Biochar amendment increases bacterial diversity and vegetation cover in trace element-polluted soils: A long-term field experiment

Paloma Campos, Ana Z. Miller, Sergio A. Prats, Heike Knicker, Nikolas Hagemann, José M. De la Rosa

PII: S0038-0717(20)30310-2

DOI: https://doi.org/10.1016/j.soilbio.2020.108014

Reference: SBB 108014

To appear in: Soil Biology and Biochemistry

Received Date: 18 June 2020

Revised Date: 5 September 2020

Accepted Date: 13 September 2020

Please cite this article as: Campos, P., Miller, A.Z., Prats, S.A., Knicker, H., Hagemann, N., De la Rosa, José.M., Biochar amendment increases bacterial diversity and vegetation cover in trace element-polluted soils: A long-term field experiment, *Soil Biology and Biochemistry* (2020), doi: https://doi.org/10.1016/j.soilbio.2020.108014.

This is a PDF file of an article that has undergone enhancements after acceptance, such as the addition of a cover page and metadata, and formatting for readability, but it is not yet the definitive version of record. This version will undergo additional copyediting, typesetting and review before it is published in its final form, but we are providing this version to give early visibility of the article. Please note that, during the production process, errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

© 2020 Published by Elsevier Ltd.



# Biochar amendment increases bacterial diversity and vegetation cover in trace element-polluted soils: a long-term field experiment

3

Authors: Paloma Campos<sup>1</sup>, Ana Z. Miller<sup>2\*</sup>, Sergio A. Prats<sup>3</sup>, Heike Knicker<sup>1</sup>, Nikolas
 Hagemann<sup>4,5</sup>, José M. De la Rosa<sup>1</sup>

6

Affiliations: 1) Instituto de Recursos Naturales y Agrobiología de Sevilla (IRNAS-CSIC), Av. Reina Mercedes 10, 41012, Seville, Spain; 2) Laboratorio Hercules, University of Évora, Largo Marquês de Marialva 8, 7000-809, Évora, Portugal; 3)
Centre for Environmental and Marine Studies (CESAM), Department of Environment and Planning, University of Aveiro, Aveiro, Portugal; 4) Agroscope Zurich, Reckenholzstr. 191, Zurich, Switzerland; 5) Ithaka Institute, Arbaz, Switzerland

13

14 \*corresponding author: anamiller@uevora.pt

15

# 16 Abstract

Application of biochar has been widely suggested as a remediation tool for trace 17 element-polluted soils, but the impact of biochar on microbial communities and on 18 native plants remain largely unknown. To overcome this knowledge gap, biochar 19 produced from rice husk and olive pit were applied at a rate of 8 t ha<sup>-1</sup> into a soil with 20 two contrasting levels of trace elements (high and moderate) to study their effects on 21 22 soil microbial community composition, vegetation cover and soil properties after 1, 6, 23 12 and 20 months under field conditions. Differences in bacterial community 24 composition were studied using the Illumina Miseg technology of the 16S rRNA gene.

25 Although variations in soil properties and ecological function were seasonal and soil-26 type dependent, biochar application enhanced soil properties and vegetation cover in the moderately polluted soil (MPS), and increased microbial diversity as well as 27 vegetation cover in the highly polluted soil (HPS). Enzymatic activities and soil 28 respiration rates were not modified with the application of biochar, but increased total 29 carbon content of soils. The application of biochar from crop residues to trace-element 30 contaminated soils provided environmental benefits, including plant diversity and 31 32 growth, as well as the increase of bacterial diversity and carbon sequestration.

33

Keywords: Soil remediation; pyrogenic carbon; heavy metals; soil microbial
 community; plant diversity

# 36 **1. Introduction**

Trace element-polluted soils is a worldwide concern comprising 37 % of the degraded 37 soils in the European Union (EEA, 2007). Ex-situ decontamination of polluted soils is 38 39 generally unfeasible due to land size and soil contamination levels, which are difficult to effectively and economically reduce with conventional soil remediation procedures 40 (Tack et al., 2018). Biochar, the C-rich porous solid residue produced by the thermal 41 42 conversion of biomass under the partial or total absence of oxygen (pyrolysis, e.g. 43 Hagemann et al. 2018), has the ability to immobilize trace elements and increase the 44 pH of acidic soils reducing trace element mobility and bioavailability. Karer et al. (2015) 45 reported a decrease in NH<sub>4</sub>NO<sub>3</sub>-extractable fraction of Pb, Zn and Cd with biochar 46 amendment, but an increase of Cu. Beesley et al. (2010) also reported an immobilization of Cd and Zn and a mobilization of Cu after biochar application. 47 Oustriere et al. (2017) showed long-term Cu stabilization due to biochar addition into a 48 contaminated soil, whereas Uchimiya et al. (2012) reported Cu immobilization but 49 mobilization of Sb. These discrepancies are probably due to the complexity of 50 immobilization mechanisms and different biochar compositions and properties, but also 51 due to differences in the soil properties, e.g. in pH. In fact, previous studies already 52 demonstrated that the efficacy of biochar as a soil amendment greatly depends on its 53 pyrolysis conditions and feedstock (Campos et al., 2020; De la Rosa et al., 2014). For 54 instance, Kammann et al. (2012) showed a significant increase in biomass yield after 55 applying 50 Mg ha<sup>-1</sup> of peanut hull biochar to a Luvisol. Gascó et al. (2016) reported 56 that β-glucosidase, phosphomonoesterase and phosphodiesterase activities were 57 58 lower when a sandy loam soil was incubated with 8 % (w/w) of biochar produced from 59 pig manure at 500 °C whereas the biochar produced at 300 °C increased 60 dehydrogenase activity. The study of Shen et al. (2019) demonstrated that biochar 61 produced at 500 °C was more effective in the removal of lead from soil solution than the biochar produced at 300 °C. Generally, biochar produced at 500 °C has high pH 62 and water holding capacity, and high degree of aromatization (Campos et al., 2020). 63

