

1	Buffered cyclodextrin extraction of ¹⁴ C-phenanthrene from
2	black carbon amended soil
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25 Abstract

26 The presence of black carbon (BC) in soil drastically reduced the mineralization 27 of ¹⁴C-phenanthrene and its extractability by hydroxylpropyl-β-cyclodextrin 28 (HPCD) extractions. This study also tested the effects of pH on the HPCD 29 extraction of ¹⁴C-phenanthrene in soils with BC. Extractions using 60 mM HPCD 30 solutions prepared in deionized water (pH 5.89) and phosphate buffers (pH 7 31 and 8) were conducted on ¹⁴C-phenanthrene-spiked soils amended with three 32 different types of BC (1% dry weight) after 1, 25, and 50 d of ageing. 33 Biodegradation assays using a *Pseudomonas* sp. strain were also carried out. 34 Results showed that after 1 and 25 d, HPCD at pH 7 extracted significantly more 35 ¹⁴C-phenanthrene (p < 0.05) from BC-amended soils than the other two solutions 36 (un-buffered and pH 8), while HPCD at pH 8 extracted statistically similar (p >37 0.05) amounts of phenanthrene compared to the un-buffered solution. At 50 d, 38 HPCD at pH 8 generally extracted more ¹⁴C-phenanthrene from all treatments. It 39 was proposed that higher pH promoted the dissolution of soil organic matter 40 (SOM), leading to a greater solubility of phenanthrene in the solvent phase and 41 enhancing the extractive capability of HPCD solutions. Although correlations 42 between extractability and biodegradability of ¹⁴C-phenanthrene in BC-amended 43 soils were poor, increasing pH was demonstrated a viable approach to enhancing 44 HPCD extractive capability from the ¹⁴C-PAH from soil.

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Keywords — black carbon, phenanthrene, hydroxylpropyl- β -cyclodextrin extraction 47 (HPCD), mineralization, pH

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50 1. Introduction

51	Massive consumption on fossil fuels and combustion of biomass in modern
52	world has dramatically increased the input of black carbon (BC) into the
53	environment [1]. BC is a group of heterogeneous carbon possessing strong
54	sorptive capabilities and recalcitrance to chemical and biological transformation
55	[4]. It is mainly produced by incomplete combustion of fossil fuels or biomass [1-
56	4]. BC is ubiquitously distributed across the environmental compartments
57	including soil, where it impacts the fate and behaviour of a range of
58	contaminants such as hydrophobic organic contaminants (HOCs) [5, 6].
59	Moreover, commercially produced BC (e.g. activated carbon, AC) has also been
60	proposed and piloted as a tool for contaminated land remediation [7].
61	Nevertheless, there is still a lack of understanding regarding the implications of
62	BC on the bioaccessibility of soil organic contaminants and risk assessment of
63	contaminated land [4].
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75 microorganisms, which HPCD extraction was not able to account for. Although 76 the mechanism involved in direct microbial uptake of sorbed substances has 77 been reported by Alexander [13], this explanation is still questionable 78 considering that the uptake of organic substances by soil microorganisms 79 predominantly takes place in the aqueous phase [14, 15]. It is also possible that 80 other microbial processes (e.g. biosurfactant production) could promote the 81 desorption of the BC-associated target chemical, while water as the solvent of 82 HPCD solution used in these researches was not capable of displacing target 83 compounds from sorption sites on BC particles [16]. 84 As it has been previously suggested [4], it is important to find a reliable chemical 85 method to estimate the bioaccessibility of HOCs in soils with BC given the 86 growing input of BC to soil from anthropogenic sources and the application of 87 commercially produced BC as a strategy for the remediation of contaminated 88 systems. For this purpose, a potential approach is to modify HPCD extraction 89 methodology by integrating a buffer of higher pH into the solvent to achieve a 90 greater displacement capacity for target compounds, as increasing pH promotes 91 the dissolution of SOM [17] which contributes to greater aqueous solubility of 92 organic pollutants [18]. This was also demonstrated by Reid et al. [10] who 93 observed enhanced extractive capability of HPCD solution prepared in 94 phosphate buffer of pH 8 for phenanthrene [10]. Therefore, this study aims to 95 investigate the effects of phosphate buffers of higher pH values on the extractive 96 capability of HPCD solutions for phenanthrene (a) in soils amended with 97 different types of commercially produced BC, (b) after different periods of soil-98 contaminant interactions. Parallel biodegradation assays with a phenanthrene-

- 99 degrading inoculum (*Pseudomonas* sp.) to measure the microbially accessible
- 100 fraction of the PAHs in the soil.

