1	The effect of organic acids on the behaviour and biodegradation of ¹⁴ C-phenanthrene in
2	contaminated soil
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19 Abstract

20 The interaction between root exudates and soil microbes has been hypothesised as the 21 primary mechanism for the biodegradation of organic pollutants in the rhizosphere. However, 22 the mechanisms governing this loss process are not completely understood. This study aimed 23 to investigate the effect of two important compounds within root exudates (citric and malic acid) on ¹⁴C-phenanthrene desorption and bioaccessibility in soil. Overall results showed that 24 the presence of both citric and malic acid (> 100 mmol l^{-1}) enhanced the desorption of $l^{-1}C$ -25 26 phenanthrene; this appeared to be concentration dependant. Increases in extractability were 27 not reflected in a higher bioaccessibility. Despite enhancing the desorption of ¹⁴Cphenanthrene in soil, there is no direct evidence indicating that citric or malic acid have the 28 29 ability to promote the biodegradation of ¹⁴C-phenanthrene from soil. Results from this study 30 provide a novel understanding of the role that substrates, typically found within the 31 rhizosphere due to root exudation, play in the bioaccessibility and biodegradation of 32 hydrocarbons in contaminated soil.

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34 Keywords: Phenanthrene, organic acids, root exudates, desorption, bioavailability, ageing

36 1 Introduction

37 The rhizosphere is defined as the soil in closest proximity to plant roots and has been 38 hypothesised to enhance the biodegradation of organic contaminants such as aliphatic and 39 aromatic hydrocarbons through different mechanisms (Anderson et al., 1993; Pilon-Smits, 40 2005). These include the promotion of (1) larger microbial populations (Anderson et al., 41 1993) and shifts on their community composition (Joner et al., 2002), (2) source of 42 biologically important substrates including nutrients and readily available sources of carbon 43 (Reilley et al., 1996; Dakora & Phillips, 2002; Martin et al., 2014; Sivaram et al., 2019), and 44 (3) increasing the bioavailability of the contaminants due to root exudates, decay and 45 turnover (Siciliano & Germida, 1998; Martin et al., 2014; Sivaram et al., 2019). The amount 46 and type of substances released by roots is highly dependent on a series of factors including 47 plant species and age, as well as particular soil and environmental conditions (Jones, 1998; 48 Shukla et al., 2011; Agnello et al., 2014; Martin et al., 2014). However, a number of low 49 molecular weight compounds, such as amino acids, sugars and organic acids, have been 50 identified as common constituents of root exudates (Jones et al., 2003; van Hees et al., 2005). 51 Organic acids, including citric, malic and oxalic are reported to be the most abundant (Jones 52 & Brassington, 1998; Ling et al., 2015); therefore, it is reasonable to consider the role that 53 these acids might play in influencing the extractability and the bioavailability of different 54 contaminants and how this might influence their biodegradation.

The use of root exudates for the dissipation of organic contaminants in soil has been reported (Miya & Firestone, 2001; Yoshitomi & Shann, 2001; Joner *et al.*, 2002). These investigations have used simulated rhizosphere conditions by the introduction of artificial or natural root exudates in order to approach the subject in a more controlled manner (Miya & Firestone, 2001; Joner *et al.*, 2002). From these studies, research has been developed to consider the 60 effect of these substances on the shifts of the microbial populations and/or communities 61 (Joner et al., 2002; Shukla et al., 2011), overall dissipation of contaminants (Joner et al., 62 2002), and their effect on soil physical and chemical properties (Shukla et al., 2011). Authors 63 such as Sun et al. (2013), Martin et al. (2014) and Gao et al. (2015) have pointed out that 64 although efforts have been directed towards investigating the effect of root exudates on the 65 biodegradation of hydrocarbons in contaminated soil, information regarding the role of single compounds from this solution is scarce. Within these few studies, it has been reported that 66 67 organic acids commonly found in root exudates can promote the desorption of phenanthrene 68 from soil (Gao et al., 2010b, 2015b; Ling et al., 2015).

69 It has been observed that changes in the extractability of polycyclic aromatic hydrocarbons 70 (PAHs) might act as a predictor of the microbial degradability of different species of PAHs. 71 Specifically, the rates of desorption of some PAHs have successfully been used as predictors 72 of their biodegradation (Cornelissen et al., 1998a). As fractions of hydrocarbons are 73 transferred from soil to solution through the desorption process; these can also become more 74 bioaccessible and susceptible to be metabolised by the soil microbial community (Semple et 75 al., 2003, 2007, 2013). Therefore, the possibility of enhancing the desorption of PAHs by 76 using organic acids to promote or enhance biodegradation in soil has been identified as a 77 promising strategy, but remains poorly explored (Martin et al., 2014). In addition, the extent 78 into which these organic acids affect the biodegradation of the desorbed hydrocarbon has not 79 been considered. Therefore, the aim of this study was to investigate the effect of two low 80 molecular weight organic acids (LOAs) commonly found within root exudates in the extractability and bioaccessibility of ¹⁴C-phenanthrene contaminated soil. Phenanthrene was 81 82 selected as a model PAH given its widespread distribution, biodegradability and persistent 83 properties in soil. For this, mineralisation, hydroxypropyl-β-cyclodextrin (HPCD) extractability and desorption kinetics of ¹⁴C-phenanthrene were assessed in the presence of 84

organic acids at a range of concentrations. Results from this experiment provide a novel
perspective of the effect of organic acids on the fate of ¹⁴C-phenanthrene soil by investigating
(1) the desorbing capacity of citric and malic acid and (2) the extent by which these can
promote a higher bioavailability and mineralisation.

89 2 Methodology

90 2.1 Soil preparation and spiking

91 An uncontaminated clay loam soil (top 20 cm, 2.7 % organic matter) was collected from 92 Myerscough Agricultural College, Preston, U.K. Partially air-dried soil (24 h) was passed 93 through a 2 mm sieve and stored in the dark at 4 ± 1 °C until needed. Main soil physical and 94 chemical characteristics have been described by Towell et al. (2011). Sieved soil was 95 rehydrated (50% water holding capacity (whc)) and spiked following the procedure proposed 96 by Doick et al. (2003). In short, this approach consists on the application of the standards to a 97 fraction of the total amount of soil (inoculum) followed by gradual mixing and incorporation of the remaining soil with a stainless steel spoon. Standards used for spiking contained ^{12/14}C 98 99 phenanthrene dissolved in acetone to deliver a final concentration of 100 mg kg⁻¹ (dw) phenanthrene with an associated ¹⁴C-activity of 83 Bq g⁻¹ (dw). Spiked soil was placed in 100 101 sealed sterilized amber jars and incubated in the dark at 21 ± 1 °C in a controlled environment room for up to 15 weeks. Determination of the total ¹⁴C-phenanthrene associated activity in 102 103 the soil was assessed at every time point by sample oxidation following the methodology 104 described by Rhodes et al. (2012).