The effects of biochar on the physical and chemical properties of agricultural soils have 64 been profoundly studied and during the last years special attention has been paid to 65 the study of biochar as soil amendment for the retention of contaminants (Uchimiya et 66 67 al., 2011; Kumar et al., 2018; De la Rosa et al., 2019). Within this context, effects of biochar addition on soil microbiota, which play a vital role in soil ecosystem stability, 68 soil quality and soil nutrient cycle (Lehmann et al., 2011), are highly relevant. Li et al. 69 (2019) reported a decrease in Actinobacteria with biochar application into purple soil, 70 71 whereas Ali et al. (2019) reported an increase for pesticide-contaminated soil. Most of 72 these studies on soil microbial diversity in polluted soils after biochar application are

pot-based experiments (Jiang et al., 2017; Han et al., 2017), which are likely to be
important in the quest to constrain the numerous influencing factors, but are less
realistic than field studies.

76 An aspect also worth further researching is the biodegradability of biochar in soils. Biochar has traditionally been considered a material of high chemical and biochemical 77 stability, which predominantly contains C in the form of condensed aromatic rings. This 78 79 fraction of C is hardly decomposed by soil biota due to its recalcitrant nature (Kuzyakov 80 et al., 2009, 2014). Nevertheless, recent studies indicate a much lower biochemical stability (Knicker et al., 2013, De la Rosa et al., 2018). Thus, the effects of biochar 81 82 application on soil CO<sub>2</sub> emissions are often ambiguous and previous studies reported 83 increases, decreases or no changes (Bamminger et al., 2014; Kolb et al., 2009; Paz-Ferreiro et al., 2012). Hence and considering that changes on soil properties promoted 84 by biochar application may affect soil microbial communities (Luo et al., 2013; Su et al., 85 2015; Xu et al., 2016) and soil CO<sub>2</sub> emissions, their assessment deserve further 86 attention. 87

The application of low degradability of biochar in trace element-polluted soils would allow an effective *in situ* remediation by enhancing soil quality and improving its capability to perform soil ecological functions. To test this hypothesis, we applied rice husk (RH) and olive pit (OP) biochar into two trace element-polluted acidic soils under field conditions to study their effects on soil physicochemical properties, soil CO<sub>2</sub> emissions and enzymatic activities, as well as soil microbial community composition and vegetation cover after 1, 6, 12 and 20 months of biochar application.

95

# 96 2. Materials and methods

# 97 2.1. Biochar samples

98 Rice husk (RH) and olive pit (OP) were used as feedstock to produce biochar due to 99 their great abundance in Mediterranean countries. RH is a siliceous-rich raw material 100 with relatively low C content, while OP is a hard-wood biomass, mainly composed of 101 cellulose and lignin. The company Orivarzea S.A. (Portugal) provided the RH biomass, 102 whereas OP was provided by *Cooperativa Nuestra Señora de los Ángeles* (Montellano, 103 Spain).

Prior to pyrolysis process, feedstock was dried at 40 °C during 48 h, homogenized and
stored in sealed plastic bags at 4 °C. The RH and OP biochar (RHB and OPB) were
produced in a continuously feed pyrolysis reactor with a screw conveyor (PYREKA,
Pyreg GmbH, Dörth, Germany, cf. Hagemann et al. 2020) under N<sub>2</sub> flux at Agroscope
Zurich (Switzerland). The pyrolysis temperature was 500 °C and the residence time

109 was 12 minutes. Biochar was not quenched with water and stored in plastic bags.110 Biochar characteristics are shown in Table 1.