101 **2. Materials and methods**

102 2.1 Chemicals

- 103 Unlabelled phenanthrene was obtained from Sigma Aldrich Co, Ltd. UK. [9-14C]
- 104 Phenanthrene was purchased from American Radiolabelled Chemicals, Inc., USA.
- 105 Liquid scintillation cocktail (Goldstar) and sample oxidation cocktails (Carbotrap

106 and Carbocount) were obtained from Meridian Biotechnologies Ltd, UK.

107 Hydroxylpropyl-β-cyclodextrin (HPCD) was purchased from Acros Organics,

108 Belgium. General purpose grade agar (GPA) was obtained from Fisher Scientific,

- 109 UK. Activated carbon (Colorsorb P3-1, Aquasorb CP2 and Aquasorb BP2) was
- 110 obtained from Jacobi Carbons, UK.
- 111 2.2 Soil collection and characterization
- 112 Pristine soil was collected (A horizon; 5 20 cm) from Myerscough Agricultural
- 113 College in Lancashire, UK, and passed through a 2 mm sieve to remove stones
- and roots. General soil properties are presented in Table 1. Particle size was
- analysed through laser diffraction (Hydro 2000MU, Malvern Instruments Ltd.,
- 116 UK). Soil organic matter content (dry weight basis) was determined by mass loss
- 117 on ignition (450 °C for 24 h). Total carbon and nitrogen content (%) were
- assessed using an Elementar Vario EL III elemental analyser (Hanau, Germany).
- 119 2.3 BC amendment and soil spiking
- 120 Prior to BC amendment, the soil was rehydrated with deionized water to field
- 121 moisture content (30 35% dry weight basis). Subsequently, soil treatments
- 122 with 1% (dry weight basis) of three different types of BC (designated as P3-1, CP
- 123 2 and BP 2, properties presented in Table 2) were prepared by blending specific

124 quantities of BC with each treatment using a stainless spoon [2]. A treatment 125 without BC was also prepared as a control. Immediately after BC amendment, 126 soils were spiked with ¹²C-/¹⁴C-phenanthrene using acetone as carrier (3.75 ml 127 per 300 g dry soil at 0.8 mg/ml for ¹²C- and 6666.67 Bq/ml for ¹⁴C-phenanthrene) 128 as described by Doick et al. [19], to achieve a ¹²C-phenanthrene concentration of 129 10 mg kg⁻¹ and ¹⁴C-phenenthrene-associated radioactivity of 64 – 78 kBg kg⁻¹ dry 130 soil. Unspiked control soils were also prepared for each BC treatment. As mineralisation of phenanthrene by both indigenous and inoculated 131 132 microorganism has been shown to be equally efficient and dependent solely on 133 the available amount of phenanthrene [20, 21], the soil samples were not 134 sterilised after spiking and were incubated in sealed amber glass jars at room 135 temperature $(21 \pm 1 \circ C)$ for 1, 25, and 50 d. 136 2.4 Preparation of phenanthrene-degrading inoculum 137 Prior to the mineralization assay, a phenanthrene-degrading inoculum of 138 *Pseudomonas* sp. was cultured in a mixture of minimal basal salts solution (MBS) 139 containing phenanthrene solution (0.1 ml l⁻¹) as the sole C-source [22] on an IKA 140 Labortechnik KS501 digital orbital shaker at 100 rpm at room temperature (21 ± 141 1 °C). On the fourth day of incubation (late exponential phase of growth), the 142 inoculum was concentrated by centrifugation at 10,000 x g for 30 minutes 143 (Hettich Zentrifugen, Rotanta 460, UK). The supernatant was then discarded and 144 the cell pellet washed and re-suspended with fresh MBS. A second centrifugation 145 was subsequently carried out to ensure the removal of any residual 146 phenanthrene, obtaining a final cell density of approximately 10⁸ cells ml⁻¹. 147 2.5 Mineralization of ¹⁴C-phenanthrene