106 **2.2 Influence of organic acids on the mineralisation of** ¹⁴C-phenanthrene

107 Mycobacterium gilvum has been previously shown to degrade a range of hydrocarbons, 108 including PAHs such as naphthalene, fluorine, phenanthrene, anthracene, fluoranthene and 109 pyrene and as sole or primary source of carbon (Xiong et al., 2017; Posada-Baquero et al., 110 2019). In addition, this strain has also been isolated from chronically contaminated 111 environments such as soil from a former coking plant (Xiong et al., 2017), indicating its 112 ability to adapt to these type of conditions, utilising hydrocarbons as their primary source of 113 carbon under nutrient limiting conditions, which are typically observed on highly 114 contaminated sites. To assess this, the mineralisation of ¹⁴C-phenanthrene was measured 115 using the methodology developed and tested by Reid et al. (2001) and Semple et al. (2006). 116 Soil (10 g dw) aged over 14 and 50 days was placed into 250 ml modified Schott bottles fitted with a 1 M NaOH ${}^{14}CO_2$ trap (n = 3). These ageing times were selected based on that 117 reported by Kelsey et al. (1997), where ¹⁴C-phenanthrene mineralisation was significantly 118 119 reduced within the first few weeks and then remained stable for 50 days. To assess the effect 120 of organic acids towards the mineralisation process, citric, malic, oxalic and succinic acids 121 were selected as representative LOAs often observed in the rhizosphere (van Hees *et al.*, 122 2005). Solutions containing individual LOAs within its naturally appearing range in the 123 rhizosphere soil solution (0.1 and 0.5 mmol 1⁻¹) (van Hees *et al.*, 2005) were used. These were 124 incorporated into the minimal basal salts (MBS) medium and used for the mineralisation 125 assay. For the mineralisation assays, soil was mixed with the MBS containing the organic acids (25 ml) and 5 ml of a bacterial inoculum of M. gilvum (10^5 cells ml⁻¹) to achieve a final 126 3:1 liquid:soil ratio. Bottles were then placed onto an orbital shaker at 100 rpm in a controlled 127 environment room at 21 ± 1 °C in the dark. ¹⁴CO₂ evolution was assessed by periodically (up 128 129 to every 24 h) replacing the trap, mixing with 5 ml liquid scintillation cocktail and assessed 130 by liquid scintillation counting (LSC) (10 min - Canberra Packard Tri-Carb 2300, U.K.).

132 **2.3 Influence of organic acids on the extractability of ¹⁴C-phenanthrene**

133 2.3.1 Preliminary tests

134 A series of preliminary tests were carried in order to optimize the general experimental 135 parameters and design of the extraction assays. Solutions of deionised water containing citric, 136 malic, oxalic and succinic acids solutions of individual organic acids were prepared at 0.1 and 0.5 mmol l⁻¹. Desorption kinetics of ¹⁴C-phenanthrene with these solutions (n = 3) were 137 138 assessed from spiked soil following the methodology described below. The temporal effect of organic acids on the bioaccessibility of ¹⁴C-phenanthrene was also 139 140 assessed (n = 3). Soil was saturated with malic acid solution (100 % whc) at two concentrations (0.5 and 500 mmol 1^{-1}) and incubated in a controlled environment room 21 ± 1 141 142 °C for 1, 3, 6, 8 or 24 h. Each experimental unit was also fitted with a 1 M NaOH ¹⁴CO₂ trap for the assessment of any possible dissipation of ¹⁴C-phenanthrene by microbial respiration 143 144 during the incubation time. Soil was extracted with 50 mM HPCD solutions after each incubation time following the methodology described below. ¹⁴CO₂ traps were assessed by 145 adding 5 ml of liquid scintillation cocktail and assessed by LSC as previously described. 146 147 Citric, malic, oxalic and succinic acids did not impact significantly on the desorption of ¹⁴Cphenanthrene from the soil (Table SI-1) when compared against the control (p > 0.05). 148 Despite of this, soil extracted with citric and malic acid at 0.5 mmol 1⁻¹ were the only two 149 treatments presenting higher rapidly desorbing fractions (F_{rap}), with (44.16 %) and (49.76 %) 150 151 respectively, than the control (40.94 %). Based on this, these two organic acids were selected 152 for further investigation at a wider range of concentrations (0.5, 100, 250, 500 and 1000 mmol 1⁻¹) in order to assess the full potential of these compounds to impact the desorption of 153

¹⁴C-phenanthrene in soil. Selected concentrations ranged from naturally appearing LOAs
concentrations (van Hees *et al.*, 2005) up to maximum tested concentrations within
experiments with similar aims (Gao *et al.*, 2015a; Ling *et al.*, 2015).

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158 2.3.2 HPCD extraction of ¹⁴C-phenanthrene from soil

159 Changes in the bioaccessibility of ¹⁴C-phenanthrene were measured by HPCD extractions from soil aged over 1 and 15 weeks. At each time point, 1.25 g soil (dw) were placed into 160 161 Teflon centrifuge tubes (n = 5); soil was saturated (100 % whc) with citric and malic acid solution (0.5, 100, 250, 500 and 1000 mmol 1⁻¹). Sealed tubes where incubated in a controlled 162 163 environment room $(21 \pm 1 \text{ °C})$ for 8 h. Then, 25 ml of 50 mM HPCD solution was added. 164 Tubes were placed onto an orbital shaker (100 rpm) for 22 h at 21 ± 1 °C. Afterwards, samples 165 were centrifuged (3000 x g for 1 h) and 5 ml of the supernatant was placed in a glass scintillation vial and mixed with 15 ml liquid scintillation cocktail. Samples were assessed 166 through LSC as described previously. Remaining ¹⁴C-associated activity in the soil was 167 168 assessed by sample oxidation (Rhodes et al., 2012).

169

170 2.3.3 Desorption of ${}^{14}C$ -phenanthrene by organic acids

Tests were performed following a randomized design (n = 5) and blind sampling. Desorption kinetics were assessed after 1 and 15 weeks soil-contaminant contact time; at each time point, 4 g soil (dw) was placed into Teflon centrifuge tubes and mixed with 25 ml of organic acid solution at a given concentration (0.5, 100, 250, 500 and 1000 mmol 1⁻¹). Tubes were placed onto an orbital shaker (100 rpm) in a controlled environment room at 21 ±1 °C. Soil samples were sequentially extracted after 1, 4, 6, 12, 24, 45, 90, 180 and 360 h by centrifuging at 3000

- 177 x g for 1 h. Aliquots (5 ml) were mixed with 15 ml liquid scintillation cocktail in a glass
- 178 scintillation vial and assessed by LSC. Residual activity in the soil after the last extraction
- 179 was assessed by sample oxidation as described by Rhodes *et al.* (2012).
- 180 Desorption of ¹⁴C-phenanthrene was examined by two- (Equation 1) and three-compartment
- 181 (Equation 2) first-order kinetics (Cornelissen *et al.*, 1998b; Rhodes *et al.*, 2010):
- 182 Equation 1:

183
$$S_t / S_0 = [F_{rap} \bullet \exp(-k_{rap} \bullet t)] + [F_{slow} \bullet \exp(-k_{slow} \bullet t)]$$

184 *Equation 2*:

185
$$S_t / S_0 = [F_{rap} \bullet \exp(-k_{rap} \bullet t)] + [F_{slow} \bullet \exp(-k_{slow} \bullet t)] + [F_{very slow} \bullet \exp(-k_{very slow} \bullet t)]$$

where S_t represents the amount of ¹⁴C-phenanthrene sorbed to the soil at the desorption time *t* (h) and S_0 is the initial total amount of ¹⁴C-phenanthrene at the beginning of the assay (time 0). F_{rap} , F_{slow} and $F_{very \ slow}$ (%) are the rapid, slow and very slow desorbing fractions and k_{rap} , k_{slow} and $k_{very \ slow}$ (h⁻¹) are the rate constants for the rapid, slow and very slow desorption, respectively. The model assumes that $k_{very \ slow} \le k_{slow} \le k_{rap}$ (Rhodes *et al.*, 2010; Clegg *et al.*, 2014), and that the addition of the desorbing fractions equals 100 % (Clegg *et al.*, 2014). The values of F_{rap} , F_{slow} , $F_{very \ slow}$, k_{rap} , k_{slow} and $k_{very \ slow}$ were obtained by exponential curve

- 193 fitting using Excel Solver add-in, using a non-linear least squares method.
- 194

195 **2.4 Statistical analysis**

196 Statistical analyses were carried using the SPSS 21 (95 % confidence interval). Normality of

197 the data was verified by Shapiro-Wilk tests, transformations were applied in cases where a

normal distribution was not observed. Analyses of the differences across time were carried by
Student's t-test and Wilcoxon test for normal and not normally distributed data respectively.
Differences between the treatments at each time point were analysed using One-Way
ANOVA (Tukey) or Kruskal-Wallis test for normal and not-normal distributed data
respectively. Graphical representations of the results were done with the software Sigma Plot
2000.