111

# 112 **2.2. Area of study and experimental design**

The field experiment was conducted at 'Las Doblas' site (37° 23' 7.152"N, 6° 13' 113 43.175") over a period of 20 months. This place is located close to the Guadiamar river, 114 10 km from the former mine "Los Frailes" close to Aznalcollar, Southern Spain. On the 115 25<sup>th</sup> of April 1998, after a major mining accident, a huge amount of toxic sludge spilled 116 117 out from a tailing reservoir of this large open-pit mine, causing high levels of heavy 118 metals to leach into the soil and groundwater. Figure 1 shows the location of the field 119 experiment. The area belongs to a typical dry Mediterranean climate region, with hot and extended summers, soft winters and a very pronounced variation in the 120 precipitation rate (AEMET, 2018). 121

The sandy loam soil of the area is classified as Fluvisol (IUSS Working Group WRB, 122 123 2015). In this study, two nearby sites were selected according to their contamination 124 level and acidity, comprising a highly polluted soil (HPS) and a moderately polluted soil (MPS). HPS is a bare soil with high acidity and concentrations of heavy metals, as 125 previously described in Cabrera et al. (1999) and Martín-Peinado et al. (2015). These 126 bare spots account over 200 ha of lands affected by the accumulation of residual toxic 127 sludge of the spill. In contrast, MPS areas were subjected to a decontamination 128 programme by the Andalusian regional government which included the removal of the 129 toxic sludge (Arenas et al., 2008). Despite the decontamination efforts, MPS also 130 131 shows relatively high concentration of Ba, Cu, Fe, Pb and Zn (Campos and De la Rosa, 132 2020). Soil pH, total carbon (TC), total nitrogen (TN) and total hydrogen (TH) contents 133 of HPS and MPS are shown in Table 1.

In April 2018, 12 plots of 1 m × 1 m each were randomly established in HPS and MPS sites (6 plots per site). RHB and OPB were applied as produced and mixed into the first 10 cm of soil at a dose of 8 t ha<sup>-1</sup> (Plots ID: RHB\_HPS, OPB\_HPS, RHB\_MPS and OPB\_MPS). In addition, control plots without amendment were stablished for both areas (C\_MPS and C\_HPS) but received the same mechanical treatment. For all the plots, ground vegetation (shrub and grass) was manually removed; the soil was then homogenised using a manual rake.

Four sampling campaigns were performed after 1, 6, 12 and 20 months of biochar incorporation into soils (hereafter:  $t_1$ ,  $t_6$ ,  $t_{12}$  and  $t_{20}$ , respectively). For each plot, five samples of soil were taken randomly from the first 10 cm depth to create a composite sample per plot. An aliquot of the composite sample was immediately used for enzymatic analyses, other aliquot was stored in sterile Whirl-pak® bags at -80 °C for DNA-based analysis and the remaining material was dried at 40 °C during 48 h, sieved
(<2 mm) and stored in sealed bags at 4 °C.</li>

148

# 149 **2.3. Chemical and biochemical analysis**

The pH was measured in triplicates in the supernatant of a 1:5 (w/v) soil:0.01 M CaCl<sub>2</sub> solution ratio mixture after 30 minutes shaking and 30 minutes resting, using a pH meter (CRISON pH Basic 20).

The soil moisture (%) was determined on the dry weight basis: 20 g of moist soil was weighed, dried at 40 °C during 24h and re-weighed. Total soil moisture (%) was determined for soil samples dried at 105 °C for 24 h.

Total C (TC) was obtained by dry combustion (1050 °C) using an elemental analyzer
(TRUSPEC CHNS MICRO, LECO, St. Joseph, MI, USA).

The water holding capacity (WHC) was measured following the procedure and formula
described in Campos et al. (2020). The WHC is expressed as the percentage relatively
to the total dry weight of the sample:

161

# $WHC (\%) = \frac{Water retained weight}{Initial weight of the dry sample} \cdot 100 (Eq. 1)$

162

For elucidating microbial oxidative activities in soil, dehydrogenase activity was determined according to the method of Trevors (1986). Briefly, soil samples were incubated for 20 h with 1 M TRIS–HCl buffer (pH 7.5) and 2(p-iodophenyl)-3-(pnitrophenyl) 5-phenyl tetrazolium chloride (INT), that was used as the electron acceptor. After adding methanol, the iodonitrotetrazolium formazan (INTF) produced was measured spectrophotometrically at 490 nm.

In addition, soil β-glucosidase activity was measured according to the method of Tabatabai (1982). Briefly, 1 g of soil was incubated 1 h at 37 °C with p-nitrophenyl-β-Dglucopyranoside. After addition of CaCl<sub>2</sub>, the p-nitrophenol was extracted by filtration and measured using a spectrophotometer (Jenway, model 6315, UK) at 400 nm. βglucosidase and dehydrogenase activities were measured in both unamended and biochar-amended soils at t<sub>1</sub>, t<sub>6</sub>, t<sub>12</sub> and t<sub>20</sub>.

- 175 All chemical and biochemical analyses of the samples were performed in triplicate.
- 176

# 177 2.4. Measurement of soil CO<sub>2</sub> efflux (Soil respiration)

178 Soil respiration (carbon decomposition by microorganisms and ground root respiration)

179 was determined by measuring the CO<sub>2</sub> effluxes and expressed as  $\mu$ mol CO<sub>2</sub> m<sup>-2</sup> s<sup>-1</sup>.