148 Mineralization assays were conducted in 'respirometers', which were modified 149 250 ml Schott bottles as described by Reid et al. [22]. After 1, 25 and 50 d of soil 150 incubation, the respirometers (n = 3) were set up with 10 ± 0.2 g soil wet weight 151 (~7.5 g dry soil), 25 ml of MBS, and 5 ml of concentrated inoculum ($10^5 - 10^6$ 152 cells per g soil) [23]. Uninoculated respirometers (n = 3) and soil incubations 153 with no 14 C-activity (n=3) were also set up for each treatment. The 154 respirometers were then incubated on an IKA Labortechnik KS501 digital orbital 155 shaker at 100 rpm for 14 days at room temperature (21 ± 1 °C). During this 156 period of time, ¹⁴CO₂ generated from microbial degradation of ¹⁴C-phenanthrene 157 was trapped in 7 ml glass scintillation vials suspended from the Teflon lined-lid 158 containing 1 ml of NaOH (1 M). The vials were replaced every 24 h, after which 5 159 ml Goldstar scintillation cocktail was subsequently added to each of the sampled 160 vials and the ¹⁴C-associated activity was quantified by liquid scintillation 161 counting (LSC, Canberra Packard Tri-Carb2250CA) after a >12h storage in the 162 dark to avoid chemo-luminescence. 163 *2.6 Extraction of ¹⁴C-phenanthrene with hydroxylpropyl-β-cyclodextrin (HPCD)* 164 solutions 165 Three different HPCD solutions (60mM) were prepared in deionized water (pH 166 5.89), and phosphate buffers of pH 7 and 8 respectively. The buffers of pH 7 and 167 8 were prepared by combining K_2 HPO₄ (0.2 M) and KH₂PO₄ (0.2 M) solutions at 168 ratios of 1.6:1 and 17.9:1 respectively. The extraction assays were carried out 169 after 1, 25 and 50 days of ageing, following the methodology described by Reid *et* 170 al. [10]. In brief, soil $(1.25 \pm 0.1 \text{ g wet weight})$ from each treatment was weighed 171 into 35 ml Teflon centrifuge tubes with 25 ml of each HPCD solution (n = 3). The

tubes were then placed onto an orbital shaker (IKA Labortechnik KS501 digital)

- 173 at 100 rpm for 22 h in darkness at room temperature (21 ± 1 °C). Subsequently,
- the tubes were centrifuged at 3000 x g for 1 h (Hettich Zentrifugen, Rotanta 460,

175 UK) and 5 ml of supernatant was then mixed with 15 ml Goldstar scintillation

- 176 cocktail. The samples were assessed by LSC as described previously.
- 177 2.7 Statistical analysis
- 178 Following blank-correction, statistical analysis of the results was carried out
- 179 with the Statistical Package for the Social Sciences (SPSS Version 22 for Mac).
- 180 The statistical significance of BC addition, BC type and ageing period to
- 181 phenanthrene biodegradability and phenanthrene extractability by HPCD
- 182 solutions, as well as the statistical significance of pH to HPCD extractive
- 183 capability, was determined using a linear model (ANOVA, Tukey Test) and/or
- 184 Student t-test at 95% confidence level (p < 0.05).
- 185 **3. Results and discussion**
- 186 *3.1 Mineralization of* ¹⁴*C*-phenanthrene in soils
- 187 ¹⁴C-Phenanthrene catabolism was drastically reduced in all BC-treated soils at all
- 188 time points. Compared to soil without BC, the fastest rates (the highest yield of
- $^{14}CO_2$ per day during mineralisation assays) and extents and of ^{14}C -
- 190 phenanthrene mineralization decreased by more than 99% at 1 and 25 d, and
- 191 more than 93% after 50 d of soil incubation (Table 3, 4). The fastest rates of
- 192 phenanthrene mineralisation did not exceed 0.10% per d at 1 and 25 d and were
- less than 0.3% per d at 50 d in BC-amended soils (Table 3). At 1 d, only 0.15%,
- 194 0.07%, and 0.11% of ¹⁴C-PAH was mineralized in soils amended with P3-1, CP 2,
- and BP 2 respectively, while 63.20% of the ¹⁴C-phenanthrene was mineralised in
- soil without BC. Furthermore, influences of BC type on biodegradation were
- 197 observed. At 25 d contact time, soil amended with CP 2 yielded significantly less