204 **3 Results**

205 **3.1 Short-term impact of organic acids on the mineralisation of** ¹⁴C-phenanthrene in soil

206 The impact of citric, malic, oxalic and succinic acids within a naturally occurring range of concentrations was tested on the mineralisation of ¹⁴C-phenanthrene. Organic acids were only 207 observed to produce significant differences on the mineralisation of ¹⁴C-phenanthrene after 208 209 14 days soil-PAH contact time, while remaining unaffected after 50 days of soil-PAH contact 210 time. The data showed that after a short soil-PAH contact time (14 d), the presence of citric acid (0.1 mmol l⁻¹) resulted in a significantly faster rate of mineralisation (29.52 % d⁻¹) than 211 the control (20.99 % d⁻¹) (F = 2.795, p = 0.016) (Table 1). At this same time point, although 212 not significant (p = 0.077), the lag phase of the control soil was longer (18.28 h) than in soil 213 214 incubated with organic acids (4.11 - 5.42 h).

215

3.2 Preliminary tests for the selection of organic acids and soil-organic acid contact time for the assessment of ¹⁴C-phenanthre bioaccessibility in soil

- 218 As significant differences were not observed within the different organic acids used in the
- 219 preliminary assay looking at their impact in ¹⁴C-phenanthrene desorption kinetics; malic acid

220 was selected as a representative organic acid for the optimisation of the methodology for the 221 assessment of bioaccessibility. Results from the test looking at the temporal impact of malic 222 acid in HPCD-extractable ¹⁴C-phenanthrene fraction showed that soil-organic acid contact time did not have a significant effect on the bioaccessible fraction of this hydrocarbon (p > p)223 0.05). However, data showed that the largest extractable proportion of ¹⁴C-phenanthrene was 224 225 obtained after 8 h of soil-organic acid incubation (control soil, 8.15 %), compared to the 226 lowest value presented after 48 hours (control soil, 1.10 %). Therefore, 8 h soil-organic acid 227 incubation was considered to be the most suitable contact time and consequently selected for 228 further investigation. Furthermore, mineralisation from the HPCD extractable experimental 229 units within the incubation time was observed to be negligible.

230

231 **3.3 Bioaccessibility of ¹⁴C-phenanthrene in soil**

Changes on the bioaccessibility of ¹⁴C-phenanthrene in soil was assessed by HPCD extractions and were observed to be significantly different over time (Table 2, t = 66.682, p < 0.001). After one week of soil-PAH contact time; saturation of soil with organic acids (100 % whc, 8 h) did not have significant effects on the bioaccessibility of ¹⁴C-phenanthrene (F = 1.981, p = 0.059). In the case of soil incubated for 15 weeks, the addition of 500 mmol l⁻¹ citric acid significantly enhanced the bioaccessible fraction of ¹⁴C-phenanthrene (14.92 %)

238 compared to the control (6.72 %) (F = 4.513, p = 0.003).

239

240 **3.4 Desorption of ¹⁴C-phenanthrene with organic acids**

The amount of ¹⁴C-phenanthrene that was desorbed from soil was significantly affected by the presence of organic acids (p < 0.001) (Table 3). Citric acid at the highest concentration

(1000 mmol 1⁻¹) consistently produced a significantly higher desorption than any other 243 244 treatment after 1 week (39.27 %) and 15 weeks (47.86 %) soil-PAH contact time (p < 0.001). 245 Furthermore, this was the only treatment capable of enhancing the desorption of ¹⁴Cphenanthrene after 1 week soil-PAH contact time. The presence of citric acid (0.5 - 250 246 mmol 1⁻¹) and malic acid (0.5 - 500 mmol 1⁻¹) significantly reduced the total desorbable 247 248 fraction of ¹⁴C-phenanthrene soil after one-week soil-PAH contact time (p < 0.001). In 249 contrast, only the lowest concentrations (0.5 mmol l⁻¹) of citric acid (13.04 %) and malic acid (12.51 %) produced significantly lower levels of desorption of ¹⁴C-phenanthrene than the 250 251 control (18.96 %) after 15 weeks soil-PAH contact time. This was in contrast to the desorption behaviour observed at concentrations above 100 mmol l⁻¹ citric acid and 250 252 mmol 1⁻¹ malic acid, where desorbed ¹⁴C-phenanthrene was significantly higher (p < 0.001). 253

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255 3.4.1 Impact of organic acids on ¹⁴C-phenanthrene desorption kinetics

Desorbing fractions (F_{rap} and F_{slow} ; F_{rap} , F_{slow} and $F_{very slow}$) and rate constants (k_{rap} and k_{slow} ; 256 k_{rap} , k_{slow} and $k_{very \ slow}$) from the two- and three-compartment model fitting, respectively, are 257 258 presented on Tables 4 and 5. Squared deviations data showed a better fit by the three-259 compartment one (Table SI-3, p < 0.001); therefore, further analysis was focused on the 260 values estimated by this desorption model. Desorbing fractions (%) and rate constants (h^{-1}) 261 obtained by the three-compartment model (Figures SI 4-9) showed significant differences for 262 all cases (p < 0.001). After one-week soil-PAH contact time, significantly higher fractions of ¹⁴C-phenanthrene were rapidly desorbed by 1000 mmol l⁻¹ citric (19.22 %) and malic acid 263 264 (20.20%) than in the control soil (12.08%). In contrast, lower concentrations of malic acid $(100 \text{ and } 250 \text{ mmol } l^{-1})$ and citric acid $(100 \text{ mmol } l^{-1})$ significantly reduced the rapidly 265 266 desorbing fractions. Rapidly desorbing rate constants were not affected by the majority of the

treatments with the exception of the effect produced by citric acid at 100 mmol l⁻¹. Slowly 267 268 desorbing fractions were significantly reduced by all treatments apart from citric acid (1000 269 mmol 1⁻¹), which was found to be similar to the control. Furthermore, rate constants of this fraction (k_{slow}) were significantly enhanced in most of the treatments (except 0.5 and 1000 270 271 mmol 1⁻¹ citric acid), with a longest slowly desorbing phase produced in the present of malic acid $(0.139 - 0.146 \text{ h}^{-1})$ when compared against the control (0.013 h^{-1}) . Very slowly 272 273 desorbing fractions accounted for the largest phase in all of the treatments. Moreover, organic acids significantly increased this fraction in all treatments (except 1000 mmol l⁻¹ citric acid), 274 ranging from 77.1 to 88.09 % against the 72.12 % when dH₂O was used as extractant. Very 275 276 slowly desorbing rate constants were also significantly higher in the presence of citric (≥ 500 277 mmol l⁻¹) and malic acid at all tested concentrations.