180 For each plot, 3 PVC collars (10 cm diameter and 5 cm high) were installed 3 cm into

the soil and measurements were conducted in triplicate using the soil  $CO_2$  flux chamber LI-COR 6400-09 (LI-COR, Nebraska, USA) at  $t_1$ ,  $t_6$ ,  $t_{12}$  and  $t_{20}$ . Soil temperature was monitored using a thermocouple probe (Li6000-09 TC, LiCor Inc) inserted to a depth of 5 cm near the soil  $CO_2$  flux chamber.

185

# 186 2.5. Effects on vegetation

187 The vegetation species were carefully identified and the number of individuals per plot 188 were accounted at  $t_{12}$ . Subsequently, the total plant biomass was determined by 189 harvesting and measuring the fresh weight per plot.

In order to determine the percentage of vegetation cover at time  $t_{20}$ , high resolution photographs were taken for each plot using a digital camera (Canon Inc., Canon 7D, Japan) installed on a tripod at a height of 1.5 m. Digital images were then analysed using the open source image-processing software Image J. The area covered by green plants was selected by adjusting the hue levels in the colour threshold tool. The percentage of vegetation cover was determined by using the following equation 2:

196

197

% vegetation cover =  $\frac{green area}{total area} \cdot 100$  (Eq. 2)

198

# 199 2.6. Soil DNA isolation and sequencing

200 Total DNA was extracted from soil samples using the DNeasy PowerSoil DNA isolation 201 kit (Qiagen, Hilden, Germany), according to the manufacturer's instructions. DNA 202 quality and quantity were tested. As a standard procedure, 1.5% agarose gel electrophoresis was performed with 1 µL of qDNA of each sample to test the integrity 203 204 and purity. DNA concentrations were verified using a Qubit 2.0 fluorometer (Life Technologies, Carlsbad, CA, USA). Qubit broad-range reagent was used for 205 determining the DNA concentration of MPS samples, whereas Qubit high-sensitivity 206 207 reagent was needed for HPS samples due to their low amount of DNA.

Library construction was performed according to the Illumina 16S Metagenomic Sequencing Library preparation protocol by STAB VIDA Sequencing Services (Portugal). MiSeq Reagent Kit v3 in the Illumina MiSeq platform was used for sequencing new generated DNA fragments. 300 bp paired-end sequencing reads were used.

The microbial community composition and diversity (alpha and beta-diversity) were determined after bioinformatics processing of the 16S rRNA gene sequences. Sequence quality control was performed using QIIME2 v2019.1.0 (Bolyen et al., 2019). The reads were denoised using the DADA2 plugin, organized in operational taxonomic

units (OTUs) and classified by taxon using the SILVA database, with a clustering threshold of 97% similarity. OTUs were considered as significant only when they contained at least 10 sequence reads. This procedure results in an abundance table with taxonomy information, which were further analysed and visualized using the online web-tool Calypso (Zakrzewski et al., 2017).

The raw reads were deposited in NCBI Sequence Read Archive (SRA) database
(https://submit.ncbi.nlm.nih.gov/about/sra/) under the accession number
PRJNA637319.

225

# 226 **2.7. Data analysis**

Data of soil and biochar characteristics are expressed as mean ± standard error (SE)
of triplicate measurements. Data of the samplings are expressed as mean ± SE of the
five composite samples per treatment. Number of species and number of individuals
are expressed as median.

231 Shapiro-Wilk test was used to verify normality and Levene test was used to test 232 homoscedasticity of the data. Normal distributed response variables were analysed by one-way ANOVA followed by the Tukey's Honestly Significant Difference test. The level 233 of significance used was 0.05. When response variables were non-normal, Kruskal 234 Wallis followed by Mann Whitney U tests were conducted. Pearson correlation (p < p235 0.05) was conducted to examine relationships between soil properties, soil microbial 236 community and vegetation data. Statistical analyses were carried out using IBM SPSS 237 Statistics 26.0 (SPSS, Chicago, USA) for Windows. 238

239

# 240 **3. Results**

# 241 3.1. Soil pH and moisture

Soil pH of HPS samples were more acidic than MPS (3.57–3.77 and 4.18–5.11,
respectively) (Table 2). Biochar addition did not significantly enhance soil pH of HPS.
For MPS, biochar amendment clearly increased soil pH, but this increase was
mitigated over the time span of the experiment.

Soil moisture determined by drying at 40 °C was greater for MPS than HPS samples with comparable treatments (Table S1). Biochar addition augmented soil moisture (40 °C) at  $t_6$  and to a lesser extent at  $t_{20}$ . As expected, seasonal changes modified the soil moisture at 40 °C and 105 °C, with a significant drop at  $t_{12}$ , followed by a considerable increase during the autumn ( $t_{20}$ ). MPS showed a greater WHC (%) than HPS, and were affected by seasonal changes, showing in general a similar trend as the soil moisture content (Table 2).

