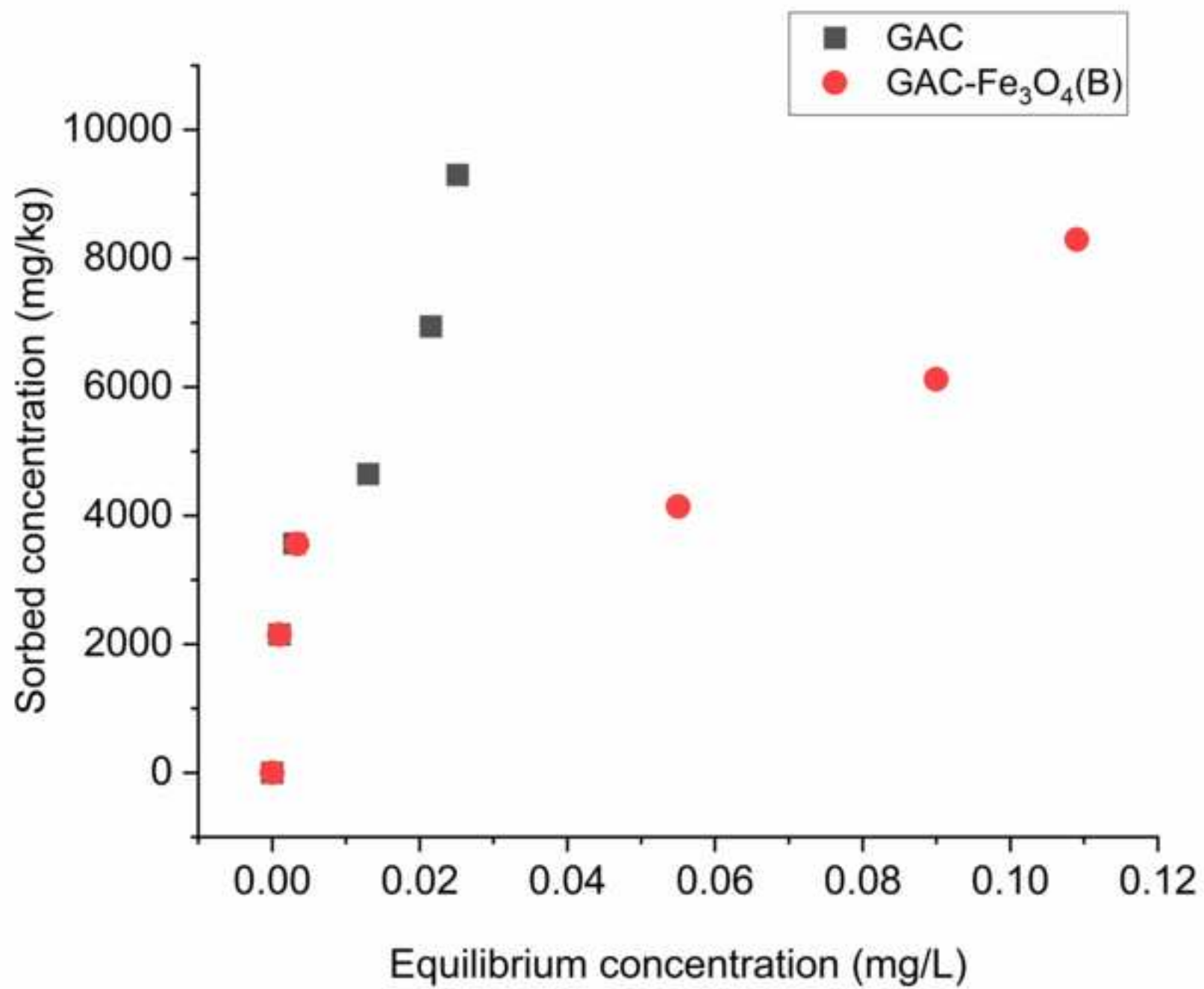
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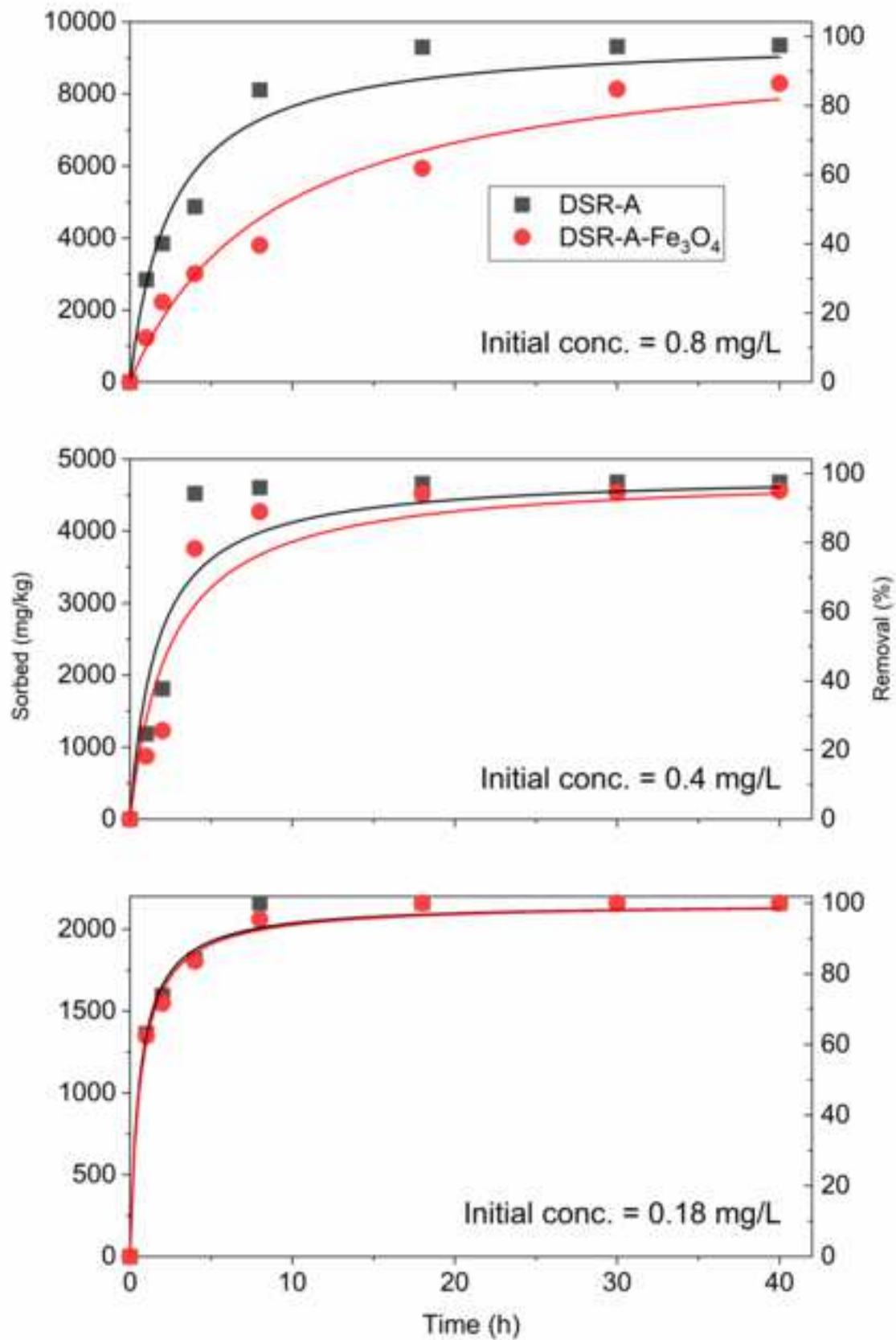