278 After 15 weeks incubation, high concentrations of citric $(500 - 1000 \text{ mmol } l^{-1})$ and malic 279 $(500 \text{ mmol } l^{-1})$ acid were found to significantly enhance the rapidly desorbing fraction of ${}^{14}C$ -280 phenanthrene, representing up to 25.12 % compared to the control (13.11 %). Moreover, low 281 concentrations of both organic acids (0.5 mmol l⁻¹) had the opposite effect, significantly reducing the fraction of ¹⁴C-phenanthrene desorbed to 3.38 and 5.50 % respectively. 282 283 Similarly, rapidly desorbing rate constants were also significantly larger when soil was extracted with citric $(0.5 - 1000 \text{ mmol } 1^{-1})$ and malic acid $(1000 \text{ mmol } 1^{-1})$. Slowly desorbing 284 fractions (F_{slow}) were significantly reduced by all tested concentrations of citric acid and 0.5, 285 100 and 1000 mmol 1⁻¹ malic acid, while the corresponding desorption rate constants 286 287 displayed the opposite behaviour. Fractions desorbed in the very slow phase were 288 significantly increased by all treatments (except 500 mmol l⁻¹malic acid) going from 3.23 % 289 in the control up to 92.50% when soil was treated with 0.5 mmol 1⁻¹malic acid. Very slowly 290 desorbing rate constants were similar to the control with the exception of 100 mmol l⁻¹malic 291 acid where significantly higher values were observed (p < 0.001).

292 4 Discussion

293 4.1 Effect of organic acids in the bioaccessibility of ¹⁴C-phenanthrene in soil

294 The bioaccessibility of PAHs can be quantified using different biological and chemical 295 approaches. For the purposes of this study, the mineralisation and extraction of ¹⁴C-296 phenanthrene by *M. gilvum* and HPCD were used, respectively. As a whole, these two 297 methods are considered acceptable methodologies to assess not only the fraction of the 298 hydrocarbon that is freely available to microorganisms, but also encompasses the fraction of 299 the contaminant that may become bioavailable, and therefore removed from the soil (Semple et al., 2004). The general absence of effects by organic acids on the mineralisation of ¹⁴C-300 301 phenanthrene reported in this study has also been observed by Cébron et al. (2011) and 302 Louvel et al. (2011), both of whom worked with root exudates containing mixtures of organic 303 acids. Despite this trend, both authors were able to observe an initial acceleration of the 304 mineralisation process (Cébron et al., 2011; Louvel et al., 2011), as was the case of citric acid (100 mmol l⁻¹) in this present study. Cébron *et al.* (2011) further discussed that this general 305 306 absence of effects might be the consequence of enhanced sorption of phenanthrene to SOM 307 and other soil inorganic fractions such as mineral clays caused by the organic acids, and that 308 ultimately reflected in a reduction in the availability of phenanthrene for microbial 309 degradation. Given the chemical characteristics of citric acid, other authors suggest that this 310 LOA is capable of forming more stable complexes with different compounds in soil (An et 311 al., 2011; Ling et al., 2015).

312 Similar trends were also observed when the bioaccessibility of ¹⁴C-phenanthrene was 313 measured through its HPCD extractability. Bioaccessibility was only significantly higher in 314 one of the treatments (500 mmol l⁻¹ citric and malic acid after 15 weeks soil-PAH contact 315 time). These findings contrast with that reported by other authors where PAH availability can 316 be significantly promoted by different LOAs assessed through *n*-butanol extractions (Ling *et* al., 2009; Sun et al., 2012, 2013; Kong et al., 2013; Gao et al., 2015b). Disagreement 317 318 between these two trends is suggested to be due to differences in the methodologies used for 319 this purpose, where *n*-butanol extracted PAH is not only the bioaccessible fraction of the 320 hydrocarbon, but also a portion of the non-bioaccessible PAH residues, as pointed by Ling et 321 al. (2009). Although *n*-butanol has been proposed to act as a predictor for the bioavailability 322 of PAHs in soil (Kelsey et al., 1997; Liste & Alexander, 2002), this extractant has also been 323 observed to exhibit greater extraction efficiencies when compared against HPCD 324 extractability (Swindell & Reid, 2006). This has become important when comparing these 325 two methods to the mineralisation of phenanthrene in soil (Reid et al., 2000; Rhodes et al., 326 2010), where close linear 1:1 relationships have been observed for the case of HPCD 327 extracted PAH. Furthermore, *n*-butanol has also been demonstrated to act as a more 328 exhaustive extractant than HPCD (Reid et al., 2000; Swindell & Reid, 2006), even extracting 329 similar quantities of PAHs than DCM, which is often use to determine total concentrations of 330 contaminants in soil (Reid et al., 2000).

331 The impact that organic acids from the rhizosphere might have towards the biodegradation of 332 PAHs remains a poorly explored area; however, this is one of the presumed mechanisms by 333 which plant-enhanced bioremediation is thought to occur (Pilon-Smits, 2005). Changes in the 334 physico-chemcial conditions in soil, such as pH, may play an important role in the microbial 335 degradation of PAHs (Kästner et al., 1998). Specifically, the acidic ranges of pH that the 336 presence of high concentrations of organic acids will produce, associated to the SOM-bound 337 PAHs that have been discussed, may be responsible to limit the microbial activity. This may 338 explain why, despite the fact that larger amounts of ¹⁴C-phenanthrene can be extracted with 339 high concentrations of organic acids, the PAHs are not being biodegraded by soil bacteria and 340 metabolised to CO₂.

341 **4.2 Impact of LOAs on the desorption of** ¹⁴**C-phenanthrene in soil**

The total desorbable fraction of ¹⁴C-phenanthrene did not decrease as a function of time over 342 343 the course of the incubation when soil was extracted with organic acids ($\geq 100 \text{ mmol } 1^{-1}$). 344 Despite the general acknowledgement of the negative correlation between the extractability 345 of organic contaminants and contact time (Hatzinger & Alexander, 1995; Semple et al., 346 2003), this behaviour was only observed in the control and the lowest tested concentration of 347 organic acids after 15 weeks of soil-PAH contact time. This trend suggests that high amounts of organic acids could potentially restrict the reduction of bioaccessibility of ¹⁴C-348 349 phenanthrene, therefore limiting the ageing process. Although not observed before, this 350 behaviour could be the reflection of a dual effect of organic acids on phenanthrene sorption 351 reported by Ouvrard et al. (2006) who described the impact of LOAs as a combined process 352 characterised by an initial short term enhanced sorption of phenanthrene by SOM, followed by an increased mass transfer of the hydrocarbon due to the destabilisation of this soil 353 354 fraction. Similarly, data from the present study showed a general reduction of the extractable 355 ¹⁴C-phenanthrene after a short period of ageing while organic acids were consistently 356 observed to promote a larger desorption after 15 weeks of soil-PAH contact time when 357 compared against the control. This increase in the extractability of PAHs by LOAs has also 358 been reported by other authors (Ling et al., 2009, 2015; Gao et al., 2010a; b, 2015b; Kong et al., 2013), but rarely considered the impact of soil-PAH ageing included in the present study. 359 360 Although not common, the reduction of ¹⁴C-phenanthrene desorption in the presence of 361 organic acids observed after a short soil-PAH contact time in the present study has been 362 reported before (Ouvrard et al., 2006; Zhu et al., 2009; Gao et al., 2015b). This initial 363 behaviour has been associated with the capacity of small amounts of oxalate, citrate and 364 malate to promote the sorption of anions to the soil (Jones & Brassington, 1998; Jones et al., 365 2003). In a similar way, phenanthrene has been hypothesised to be also sorbed through the

development of new sorption sites by these sorbed organic acids (Ouvrard *et al.*, 2006; Gao *et al.*, 2015a).