253

# **3.2. Total carbon content and soil respiration**

Biochar addition caused a non-significant increase (p > 0.05) of the C content of the amended soils when compared with control plots (Table 2). At t<sub>12</sub>, the C content significantly increased due to OPB amendment in HPS samples (9.6 to 16.6 g kg<sup>-1</sup>).

The CO<sub>2</sub> emission rates were always higher in MPS than in HPS. The latter showed very low respiration rates. Significant differences were not observed with biochar application into the soils, nor differences between both biochar samples for the same soil type (Table 2).

262

# 263 3.3. Soil enzymatic activities

β-glucosidase activity at  $t_1$  of MPS was greater than HPS (0.69-1.45 *vs* 0.17-0.39 µmol PNF g<sup>-1</sup> h<sup>-1</sup>) (Fig. 2). At this time, the HPS control soil showed a greater β-glucosidase activity than biochar amended soils. Nevertheless, this difference disappeared at  $t_6$  and  $t_{12}$ . This enzymatic activity was greater for both amended treatments than for control in MPS at  $t_1$ . Nevertheless, at  $t_6$  only OPB addition maintained a greater β-glucosidase activity than control soils and at  $t_{20}$  no significant differences were observed.

270 MPS plots showed greater dehydrogenase activity than in HPS in all the cases. 271 Similarly to the trend observed for  $\beta$ -glucosidase activity, dehydrogenase activity 272 showed seasonal changes and during the first 6 months of the experiment MPS soils 273 amended with biochar showed lower values than control soils.

274

# **3.4. Effects on vegetation development**

276 A total of 14 different species were observed in MPS plots which were not found in 277 HPS plots (Echium gaditanum, Lotus parviflorus, Trifolium arvense, Ornithopus 278 compresus, Anagallis arvensis, Bartsia trixago, Trifolium sp., Vulpia ciliata, Trifolium 279 vesiculosum, Trifolium striatum, Hypochaeris glabra, Astragalus pelecinus, Trifolium campestre, Spergularia media, Silene sclerocarpa and Petrorhagia nanteuilii). In 280 contrast, solely one plant species (Sonchus oleraceous) was found in HPS plots which 281 was not found in MPS (Table S2). Rosmarinus officinalis, Chamaemelum mixtum, 282 Agrostis truncatula, Spergularia rubra, Logfia minima and Cynodon dactylon were 283 found in HPS and MPS plots. Furthermore, biochar application enhanced vegetation 284 285 diversity, as Trifolium campestre, Spergularia media, Silene scleorocarpa and Petrorhagia nanteuilii solely grew in MPS biochar-amended plots. Logfia minima was 286 strictly found in OPB plots, but not in the unamended ones (Table S2). 287

Figure 3a and b shows the average number of different plant species and individuals, respectively, in biochar-amended and unamended plots 12 months after the setup of the experiment. A greater diversity of vegetation species was observed in MPS than in

HPS plots (Fig. 3a), as also occurred for the number of individuals (Fig. 3b). One-way ANOVA showed that the number of individuals per square meter (Fig. 3b) and vegetation cover (Fig. 3c) in HPS were significantly (p < 0.05) lower than in MPS. The application of OPB in HPS plots significantly increased the area of vegetation cover in comparison with the control plots (C\_HPS; Fig. 3c). In contrast, although an increase in vegetation cover was observed for HPS amended with RHB, it was not statistically significant (Fig. 3c).

Concerning the fresh weight per plot (Fig. 3d), it increased significantly due to OPB application into HPS, but no significant differences were found for MPS plots. However, the increase of plant fresh weight in OPB\_HPS was statistically similar to OPB\_MPS. For MPS plots, the application of biochar did not promote statistical differences (p >0.05) among samples for all parameters (Fig. 3 b–d).

303

# 304 **3.5. Pearson correlations of soil properties**

305 Table S3 shows that pH was positively correlated with soil moisture measured at 40 °C 306 and soil respiration after 6 months (p < 0.05; Pearson coefficients were 0.877 and 307 0.917, respectively) and 20 months after setup (p < 0.05; Pearson coefficients were 0.903 and 0.964, respectively). Soil moisture correlated with WHC at month 6 of the 308 experiment (p < 0.05; Pearson coefficient 0.835) and with soil respiration after 20 309 months (p < 0.05; Pearson coefficient 0.862). Nevertheless, 12 months after biochar 310 application, only the pH was negatively correlated with soil moisture measured at 40 °C 311 (p < 0.05; Pearson coefficient -0.984), indicating variability of soil properties with time 312 313 and climate conditions.