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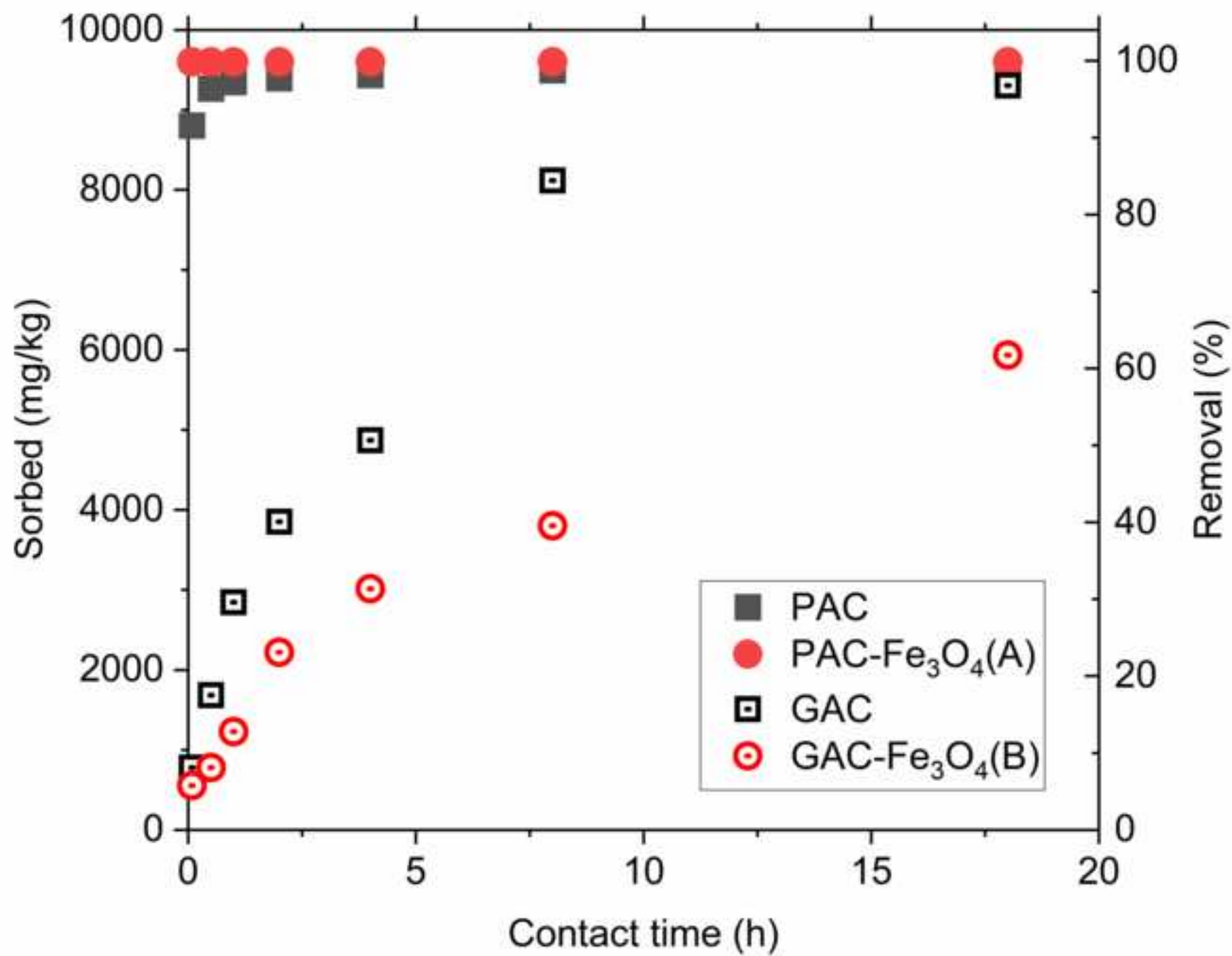