368 LOAs have been acknowledged to significantly influence the physical, chemical and 369 biological properties of soil (Jones & Darrah, 1994; Jones, 1998). As such, the main 370 mechanism behind the enhancement of the desorption of PAHs in soil impacted by organic 371 acids has been proposed to be the solubilisation of soil organic matter (SOM) with a 372 subsequent release SOM-associated hydrocarbons (Ouvrard et al., 2006; Agnello et al., 373 2014). This explanation is supported by findings from different authors who have reported 374 consistently higher amounts of dissolved organic matter and certain minerals when artificial 375 root exudates (Gao et al., 2010a) and single LOAs (Ling et al., 2009, 2015; Gao et al., 2010b, 376 2015a; Sun et al., 2012; Kong et al., 2013) were used to extract PAHs from contaminated 377 soil. In a similar way, previously immobilised aromatic compounds have been observed to be 378 released from soil to pore water after the introduction of organic acids solution (White et al., 379 2003; Gao et al., 2015b; Keiluweit et al., 2015).

380 4.2.1 Desorption kinetics

381 ¹⁴C-Phenanthrene desorption kinetics in the presence of the organic acids displayed a 3-382 compartment desorption behaviour. Although desorption of organic contaminants has been 383 widely observed to behave with an initial rapid desorption followed by a slower phase 384 (Cornelissen et al., 1998b), this rapid/slow/very slow release from the soil has also been 385 observed (Rhodes et al., 2010). Typically, studies investigating desorption kinetics are 386 performed using extractants that are known (or suspected) to correlate with the biodegradable 387 fraction of the contaminant in question (Cornelissen et al., 2001; Rhodes et al., 2010). 388 However, this current research study was focused on the assessment of the desorbing 389 potential that organic acids might be able to provide. Bearing this in mind, the proportion of

¹⁴C-phenanthrene desorbed at each of these phases should be considered as a measure of the
behaviour of this PAH under the influence of organic acids rather than an indication of its
bioaccessibility.

393 The different fractions described by the desorption kinetics can be interpreted as the 394 biodegradable (F_{rap}), and less accessible F_{slow} and/or $F_{very slow}$ fractions of the organic 395 contaminant (Cornelissen et al., 1998b; Rhodes et al., 2010). Results from this investigation 396 showed that the majority of the treatments had a tendency to enhance the very slowly 397 desorbing fraction ($F_{verv \ slow}$). These results could be interpreted as that the presence of organic 398 acids might be able to mobilise a significant proportion of the readily bioaccessible fraction of ¹⁴C-phenanthrene (F_{rap}) towards a less accessible form (F_{slow} and $F_{very slow}$), therefore 399 400 limiting the biological degradation of the contaminant or the rate at which this process takes 401 place (Pignatello & Xing, 1996; Clegg et al., 2014). Moreover, similar behaviour has been 402 observed to occur during the mineralisation of organic acids, where these compounds have 403 been observed to induce shifts of ¹⁴CO₂ production from a rapid to a slower phase (Oburger 404 et al., 2009).

405 **5** Conclusions

406 Organic acids found within the rhizosphere play an important role on the behaviour of 407 phenanthrene in soil. It was found that the total extractable fraction of ¹⁴C-phenanthrene can 408 be significantly enhanced by citric and malic acid. This effect is most likely to be observed at 409 a longer soil-PAH contact time, where organic acids showed to restrict the ageing effect. 410 Despite these enhancing effects, desorption kinetics indicated that the desorbed phenanthrene 411 was readily available given the behaviour as slow and very slow desorbing fractions. These trends were confirmed when accessibility and mineralisation of ¹⁴C-phenanthrene where 412 413 assessed. In this case, despite the enhancement of the total hydrocarbon extractable fractions

414 in the presence of citric and malic acid; there is no clear evidence suggesting that this condition can promote the microbial utilisation of ¹⁴C-phenanthrene. This study contributes to 415 416 the understanding of the role of root exudation within the rhizosphere towards the 417 bioaccessibility and biodegradation of hydrocarbons in contaminated soil. It is important to 418 note that organic acids may be able to remobilise contaminants, which were considered to be 419 non-bioaccessible. This may be important from a risk assessment perspective; however, the 420 concentrations of remobilised PAHs may be low and not represent a risk to environmental or 421 human health (Umeh et al., 2018).

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Table 1. Mineralisation kinetics of ¹⁴C-phenanthrene from soil affected by organic acids after 14 and 50 days ageing. Values of the lag phases (h), maximum rates (% d⁻¹) and total extents (%) represent the mean \pm standard error of the mean (n = 3). Different letters indicate significant differences between the treatments at each time point assessed by post hoc Tukey tests.

				14 day	vs ageing		
Treatment	Concentration (mmol l ⁻¹)	Lag	phase*	Faste	est rate	Total	extent
Control	0	18.288	± 0.358	^a 20.991	± 0.564	^{ab} 57.483	± 0.617
Citric acid	0.1	4.104	± 0.417	^b 29.520	± 2.316	^{ab} 65.325	± 3.590
	0.5	4.728	± 0.382	^{ab} 25.519	$\pm \ 1.439$	^{ab} 63.863	± 2.559
Malic acid	0.1	4.440	± 0.347	^{ab} 27.114	± 1.268	^{ab} 63.741	± 1.430
	0.5	4.320	± 0.364	^{ab} 27.902	± 1.050	^b 68.370	± 0.101
Oxalic acid	0.1	5.424	± 0.564	^{ab} 22.562	± 2.125	^a 54.085	± 3.386
	0.5	4.632	$\pm \ 0.397$	ab26.093	± 1.576	^{ab} 58.915	± 1.694
Succinic acid	0.1	4.656	± 0.260	^{ab} 25.806	± 0.664	^{ab} 63.543	± 2.409
	0.5	4.800	± 0.445	^{ab} 25.270	± 1.938	^{ab} 60.683	± 3.183
				50 day	vs ageing		
Treatment	Concentration (mmol 1 ⁻¹)	Lag	phase	Faste	est rate	Total	extent
Control	0	^a 84.350	± 2.400	^a 1.613	± 0.257	^a 11.581	± 1.321
Citric acid	0.1	^a 81.128	± 1.945	^a 1.607	± 0.128	^a 11.419	± 0.560
	0.5	^a 48.488	± 1.400	^a 1.651	± 0.020	a11.688	± 0.332
Malic acid	0.1	^a 69.532	± 2.020	a1.592	± 0.108	a12.535	± 0.474
	0.5	^a 85.519	± 2.319	a1.345	± 0.089	^a 10.590	± 0.667
Oxalic acid	0.1	^a 69.922	$\pm \ 0.894$	a1.553	± 0.026	a12.387	± 0.410
	0.5	a72.237	± 1.494	^a 1.670	± 0.128	a12.959	± 0.935
Succinic acid	0.1	^a 78.770	± 1.819	^a 1.661	± 0.140	a11.093	± 0.591
	0.5	^a 56.177	± 2.119	a1.959	± 0.154	a13.641	± 0.159

*Not normally distributed data analysed by Kruskal-Wallis non-parametric test (p = 0.077)

Table 2. HPCD extractable fraction of ¹⁴C-phenanthrene from soil after 1 and 15 weeks ageing following saturation with organic acids solution (100 % whc, 8 h). Values represent the mean \pm standard error of the mean (n = 5). Different letters indicate significant differences between the treatments at each time point (Tukey)

Treatment	Concentration (mmol l ⁻¹)	Bioac	cessible ¹⁴ C	C-pheannthre	ene (%)
		1 w	veek*	15 w	reeks
Control	0	^a 79.841	± 0.717	^{ab} 6.721	± 0.970
Citric acid	0.5	^a 78.964	± 2.070	^a 5.033	± 0.955
	100	^a 78.769	± 1.346	^{ab} 6.492	± 1.646
	250	^a 79.239	± 1.243	abc9.804	± 1.033
	500	^a 79.852	± 0.633	°14.929	± 0.582
	1000	^a 76.745	± 0.164	^{bc} 11.223	± 1.524
Malic acid	0.5	^a 81.138	± 0.611	^{ab} 7.992	± 0.690
	100	^a 80.517	± 0.366	abc 8.938	± 1.509
	250	^a 78.540	± 0.793	^{ab} 8.764	± 2.035
	500	^a 77.937	± 0.579	abc10.726	± 0.871
	1000	^a 76.737	± 1.003	^{ab} 8.594	± 1.284