314 Pearson correlations were performed between vegetation results and between 315 vegetation results and soil properties (Table S4). Fresh weight correlated positively 316 with dry weight, number of species and number of individuals (p < 0.05; Pearson coefficients between 0.839-0.877). Plots with greater number of species also showed 317 greater number of individuals (p < 0.05; Pearson coefficient 0.959). Positive correlation 318 was found between pH and fresh weight, number of individuals and number of species 319 (p < 0.05, Pearson coefficients 0.851, 0.867 and 0.949, respectively). In addition, 320 positive correlation was found between WHC and fresh weight (p < 0.05; Pearson 321 322 coefficient 0.870). Negative correlations were found between soil moisture and vegetation results. 323

324

#### 325 **3.6. Bacterial community composition**

326 3.6.1 Sequence data

The number of raw sequence reads ranged from 113072 to 237356 for samples 327 328 collected at  $t_6$ , from 377786 to 704600 for  $t_{12}$  and from 291568 to 415478 for  $t_{20}$ . After 329 guality filtering and denoising, a total of 2354783 paired-end sequences were obtained 330 for all samples. These sequences were clustered into 16964 OTUs at 97% similarity, 331 containing both assigned and non-identified bacteria. Samples from t<sub>12</sub> showed the 332 highest number of OTUs (8000), followed by  $t_{20}$  (4863) and  $t_6$  with 4101 OTUs. For all 333 the samples, the rarefaction curves reached the plateau (data not shown), suggesting 334 that we obtained a good representation of the microbial communities from both soil 335 types (HPS and MPS).

336

### 337 3.6.2. Differences between HPS and MPS

The microbial communities from all the samples were almost exclusively composed of bacteria, with the exception of the control sample HPS (C\_HPS), where Archaea accounted for 0.47%, being represented by the *Thaumarchaeota* and *Euryarchaeota* phyla.

Differences in the taxonomic composition were clearly observed between both types of 342 soil and between sampling campaigns, particularly between t<sub>6</sub> and t<sub>12</sub> (Fig. 4 and Table 343 344 S5). MPS plots at t<sub>6</sub> showed to be more diverse at the phylum level than HPS. At t<sub>6</sub>, the most abundant phyla found in MPS plots were Proteobacteria, Planctomycetes, 345 Acidobacteria, 346 Actinobacteria, Bacteroidetes, Gemmatimonadetes and Verrucomicrobia, whereas in HPS Chloroflexi, Proteobacteria, Actinobacteria, 347 Acidobacteria, Firmicutes and Saccharibacteria were the most abundant (Fig. 4a). 348

The *Planctomycetes* phylum was mostly observed in MPS and in very low relative abundance in HPS. *Bacteroidetes*, *Gemmatimonadetes* and *Verrumicrobia* were solely found in MPS (amended and control plots), whereas *Firmicutes* was mostly found in HPS (7% in C\_HPS and 2-3% in amended HPS samples). Interestingly, *Saccharibacteria* was solely found in the amended HPS plots.

After 1 year ( $t_{12}$ ), the relative abundance of bacterial phyla changed considerably in comparison with  $t_6$  (Fig. 4b). *Actinobacteria*, *Chloroflexi*, *Proteobacteria*, *Acidobacteria*, *Firmicutes* and *Planctomycetes* contributed to 80% and 90% of the total bacterial sequences in MPS and HPS, respectively. *Verrumicrobia* and *Gemmatimonadetes* were solely found in MPS (amended and control plots). *Patescibacteria* was found in the control and amended MPS samples, as well as in amended HPS.

Differences in microbial community composition at the phylum level were not significantly observed between  $t_{12}$  and  $t_{20}$  (Fig. 4b and c). At  $t_{20}$  the predominant phyla were Actinobacteria, Chloroflexi, Proteobacteria, Acidobacteria, Firmicutes, Planctomycetes and Bacteroidetes for both soils (Fig. 4c).

Remarkable differences in microbial community composition at the order level were noticeable between HPS and MPS plots, as the bacterial sequences in MPS were almost absent in HPS samples (Fig. S1).

Principal components analysis (PCA) was computed to explain differences between 367 samples (Fig. 5). At t<sub>6</sub>, the first two components explained 95% of the variation 368 observed (Fig. 5a). The plot of the loadings of PC-1 vs PC-2 defined two clusters, 369 corresponding to each soil type. This showed that microbial diversity from HPS 370 371 samples is significantly different from MPS samples. However, within cluster 1 (HPS 372 samples), PC2 significantly separates the HPS control sample from the biochar-treated 373 HPS samples along the projected plane (Fig. 5a). In contrast, no significant differences of bacterial community composition were noted between amended and unamended 374 375 MPS plots at t<sub>6</sub>.

At  $t_{12}$  (Fig. 5b), the plot also displays clear discrimination between both types of soils, but in addition contains separation within cluster 1 among biochar treatments.

At  $t_{20}$  (Fig. 5c), PC-1 (70%) vs PC-2 (12%) scores of the control and biochar-treated soils also define two clusters, reinforcing that microbial diversity from both soil types is significantly different along the time span of the field experiment. However, PC-2 discriminates samples within each cluster. Specifically, samples treated with OPB for both types of soils were separated from their corresponding control and RHB-treated samples (Fig. 5c), revealing changes in the microbial diversity for OPB-treated soils after 20 months of incubation.

Venn diagrams were plotted to calculate the number of unique and shared OTUs among the HPS and MPS samples at  $t_6$ ,  $t_{12}$  and  $t_{20}$  (Fig. S2). Interestingly, the number of shared taxa between treatments (control and amended plots) was remarkably higher than the number of unique taxa, at  $t_6$ ,  $t_{12}$  and  $t_{20}$ , especially in MPS.