Table 1. Selected physical properties of the activated carbon materials

Material	Supplier	Source	Surface Area (m <sup>2</sup> /g)	Mesopore vol. g/cc (% of total)	Micropore vol. g/cc (% of total)	<sup>a</sup> Magnetization after Syn. Method A (emu/g)	<sup>b</sup> Magnetization after Syn. Method B (emu/g)
G60	Cabot	Lignite	987	0.38 (57%)	0.29 (43%)	0.46	
FM-1	Cabot	Lignite	520	0.36 (71%)	0.15 (29%)	0.54	
TOG-LF	Calgon	Coal	916	0.15 (33%)	0.30 (67%)	0.49	
F-400	Calgon	Coal	1044	0.16 (31%)	0.36 (69%)	0.35	
WPC (AC)	Calgon	Coconut	802	0.03 (9%)	0.29 (91%)	9.7	
DSRA (GAC)	Calgon	Pool Rej.	822	0.11 (22%)	0.38 (78%)	0.61	
DSRA-Fe <sub>3</sub> O <sub>4</sub>	Calgon	Pool Rej.	633	0.06 (18%)	0.26 (82%)		5.38

Note:

<sup>a</sup>: Magnetization after Fe<sub>3</sub>O<sub>4</sub> impregnation at conditions of 338 K and 0.01 M FeSO<sub>4</sub>.

<sup>b</sup>: Magnetization after Fe<sub>3</sub>O<sub>4</sub> impregnation at conditions of 298 K and 0.1 M FeSO<sub>4</sub>.

Table 2. The pseudo-second order kinetic model for GAC (DSRA) and GAC-Fe<sub>3</sub>O<sub>4</sub>(B)

	GAC (DSRA)			GAC-Fe <sub>3</sub> O <sub>4</sub> (B)		
	q <sub>e</sub>	k <sub>2</sub>	Adjusted R Square	q <sub>e</sub>	k <sub>2</sub>	Adjusted R Square
0.18 mg/L	2160	7.62E-04	0.990	2160	7.02E-04	0.994
0.4 mg/L	4800	1.26E-04	0.893	4800	8.47E-05	0.916
0.8 mg/L	9600	4.10E-05	0.967	9600	1.17E-05	0.973