Treatment	Concentration (mmol l ⁻¹)	Desorbed ¹⁴ C-	pheannthrene (%)
		1 week	15 weeks
Control	0	$^{de}27.980 \pm 1.636$	$^{b}18.958 \pm 0.931$
Citric acid	0.5	$^{ab}20.755 \pm 0.432$	$^{a}13.038 \pm 1.010$
	100	$^{a}17.221 \pm 0.211$	$^{cd}26.462 \pm 0.431$
	250	$^{ab}19.709 \pm 0.081$	$^{de}31.264 \pm 1.851$
	500	$^{cd}26.579 \pm 0.795$	${}^{\rm f}\!40.006 \pm 0.655$
	1000	$^{\rm f}39.274$ ± 1.921	$^{ m g}47.856 \pm 1.060$
Malic acid	0.5	$^{ab}20.068 \pm 0.376$	$^{a}12.507 \pm 0.176$
	100	$^{a}16.552 \pm 0.119$	$bc21.986 \pm 1.363$
	250	$^{ab}18.820 \pm 0.256$	$^{cd}25.923 \pm 0.983$
	500	$bc22.558 \pm 0.266$	$e32.955 \pm 1.134$
	1000	$e31.720 \pm 1.249$	$^{ef}36.184 \pm 2.054$

Table 3. Total ¹⁴C-phenanthrene desorbed from soil after 1 and 15 weeks ageing. Values represent the mean \pm standard error of the mean (n = 5). Different letters indicate significant differences between the treatments at each time point (Tukey).

Treatment	Concentration F_{rap} (%)		krap	(h ⁻¹)	Fslow	(%)	kslow	k_{slow} (h ⁻¹)	
1 week ageing									
Control	0	^{cd} 16.571	± 0.442	^a 0.128	± 0.004	^{cd} 83.428	± 0.442	^c 0.001	< 0.001
Citric acid	0.5	abc14.393	± 0.381	^{ab} 0.142	± 0.004	def85.606	± 0.381	^{ab} <0.001	< 0.001
	100	^a 12.313	± 0.160	^b 0.145	± 0.001	^f 87.687	± 0.160	^a <0.001	< 0.001
	250	^{abc} 14.423	± 0.108	^{ab} 0.134	± 0.001	def85.576	± 0.108	^a <0.001	< 0.001
	500	bcd19.090	± 0.706	^{ab} 0.142	± 0.006	°80.909	± 0.706	^b 0.001	< 0.001
	1000	e28.381	± 1.587	^{ab} 0.140	± 0.005	^a 71.618	± 1.587	^d 0.001	< 0.001
Malic acid	0.5	^{abc} 15.123	± 0.104	^b 0.139	± 0.002	def84.876	± 0.104	^a <0.001	< 0.001
	100	^a 11.901	± 0.143	^{ab} 0.142	± 0.002	f88.098	± 0.143	^a <0.001	< 0.001
	250	^{ab} 13.447	± 0.119	^{ab} 0.146	± 0.002	ef86.553	± 0.119	^a <0.001	< 0.001
	500	^d 16.122	± 0.272	^{ab} 0.141	± 0.003	^{cde} 83.877	± 0.272	^{ab} <0.001	< 0.001
	1000	^f 22.899	± 0.936	^{ab} 0.140	± 0.003	^b 77.100	± 0.936	^c 0.001	< 0.001
			15	weeks age	eing				
Control	0	^b 14.683	± 3.591	^{ab} 0.195	± 0.009	^f 85.317	± 3.591	^a 0.001	< 0.001
Citric acid	0.5	^a 6.789	± 0.247	^{ab} 0.170	± 0.012	^g 93.211	± 0.247	^a <0.001	< 0.001
	100	^{bc} 14.566	± 0.436	^{bc} 0.217	± 0.020	ef85.434	± 0.436	^b 0.001	< 0.001
	250	^b 19.534	± 0.493	^b 0.186	± 0.014	^{bc} 80.466	± 0.493	^{bc} 0.001	< 0.001
	500	^f 25.150	± 0.876	^{bc} 0.213	± 0.009	°74.850	± 0.876	^{cd} 0.001	< 0.001
	1000	^g 30.907	± 0.377	°0.262	± 0.006	^a 69.093	± 0.377	^d 0.002	< 0.001
Malic acid	0.5	^a 7.533	± 0.333	^{bc} 0.206	± 0.023	^g 92.467	± 0.333	^a <0.001	< 0.001
	100	^{bcd} 15.906	± 0.770	^a 0.108	± 0.021	def84.094	± 0.770	^a <0.001	< 0.001
	250	^{cde} 17.146	± 0.621	^{ab} 0.156	± 0.021	^{cde} 82.854	± 0.621	ab0.001	< 0.001
	500	e20.412	± 0.598	^{bc} 0.209	± 0.007	^a 79.588	± 0.598	^{bc} 0.001	< 0.001
	1000	^f 24.585	± 1.845	^{ab} 0.156	± 0.009	^b 75.415	± 1.845	^{bc} 0.001	< 0.001

Table 4. Desorbing fractions (F_{rap} and F_{slow}) and constant rates (k_{rap} and k_{slow}) calculated by a two-compartment model. Values represent the mean \pm standard error of the mean (n = 5). Different letters indicate significant differences between the treatments assessed by post hoc Tukey tests

Table 5. Desorbing fractions (F_{rap} , F_{slow} and $F_{very slow}$) and constant rates (k_{rap} , k_{slow} and $k_{very slow}$) calculated by a three-compartment model. Values represent the mean \pm standard error of the mean (n = 5). Different letters indicate significant differences between the treatments assessed by post hoc Tukey tests.