389 The number of shared taxa in HPS increased over time (Fig. S2c and e). The largest OTUs numbers were shared between all the MPS samples (260 at t<sub>6</sub>, 504 at t<sub>12</sub> and 390 330 at  $t_{20}$ ), whereas the unique taxa ranged between 5 (at  $t_6$  for C\_MPS) and 104 (at  $t_{12}$ 391 392 for C\_MPS). At t<sub>6</sub>, 34 OTUs (50%) were uniquely present in HPS plots, while in MPS 56 OTUs (16%) corresponded to unique taxa (Fig. S2a). At t<sub>20</sub> (Fig. S2e and f), the 393 394 percentage of unique taxa was 45% in HPS against 34% in MPS, suggesting that the 395 bacterial communities became in general more similar between the two soil types over time. Focusing on the biochar treatments, there were more overlapped OTUs among 396 the amended plots than between biochar-treated plots and the controls at t<sub>6</sub> (Fig. S2a 397 398 and b) and  $t_{12}$  (Fig. S2c and d).

399

## 400 **3.6.3.** Impact of biochar amendment on HPS and MPS

At month 6 (Fig. 4a), the relative abundance of soil microbiota from the HPS control plot (C\_HPS) differed markedly from those treated with biochar (RHB\_HPS and OPB\_HPS). The *Chloroflexi* phylum was the most abundant in the C\_HPS plot, but decreased notably from 83% to 50% in the biochar-amended HPS samples, as well as *Firmicutes*. In contrast, the relative abundance of *Proteobacteria* in HPS increased from 2 to 16% with biochar application, as well as *Acidobacteria* (Fig. 4a and Table S5).

For MPS plots, no significant changes were noted on the relative abundance of soilmicrobiota between biochar-amended and unamended plots.

After 1 year of biochar application into HPS (Fig. 4b), the relative abundance of *Actinobacteria* slightly decreased with biochar amendment from 47 to 30-37%. *Chloroflexi* increased from 20% in the control to 40% in the biochar-amended HPS
samples.

At  $t_{20}$  (Fig. 4c), an increase was observed on the relative abundance of *Proteobacteria* (from 14 to 17-23%) and *Bacteroidetes* (from 0.4 to 4-14%) for HPS plots amended with biochar. In contrast, biochar application reduced the relative abundance of *Chloroflexi* (from 35 to 27-28%), *Acidobacteria* (from 12 to 5%) and *Firmicutes* (from 7 to 3-5 %) in the HPS samples.

In MPS plots at  $t_{20}$ , the application of biochar increased the relative abundance of *Actinobacteria* (from 14 to 17-18%) and reduced the abundance of cyanobacteria (from 9 to 0.4-1%).

At the order level, the most abundant taxa, representing Chloroflexi in C\_HPS at t<sub>6</sub>, 422 423 belonged to the order Ktedonobacterales (71%), followed by the enigmatic phylotypes 424 JG30-KF-AS9 and B12-WMSP1 also within the class Ktedonobacteria, both 425 contributing to 9% of the total bacterial sequences (Fig. S1a). The relative abundance 426 of this Ktedonobacterial community was almost reduced by half (from 80% to 46%) with the incorporation of biochar. However, the relative contribution of B12-WMSP1 and 427 JG30-KF-AS9 increased significantly from 4% to 32-37% and from 5% to 12-16%. 428 429 respectively. In contrast, a sharp decrease was observed for members of the order Ktedonobacterales (from 71% to 2% in RHB\_HPS and 0.5% in OPB\_HPS). The 430 431 abundances of Rhodospirillales (within the Proteobacteria phylum) and 432 Acidobacteriales (within Acidobacteria) were also higher in biochar-amended HPS. The order Bacillales, belonging to the phylum Firmicutes and solely represented by the 433 genus Alicyclobacillus in C\_HPS, decreased from 7% to 3% in OPB-amended HPS 434 and to 1.6% in RHB-amended HPS. 435

436 The most abundant orders found across all treatments in MPS plots at  $t_6$  (Fig. S1a) 437 were *Tepidisphaerales* (within *Planctomycetes*), *Sphingomonadales* (within

Alphaproteobacteria), Shingobacteriales (within Bacteroidetes), Burkholderiales (within
Betaproteobacteria) and Rhizobiales (within Alphaproteobacteria).