198 (p < 0.05) ¹⁴CO₂ than the other two BC-amended soils, while significantly more 199 ¹⁴C-phenanthrene (p < 0.05) was mineralized in soil with P3-1 at 50 d than the 200 other two BC-treated soils. Overall, these results were in agreement with 201 previous studies by Rhodes et al. [2, 8]. These trends have been attributed to the 202 strong sorptive capacity of BC [24, 25]. Consequently, the aqueous concentration 203 and biodegradation of target compound was reduced, as the microbial uptake of 204 organic substances mainly takes place in soil aqueous phase [14, 15]. Moreover, 205 a fraction of ¹⁴C-phenanthrene may have become inaccessible to microorganisms 206 due to entrapment in collapsed pores on BC particles [4, 26]. However, the extent to which ¹⁴C-phenanthrene mineralization was inhibited in BC-amended soils 207 208 was much greater than those observed by Rhodes et al. [2, 8]. At least 6% of 209 spiked ¹⁴C-phenanthrene was mineralised in each soil treatment in research by 210 Rhodes *et al.* (2008) all treatments (0 - 5% AC dry weight) in the study by 211 Rhodes et al. [2], while Rhodes et al. [8] only obtained biodegradation extents 212 lower than 1% in soils treated with 5% AC. It appears that the types of BC used 213 in this study possessed greater sorptive capacity than those used in studies by 214 Rhodes et al. [2, 8]. The BC type also influenced the rates and extents of 215 mineralisation of ¹⁴C-phenanthrene in the present study (Table 3, 4). These 216 variations may be attributed to the specific properties of each BC such as surface 217 heterogeneity and functional groups, pore volume, activation and production 218 methods, source material, as well as processing temperature [3, 4, 27-29]. 219 After 50 d ageing, all BC-amended soils yielded greater ¹⁴C-phenanthrene 220 mineralization compared to 1 and 25 d. Unlike BC-amended soils, biodegradation 221 of ¹⁴C-phenanthrene in soil without BC decreased significantly (p < 0.05) over 222 time (Table 4). Apparently, ageing effect, where biodegradability of HOCs

diminishes over time [30], was absent in BC-amended soils. Similar results were

also obtained by Rhodes *et al.*, who attributed such findings to sorptive

attenuation, where soil organic matter (SOM) competes for limited sorption sites

- on BC particles and blocks them from the spiked chemical, thus lowering
- sorptive capacity of BC for target substances [2, 31]. Such competitive sorption

has also been observed by other researchers; for example, Wang *et al.* [28] found

that organic chemicals with larger molecular sizes covered the binding sites on

230 BC particles and blocked them from smaller molecules.

231 3.2 HPCD extraction of ¹⁴C-phenanthrene in soils

232 Addition of BC also led to drastic reduction in the extractability of the ¹⁴C-PAH by

233 HPCD solutions in each soil treatment at all soil-contaminant contact times

234 (Table 5). Compared to the soil without BC, ¹⁴C-activity extracted by unbuffered

aqueous HPCD solution decreased by more than 99% at 1 and 25 d, while 93 –

236 98% less ¹⁴C-phenanthrene was extracted by the unbuffered solution after 50 d