Treatment	Concentration (mmol 1 ⁻¹)	F_{rap} (%)	k _{rap}	(h ⁻¹)	F_{slow}	(%)	kslow	(h ⁻¹)	$F_{very \ slop}$	w(%)	k _{very slo}	h^{-1}
1 week ageing													
Control	0	de12.078	± 0.418	ab0.210	± 0.008	e15.801	± 0.850	^a 0.013	± 0.001	^b 72.122	± 1.251	<ª0.001	< 0.001
Citric acid	0.5	bcd10.164	± 0.564	^b 0.226	± 0.014	^{cd} 9.162	± 0.224	^{ab} 0.021	± 0.004	^d 80.674	± 0.708	<ab0.001< td=""><td>< 0.001</td></ab0.001<>	< 0.001
	100	^a 7.099	± 0.457	°0.319	± 0.037	^b 7.136	$\pm \ 0.418$	°0.042	± 0.003	efg85.765	± 0.224	<abc display="block"><abc display="block">abc 0.001</abc></abc>	< 0.001
	250	^b 8.984	± 0.423	^b 0.242	± 0.013	^{bc} 7.992	$\pm \ 0.219$	^{bc} 0.035	± 0.003	de83.025	± 0.254	<abc display="block"><abc display="block">abc 0.001</abc></abc>	< 0.001
	500	^{cde} 11.822	$\pm \ 0.656$	^{bc} 0.271	± 0.029	^d 10.589	$\pm \ 0.536$	°0.036	± 0.004	°77.589	± 0.563	<cde0.001< td=""><td>< 0.001</td></cde0.001<>	< 0.001
	1000	f19.220	± 0.344	^b 0.234	± 0.007	°16.563	± 0.614	^{abc} 0.027	± 0.005	^a 64.217	± 0.657	<cde0.001< td=""><td>< 0.001</td></cde0.001<>	< 0.001
Malic acid	0.5	^{cde} 11.379	± 0.524	^a 0.139	± 0.002	^a 3.686	$\pm \ 0.473$	^d 0.139	± 0.002	ef84.935	± 0.086	<de0.001< td=""><td>< 0.001</td></de0.001<>	< 0.001
	100	^{bc} 9.614	± 0.136	^a 0.142	± 0.002	^a 2.288	$\pm \ 0.008$	^d 0.142	± 0.002	^g 88.098	$\pm \ 0.143$	<cde0.001< td=""><td>< 0.001</td></cde0.001<>	< 0.001
	250	^{bcd} 11.074	± 0.115	^a 0.146	± 0.002	^a 2.373	$\pm \ 0.005$	^d 0.146	± 0.002	^{fg} 86.553	± 0.119	<de0.001< td=""><td>< 0.001</td></de0.001<>	< 0.001
	500	°13.632	± 0.260	^a 0.141	± 0.003	^a 2.491	± 0.012	^d 0.141	± 0.003	ef83.877	± 0.272	<e0.001< td=""><td>< 0.001</td></e0.001<>	< 0.001
	1000	^f 20.196	± 0.914	^a 0.140	± 0.003	^a 2.705	± 0.022	^d 0.140	± 0.003	°77.100	± 0.936	<f0.001< td=""><td>< 0.001</td></f0.001<>	< 0.001
					15 v	veeks ageii	ng						
Control	0	bcd13.115	± 0.712	^b 0.172	± 0.010	^d 83.646	$\pm \ 3.300$	<ª0.001	< 0.001	a3.239	± 3.234	<al> a0.001 </al>	< 0.001
Citric acid	0.5	^a 6.389	± 0.180	^{bc} 0.180	± 0.019	°14.050	± 4.178	^{bc} 0.005	± 0.001	^b 79.561	± 4.036	<ab0.001< td=""><td>< 0.001</td></ab0.001<>	< 0.001
	100	^{bc} 12.466	± 0.512	^{cd} 0.298	± 0.023	°16.386	± 0.491	^{cd} 0.010	± 0.001	^b 71.149	± 0.609	< ^b 0.001	< 0.001
	250	^{bc} 11.753	± 1.008	ef0.511	± 0.053	°18.569	± 1.781	de0.027	± 0.007	^b 69.678	± 2.345	<ab0.001< td=""><td>< 0.001</td></ab0.001<>	< 0.001
	500	ef17.595	± 1.768	de0.521	± 0.126	°19.782	± 1.110	^d 0.020	± 0.007	^b 62.623	± 2.536	<ab0.001< td=""><td>< 0.001</td></ab0.001<>	< 0.001
	1000	^g 25.120	± 1.153	de0.398	± 0.046	°15.909	± 2.951	de0.026	± 0.005	^b 58.971	± 2.787	^{ab} 0.001	< 0.001
Malic acid	0.5	^a 5.504	± 0.526	^{bc} 0.190	± 0.021	^b 1.994	± 0.305	f0.190	± 0.021	^b 92.502	± 0.324	<al> a0.001 </al>	< 0.001
	100	^{cde} 14.762	$\pm \ 0.759$	^a 0.090	± 0.012	^a 1.057	± 0.006	f0.090	± 0.012	^b 84.181	$\pm \ 0.765$	<ª0.001	< 0.001
	250	def16.916	± 0.492	^b 0.150	± 0.010	^d 58.727	± 1.719	^a 0.001	± 0.000	^b 24.357	± 1.776	<ª0.001	< 0.001
	500	^f 20.465	± 0.659	^{bc} 0.210	± 0.007	^d 58.849	± 1.998	^{ab} 0.001	< 0.001	^{ab} 20.686	± 1.732	^a 0.001	< 0.001
	1000	^{ab} 9.066	± 0.771	f0.830	± 0.032	°18.118	± 1.007	ef0.061	± 0.002	^{ef} 72.816	± 1.353	^a 0.001	< 0.001

Supplementary information

The effect of organic acids on the fate of ¹⁴C-phenanthrene in contaminated soil

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Supplementary information

					Two-c	ompartment	fitting						
Treatment	Concentration (mmol l ⁻¹)	Frap	(%)	<i>k</i> _{rap}	(h ⁻¹)	Fslow	w (%)	kslov	(h^{-1})				
Control	0	^{ab} 40.939	± 3.861	^a 0.090	± 0.019	^{ab} 59.061	± 3.861	^a 0.001	< 0.001				
Citric acid	0.1	^a 25.905	± 6.193	a0.133	± 0.020	^b 74.095	± 6.193	^a 0.001	< 0.001				
	0.5	^{ab} 44.164	± 7.875	^a 0.119	± 0.010	^{ab} 55.836	± 7.875	^a 0.001	< 0.001				
Malic acid	0.1	^a 25.356	± 3.052	^a 0.179	± 0.007	^b 74.644	± 3.052	^a 0.001	< 0.001				
	0.5	^b 49.763	± 3.847	^a 0.110	± 0.015	^a 50.237	± 3.847	^a 0.001	< 0.001				
Oxalic acid	0.1	^{ab} 37.096	± 5.009	^a 0.107	± 0.017	^{ab} 62.904	± 5.009	^a 0.001	< 0.001				
	0.5	^{ab} 34.857	± 1.603	^a 0.151	± 0.027	^{ab} 65.143	± 1.603	^a 0.001	± 0.001				
Succinic acid	0.1	^{ab} 30.451	± 2.869	^a 0.115	± 0.036	^{ab} 69.549	± 2.869	^a 0.000	< 0.001				
	0.5	^{ab} 28.999	± 1.983	^a 0.123	± 0.007	^{ab} 71.001	± 1.983	^a 0.001	< 0.001				
					Three-c	compartmen	t fitting						
Treatment	Concentration (mmol l ⁻¹)	Frap	(%)	<i>k</i> _{rap}	(h ⁻¹)	Fslow	w (%)	kslov	v (h ⁻¹)	Fvery s	_{low} (%)	kvery slow	(h ⁻¹)
Control	0	^{ab} 36.112	± 5.067	^{ab} 0.138	± 0.023	^a 56.511	± 2.855	^{ab} 0.003	< 0.001	a7.377	± 2.458	^a 0.003	< 0.001
Citric acid	0.1	^a 22.920	± 6.498	ab0.168	± 0.035	°51.399	± 18.98	ab0.003	± 0.001	^a 25.682	± 13.80	^a 0.002	± 0.001
	0.5	^b 56.929	± 9.031	^a 0.130	± 0.083	^a 30.697	± 10.45	^{ab} 0.007	$\pm \ 0.005$	^a 12.374	± 12.37	^a 0.001	± 0.001
Malic acid	0.1	^a 23.902	± 2.529	^{ab} 0.289	± 0.045	^a 61.384	± 9.804	ab0.002	± 0.001	^a 14.714	± 8.010	^a 0.001	± 0.001
	0.5	^{ab} 43.770	± 2.514	^{ab} 0.193	± 0.035	ª46.991	± 8.865	^{ab} 0.005	± 0.003	^a 9.240	± 6.792	^a 0.002	± 0.001

Table SI 1. Desorption kinetics of ¹⁴C-phenanthrene from mildly aged soil (50 d). Values represent the mean \pm standard error of the mean (n = 3). Different letters indicate significant differences between the treatments assessed by post hoc Tukey tests