237 of ageing. Extractions with buffered HPCD solutions were also strongly

influenced by the presence of BC, but the phosphate buffer also resulted in

changes in amounts of ¹⁴C-phenanthrene extracted by HPCD. At 1 and 25 d,

240 HPCD at pH 7 extracted 1.03 – 1.56% of spiked phenanthrene from all BC-treated

soils (Table 5), which was statistically higher (p < 0.05) than the amounts

extracted by the other two solutions. Further increase in pH of HPCD solution to

243 8 led to statistically similar (p > 0.05) yield of extracted ¹⁴C-activity from all BC-

amended soils compared to its aqueous counterpart at 1 and 25 d (Table 5).

245 However, after 50 days of soil incubation, HPCD at pH 8 extracted significantly

246 more (p < 0.05) ¹⁴C-activity than the other two solutions in soils amended with

247 CP 2 and BP 2, while the amounts of ¹⁴C-phenanthrene extracted by HPCD at pH

248 7 were statistically similar (p > 0.05) to the values from extractions using
249 aqueous HPCD solution for all BC-amended soils (Table 5).

250 HPCD is a well-established non-exhaustive extraction technique to measure 251 microbial bioaccessibility of numerous HOCs in soils under different conditions 252 [11, 20, 32-38]. The HPCD molecules are able to separate organic compounds 253 from water solution, thus mimicking microbial uptake of organic substances and 254 driving mass-transfer of target compound from soil matrix to dissolved phase [10, 39-42]. However, in presence of BC, sorption of the ¹⁴C-PAH resulted in 255 256 reduction of dissolved phenanthrene for HPCD molecules to separate. Moreover, 257 Jonker and Koelmans [16] suggested that water, as the solvent of aqueous HPCD 258 solution, was not capable of displacing BC-associated phenanthrene molecules 259 from binding sites on BC particles. In the present study, buffered solutions of pH 260 7, at 1 and 25 d, as well as that of pH 8, at 50 d, enhanced the extractive 261 capability of HPCD solutions in soils amended with BC. Additionally, the buffered 262 extracts from each soil treatment at each time point were highly coloured. This 263 was consistent with the observations made by Reid et al. [10] and was indicative 264 of the existence of dissolved organic matter in the extracts [10, 43]. The 265 promotion of dissolution of SOM by phosphate buffers at higher pH has been 266 reported in previous studies [44-46]. It was suggested that the deprotonation 267 under basic conditions brought by phosphate buffers attenuated the association 268 between SOM and soil minerals, thus increasing the amount of dissolved organic 269 matter (DOM) [44]. Although other researchers also demonstrated the ability of 270 phosphate to inhibit the sorption of phenanthrene in soils [47], the effects of 271 phosphate itself in this research on the release of phenanthrene from soil are 272 considered minimal given the amount of phosphate used and its contact time

273 with spiked soils. DOM subsequently contribute to greater solubility of

phenanthrene [18], so that there were more PAH molecules in the aqueous phasefor HPCD molecules to separate.

276 Interestingly, increases in pH did not always result in increases in extraction 277 using HPCD solutions, as a biphasic feature of increasing pH was identified in BC 278 amended soils at 1 and 25 d, and was absent after 50 days of ageing. This 279 observation reflects the complex interactions between soil, BC, phenanthrene and HPCD solutions. It is therefore postulated that extensive sorption of both 280 281 SOM and ¹⁴C-phenanthrene to BC particles was achieved shortly after BC 282 amendment and PAH spiking, while at 1 and 25 d, HPCD at pH 8 dissolved so 283 much BC-associated SOM that sorption sites on BC particles were exposed to 284 phenanthrene. Consequently, greater sorption of phenanthrene to BC was 285 facilitated. At 50 d, however, greater amount of SOM was attached to BC particles 286 and phenanthrene molecules partitioned deeper into BC. As a result, buffer of pH 287 8, which was able to dissolve more SOM, released more BC- and SOM-bound 288 phenanthrene than buffer of pH 7 (Fig. 1). 289 A simple comparison between HPCD extraction assays and mineralisation assays 290 conducted in BC-amended soils was carried out by calculating the ratios of

extraction to mineralisation. The results indicated that in most cases aqueous

HPCD extracted underestimated mineralisation of ¹⁴C-phenanthrene (Table 6).

293 However, the extents to which biodegradation was underestimated were not as

294 great as those reported by Rhodes et al. [2] except for few cases (Table. 6). Such

295 findings suggest that the differences between HPCD extractive capability and

biodegradability of phenanthrene in soils with BC may not be as great as they

297 were previously observed. The amounts of phenanthrene extracted by HPCD in

298 pH 7 were 3 to 15 times greater those degraded by microorganisms at 1 and 25 d, 299 and mildly deviated from the degraded amounts at 50 d (Table 6). HPCD in pH 8 300 provided mixed results in ratios of extraction to mineralisation in all BC-treated 301 soils, but the deviations of extractability from biodegradability were not as great 302 as those demonstrated by aqueous HPCD and HPCD in pH 7. These findings have 303 two implications. Firstly, the mechanism which was direct degradation of BC-304 associated phenanthrene proposed in previous studies may not be actually 305 involved in mineralisation assays. Secondly, increasing pH enhances the 306 extractive capability of HPCD and could improve this method in predicting 307 microbial accessibility of phenanthrene in soils with BC after substantial ageing 308 period, as the ratios of HPCD extraction in buffers were approach 1 compared to 309 those of aqueous HPCD (Table 6). However, due to the size of the data acquired 310 in the current study, further verification of these findings are required to 311 optimise this modification of HPCD extraction under various conditions 312 including different pH values, and soil and BC types. Besides, the order of BC 313 amendment and PAH spiking should also be considered as the faster and greater 314 binding of PAH with BC particles in pre-amended soils may occur, thus bringing 315 differences to the results obtained. 316 4. Conclusion

317 Addition of BC significantly reduced mineralisation and extraction of ¹⁴C-

318 phenanthrene after different periods of soil-contaminant interactions, where

319 variations brought by BC type were identified among soil treatments.

320 Introduction of phosphate buffers produced varying effects to the extractive

321 capability of HPCD solutions, as HPCD at pH 7 extracted significantly more

phenanthrene at 1 and 25 d, and HPCD at pH 8 yielded more extracted ¹⁴Cactivity at 50 d.

324 A biphasic feature of increasing pH on HPCD extractive capability was observed 325 at 1 and 25 d but not at 50 d. overall, these findings reflected the complex 326 interactions between SOM, BC, HPCD, and phenanthrene. Aqueous HPCD 327 extractions did not always underestimate biodegradation of phenanthrene in BC-328 amended soil, rejecting previously proposed mechanism for the incompatibility 329 between mineralisation and HPCD extraction. More studies should be carried out 330 to find out whether presence of BC indeed leads to underestimation of biodegradation by HPCD extraction. If yes, increasing pH of HPCD solution, as it 331 332 has been demonstrated in this study, is a viable approach to modifying this

technique for better prediction of the bioaccessibility of organic contaminants in

soils with BC.

Table 1. Physical-chemical properties of soil used in this study. Errors are shown as 1 SEM (n = 3).

336	SEM $(n = 3)$.			
	Soil Properties		Pai	rameter Value
	pH (in dH ₂ O)			5.36 ± 0.01
	Organic matter (%)			9.15 ± 0.06
	Nitrogen (%)			0.20 ± 0.02
	Carbon (%)			2.24 ± 0.01
	Particle size*	Clay		23.42%
		Silt		75.26%
		Sand		1.27%
		Soil texture		Silt loam
005	* 4 1	11.00		C

337 * Analysis of particle size by laser diffraction reflected the distribution of particles with

diameter < 1 mm, using total surface area as baseline.