Oxalic acid	0.1	^{ab} 31.645	± 7.967	^{ab} 0.321	± 0.179	^a 44.376	± 9.240	^{ab} 0.007	$\pm \ 0.005$	^a 23.979	± 15.78	^a 0.001	< 0.001
	0.5	^{ab} 29.134	± 1.809	^{ab} 0.375	± 0.082	^a 38.480	± 9.898	^{ab} 0.007	± 0.003	^a 32.386	± 10.93	^a 0.001	± 0.001
Succinic acid	0.1	^{ab} 29.880	± 2.748	^a 0.147	± 0.058	^a 64.678	± 5.535	^a 0.001	< 0.001	^a 5.443	± 4.019	^a 0.001	± 0.001
	0.5	^a 19.530	± 0.664	^b 0.577	± 0.106	^a 24.270	± 5.003	^b 0.016	± 0.006	^a 56.200	± 5.500	^a <0.001	< 0.001

Contact time (h)	Treatment	Extracted (%) ¹	Mineralised $(\%)^2$
1	Control	7.269 ± 0.993	1.454 ± 0.199
	0.5 mmol l ⁻¹	5.777 ± 0.767	1.155 ± 0.153
	500 mmol 1 ⁻¹	7.854 ± 0.257	1.571 ± 0.051
3	Control	6.062 ± 0.943	0.505 ± 0.236
	0.5 mmol 1 ⁻¹	6.101 ± 0.358	0.508 ± 0.089
	500 mmol 1 ⁻¹	7.268 ± 1.315	0.605 ± 0.329
6	Control	5.843 ± 1.035	0.243 ± 0.259
	0.5 mmol 1 ⁻¹	6.264 ± 0.954	0.261 ± 0.238
	500 mmol 1 ⁻¹	5.528 ± 0.818	0.230 ± 0.205
8	Control	8.146 ± 1.876	0.254 ± 0.469
	0.5 mmol 1 ⁻¹	5.978 ± 1.104	0.186 ± 0.276
	500 mmol 1 ⁻¹	8.495 ± 1.182	0.265 ± 0.296
24	Control	5.250 ± 3.213	0.054 ± 0.803
	0.5 mmol 1 ⁻¹	3.805 ± 1.050	0.039 ± 0.263
	500 mmol 1 ⁻¹	6.282 ± 1.327	0.065 ± 0.332
48	Control	1.102 ± 0.108	0.091 ± 0.432
	0.5 mmol 1 ⁻¹	0.929 ± 0.169	0.077 ± 0.676
	500 mmol l ⁻¹	1.257 ± 0.157	0.104 ± 0.628

Table SI 2. Proportion of ¹⁴C-phenanthrene (1) extracted with 50 mM HPCD solution, (2) mineralisation rate (%, h⁻¹) within the assessed contact time. Values represent the mean \pm standard error of the mean (n = 3)

Treatment	Concentration (mmol l ⁻¹)	Sum of squared differen 2 compartment fitting	ce Sum of squared difference 3-compartment fitting				
1 week ageing							
Control	0	9.05E-04 ± 1.0E-0	04 2.24E-04 $\pm 2.8E-05$				
Citric acid	0.5	4.79E-04 ± 3.6E-0	05 $2.53E-04 \pm 7.8E-05$				
	100	3.97E-04 ± 2.6E-0	$1.39E-04 \pm 1.3E-05$				
	250	$4.19E-04 \pm 2.4E-0$	$1.28E-04 \pm 1.1E-05$				
	500	8.97E-04 ± 1.0E-0	04 $2.73E-04 \pm 5.1E-05$				
	1000	2.40E-03 ± 2.8E-0	$04 7.09E-04 \pm 9.4E-05$				
Malic acid	0.5	$5.44E-04 \pm 4.0E-0$	$05 \qquad 5.44 \text{E-}04 \pm 4.0 \text{E-}05$				
	100	3.13E-04 ± 1.3E-0	$3.13E-04 \pm 1.3E-05$				
	250	4.98E-04 ± 2.0E-0	$05 \qquad 4.98E-04 \pm 2.0E-05$				
	500	6.71E-04 ± 1.4E-0	$05 \qquad 6.71E-04 \pm 1.4E-05$				
	1000	1.49E-03 ± 1.7E-0	04 $1.49E-03 \pm 1.7E-04$				
		15 weeks ageing					
Control	0	4.50E-04 ± 6.0E-0	$05 \qquad 4.60E-04 \pm \ 6.0E-05$				
Citric acid	0.5	9.59E-05 ± 4.6E-0	$05 \qquad 6.00E-05 \pm 4.0E-05$				
	100	1.17E-03 ± 2.0E-0	04 $2.20E-04 \pm 9.3E-05$				
	250	2.50E-03 ± 3.9E-0	$1.48E-04 \pm 5.0E-05$				
	500	3.59E-03 ± 4.5E-0	04 9.20E-04 $\pm 1.9E-04$				
	1000	2.99E-03 ± 7.8E-0	$04 \qquad 3.70E-04 \pm 4.8E-05$				
Malic acid	0.5	3.92E-04 ± 6.4E-0	$05 \qquad 4.20E-04 \pm \ 6.6E-05$				
	100	4.63E-04 ± 1.3E-0	04 $4.63E-04 \pm 1.3E-04$				
	250	1.98E-03 ± 1.2E-0	04 $1.28E-03 \pm 4.1E-04$				
	500	1.75E-03 ± 1.0E-0	04 $1.58E-03 \pm 2.3E-04$				
	1000	3.70E-03 ± 4.7E-0	04 $2.80E-04 \pm 6.8E-05$				

Table SI 3. Sums of squared deviations of desorbed ¹⁴C-phenanthrene fitted to a two- and three-compartment model. Values represent the mean \pm standard error of the mean (n = 5)



Figure SI 4. Rapid desorbing fractions (F_{rap}) of ¹⁴C-phenanthrene from soil aged for 1 (A) and 15 (B) weeks extracted with citric (•) and malic (\circ) acid. Values were obtained using a three-compartment model fitting. Error bars represent the standard error of the mean (n = 5).



Figure SI 5. Rate constants of the rapid desorbing fractions (k_{rap}) of ¹⁴C-phenanthrene from soil aged for 1 (A) and 15 (B) weeks extracted with citric (•) and malic (\circ) acid. Values were obtained using a three-compartment model fitting. Error bars represent the standard error of the mean (n = 5).



Figure SI 6. Slow desorbing fractions (F_{slow}) of ¹⁴C-phenanthrene from soil aged for 1 (A) and 15 (B) weeks extracted with citric (•) and malic (\circ) acid. Values were obtained using a three-compartment model fitting. Error bars represent the standard error of the mean (n = 5).



Figure SI 7. Rate constants of the slow desorbing fractions (k_{slow}) of ¹⁴C-phenanthrene from soil aged for 1 (A) and 15 (B) weeks extracted with citric (•) and malic (\circ) acid. Values were obtained using a three-compartment model fitting. Error bars represent the standard error of the mean (n = 5).



Figure SI 8. Very slow desorbing fractions ($F_{very slow}$) of ¹⁴C-phenanthrene from soil aged for 1 (A) and 15 (B) weeks extracted with citric (•) and malic (•) acid. Values were obtained using a three-compartment model fitting. Error bars represent the standard error of the mean (n = 5).



Figure SI 9. Rate constants of the very slow desorbing fractions ($k_{very slow}$) of ¹⁴C-phenanthrene from soil aged for 1 (A) and 15 (B) weeks extracted with citric (•) and malic (•) acid. Values were obtained using a three-compartment model fitting. Error bars represent the standard error of the mean (n = 5).