	Activated Carbon	Source	Activation method	Processing Temperature	Surface area (m² g ⁻ ¹)	Pore volume (cm ³ g ⁻¹)	Mean particle diameter (µm)
_	P3-1	Wood	Chemical activation	700°C	1150	Not provided	Not applicable
	BP 2	Coal	Steam activation	850-950°C	1000	1.56	21
	CP 2	Coconut shell	Steam activation	850-950°C	950	0.55	21

339 Table 2. Properties of black carbon used for soil amendments.

340 a Particle size of this grade was expressed as distribution of powder size: $<150 \mu m = 95 - 100\%$, $<75 \mu m = 85 - 95\%$, $<45 \mu m = 65 - 85\%$.

Table 3. Fastest rates of ¹⁴C-phenanthrene mineralization in soils amended with 0% black carbon and 1% P3-1, CP 2, and BP 2 at 1, 25

342 and 50 days of soil-phenanthrene interactions. Values are the % ¹⁴CO₂ per d mean (n = 3) ± standard error of the mean (SEM). Values in

343 the same column followed by the same letter, or row followed by the same number are statistically similar (student t-test and ANOVA

Ageing		Black carb	on treatment	
period (days)	0% BC	1% P3-1	1% CP2	1% BP2
1 day	26.13 ± 1.73 ^{a1}	0.03 ± 0.01^{a2}	0.02 ± 0.00^{a2}	0.03 ± 0.00^{a2}
25 day	13.78 ± 1.43^{b1}	0.05 ± 0.01^{a2}	0.02 ± 0.01^{a2}	0.10 ± 0.01^{b3}
50 day	3.85 ± 0.16^{c1}	0.25 ± 0.00^{b2}	0.04 ± 0.01^{a3}	0.06 ± 0.01^{c3}

344 Tukey test, n = 3, p < 0.05).

Table 4. Total extents of ¹⁴C-phenanthrene mineralised by microorganisms in soils amended with 0% black carbon and 1% P3-1, CP 2,

and BP 2 after 1, 25, and 50 days of soil-phenanthrene interactions. Values are the % mean (n = 3) ± standard error of the mean (SEM).

347 Values in the same column followed by the same letter, or row followed by the same number are statistically similar (student t-test and

348 ANOVA Tukey test, *n* = 3, p < 0.05).

Ageing		Black carb	oon treatment	
period (days)	0% BC	1% P3-1	1% CP2	1% BP2
1 day	63.20 ± 0.52^{a1}	0.15 ± 0.03^{a2}	0.07 ± 0.01^{a2}	0.11 ± 0.02^{a2}
25 day	38.79 ± 1.01^{b1}	0.24 ± 0.04^{a2}	0.09 ± 0.04^{ab3}	0.29 ± 0.02^{b2}
50 day	21.29 ± 0.98^{c1}	1.46 ± 0.00^{b2}	0.21 ± 0.03^{b3}	0.38 ± 0.06^{b3}

Table 5. ¹⁴C-Phenanthrene extracted by HPCD solutions from soils amended with 0% black carbon and 1% P3-1, CP2, and BP2 after 1, 25 and 50 days of soil-phenanthrene interactions. Values are the % mean (n = 3) ± standard error of the mean (SEM). At each time point,

values in the same column followed by the same letter are statistically similar; values in the same column generated from the extraction assays with the same HPCD solution followed by the same Greek letter are statistically similar; values in the same row followed by the

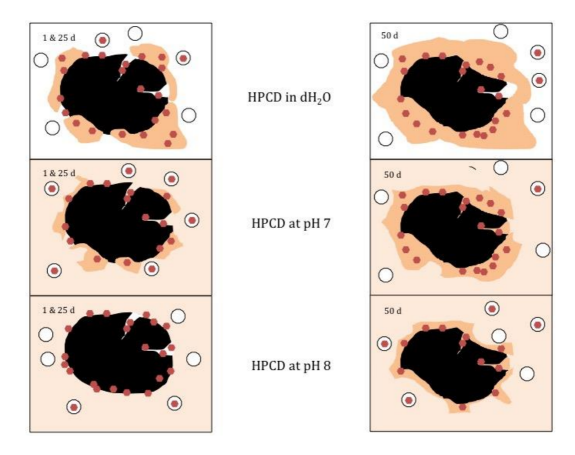
353 same number are statistically similar (student t-test and ANOVA Tukey test, n = 3, p <0.05).

Ageing	HPCD	Black carbon treatment				
period (days)	solution	0% BC	1% P3-1	1% CP2	1% BP2	
1 day	dH ₂ O	74.16 ± 0.39 ^{aα1}	$0.08 \pm 0.08^{a\alpha 2}$	$0.08 \pm 0.02^{a\alpha 2}$	$0.10 \pm 0.10^{a\alpha 2}$	
	рН 7	$74.96 \pm 0.80^{a\alpha 1}$	$1.37 \pm 0.07^{b\alpha 2}$	$1.03 \pm 0.21^{b\alpha 3}$	$1.10 \pm 0.14^{b\alpha 3}$	
	pH 8	$72.70 \pm 1.46^{a\alpha 1}$	$0.05 \pm 0.05^{a\alpha 2}$	$0.11 \pm 0.08^{a\alpha 2}$	$0.11 \pm 0.11^{a\alpha 2}$	
25 day	dH ₂ O	$14.29 \pm 1.05^{a\beta 1}$	$0.01 \pm 0.01^{a\alpha 2}$	$0.06 \pm 0.06^{a\alpha 2}$	$0.02 \pm 0.02^{a\alpha 2}$	
	pH 7	$24.71 \pm 1.35^{b\beta1}$	$1.56 \pm 0.17^{b\alpha 2}$	$1.11 \pm 0.09^{b\alpha 3}$	$1.01 \pm 0.17^{b\alpha 3}$	
	pH 8	$31.69 \pm 0.06^{c\beta 1}$	$0.20 \pm 0.12^{a\alpha 2}$	$0.20 \pm 0.13^{a\alpha\beta^2}$	$0.13 \pm 0.10^{a\alpha^2}$	
50 day	dH ₂ O	$3.76 \pm 0.35^{a\gamma 1}$	$0.24 \pm 0.15^{a\alpha 2}$	$0.05 \pm 0.01^{a\alpha 2}$	$0.08 \pm 0.06^{a\alpha 2}$	
	pH 7	$8.90 \pm 0.89^{b\gamma 1}$	$0.52 \pm 0.07^{a\beta 2}$	$0.09 \pm 0.07^{a\beta3}$	$0.05 \pm 0.04^{a\beta3}$	
	pH 8	$12.55 \pm 0.28^{c\gamma 1}$	$0.64 \pm 0.33^{a\alpha 2}$	$0.61 \pm 0.12^{b\beta 2}$	$0.44 \pm 0.06^{b\alpha 2}$	

Table 6. The ratios of the amounts of extracted ¹⁴C-activity to that of mineralised

HPCD	Ageing		BC treatme	nt
solution	period	1% P3-1	1% CP2	1% BP2
dH2O	1 day	0.55	1.14	0.85
	25 day	0.05	0.68	0.07
	50 day	0.26	0.06	0.42
рН 7	1 day	9.30	15.18	9.73
	25 day	6.50	12.36	3.49
	50 day	0.48	0.75	1.15
pH 8	1 day	0.31	1.61	1.01
	25 day	0.83	2.22	0.45
	50 day	0.67	2.73	1.78

355 ¹⁴C-activity in BC-amended soils.



356

Fig. 1. Proposed mechanism for the biphasic feature of increasing pH on

extractive capability of HPCD solutions at 1 and 25 d, and the absence of this

- feature at 50 d. At 1 and 25 d, HPCD at pH 8 dissolved large quantity of BC-
- associated SOM and exposed sorption sites on BC to phenanthrene, leading to
- 361 greater sorption of phenanthrene to BC particles. At 50 d, more SOM and
- 362 phenanthrene was attached to BC, HPCD at pH was more capable of dissolving
- 363 SOM and therefore released more BC- and SOM- bound phenanthrene.

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