440 At  $t_{12}$  and  $t_{20}$  (Fig. S1b and c), a greater bacterial diversity at the order level is observed for all the treatments in comparison with t<sub>6</sub>, and bacterial communities in biochar-441 442 amended plots became more similar to their corresponding control plots, particularly at t<sub>20</sub> (Fig. S1c). Nevertheless, at t<sub>12</sub>, the relative abundance of B12-WMSP1, representing 443 444 the Ktedonobacterial community, was higher in the biochar-amended HPS plots (Fig. 445 S1b), as also observed at  $t_6$ . The order *Frankiales*, belonging to the *Actinobacteria* 446 phylum, increased markedly at t<sub>12</sub> across all treatments, independently of biochar 447 application. It is also worth noting the increase of Acetobacterales, representing 448 Alphaproteobacteria, with biochar amendments in HPS plots. At t<sub>20</sub>, communities in biochar-amended plots became in general more similar to their corresponding control 449 plots (Fig. S1c). However, after 20 months differences in the relative abundance of 450 451 bacterial taxa within MPS samples were noticed for OPB-treated MPS (Fig. S1c).

452

### 453 **3.7. Bacterial diversity**

The diversity of microbial community structure in the HPS and MPS samples was estimated by alpha diversity and richness indices, revealing values significantly different between the samples. MPS samples (control and amended) showed higher alpha diversity (Shannon and Simpson) and richness (Chao1 and OTU count) than HPS samples (Table 3). Shannon index values ranged from 2.00 to 4.24 in HPS samples, with an average of 3.23, and from 4.47 to 5.33 in MPS samples, with an average of 4.73.

The observed Simpson index of diversity ranged from 0.71 to 0.97, with an average of 0.91 for HPS samples, and from 0.97 to 0.99, with an average of 0.98 for MPS (Table 3).

464 Shannon and Simpson index values increased in the HPS due to biochar addition at  $t_6$ . 465 This increase in alpha diversity indices was also observed for MPS plots at  $t_6$ . 466 Regardless of the presence or absence of biochar, alpha diversity increased through 467 the time span of the experiment (Table 3).

468

# 469 **3.8 Correlation between soil properties and microbial community composition**

Soil physicochemical properties and bacterial abundance variables were used to generate correlation heatmaps for  $t_6$ ,  $t_{12}$  and  $t_{20}$  (Fig. 6). At  $t_6$ , pH, WHC and soil respiration were significantly (p < 0.05) and positively correlated with the most abundant bacterial phyla found in MPS plots (*Gemmatimonadetes*, *Planctomycetes*, *Verrucomicrobia*, *Bacteroidetes*), and negatively correlated with *Chloroflexi* and

*Firmicutes*, which were the most abundant phylum in HPS plots (Fig. 6a and Table S6). Soil moisture measured at 40 °C and total moisture (measured at 105 °C) were also positively correlated with most of bacterial phyla commonly found in MPS samples, particularly *Proteobacteria* and *Acidobacteria*. Soil TC and dehydrogenase activity showed weak correlation (positive or negative) with the most abundant phyla retrieved in the soil samples. Similarly, glucosidase activity showed almost no correlation with soil microbial communities (Fig. 6a).

At t<sub>12</sub>, the most abundant phyla detected in MPS plots were strongly and positively correlated with soil pH, WHC, dehydrogenase and glucosidase activities, as well as with the botanical variables measured at t<sub>12</sub> (fresh weight, number of plant species and individuals), but negatively correlated with *Actinobacteria* (Fig. 6b and Table S6). *Chloroflexi* was negatively correlated with pH, whereas it was positively correlated with soil moistures measured at 40 and 105 °C (p < 0.05). Soil respiration and TC showed no significant relationship with soil microbial communities.

489 At  $t_{20}$ , all the soil physicochemical parameters measured in this study were positively 490 correlated with MPS microbial communities, and negatively correlated with *Chloroflexi*, 491 which was almost exclusively found in HPS samples (Fig. 6c and Table S6).

492

#### 493 **4. Discussion**

Physical, chemical and biological parameters were monitored for 6, 12 and 20 months in biochar-amended soils with two different levels of trace-element contamination under field conditions. These parameters (pH, carbon content, WHC, soil moisture, enzymatic activities, soil respiration, vegetation cover and microbial diversity) were selected to integrate the three types of soil quality indicators, which allow assessing the capability of a soil to perform its ecological functions (Arias et al., 2005).

500 Soil properties and, consequently, their plant and microbial diversity were very different 501 in HPS and MPS, independently of biochar addition. The soil properties of HPS plots 502 measured before biochar application indicated a very degraded soil with extreme 503 difficulties to sustain ecological functions. Although biochar application induced 504 changes on soil properties, climatic conditions need to be considered, as changes 505 between samplings were notable.

The dehydrogenase activity (DHA) has been also proposed as a good indicator of the toxicity of trace elements (Dick et al., 1996). Under acidic conditions, this enzymatic activity can be inhibited due to the destruction of ion and hydrogen bonds in the enzyme active centre and the alteration of its three-dimensional shape (Frankenberger and Johanson, 1982). This explains the greater values of dehydrogenase activity observed for the less acidic MPS plots, in comparison with HPS, and the positive

512 correlation between soil pH and dehydrogenase activity (Fig. S3). The low β-513 glucosidase activity measured for all HPS plots, regardless of biochar addition, can be related to soil pH (Eivazi and Tabatai, 1990), and the low abundance of labile organic 514 matter (Ferraz de Almeida et al., 2015). This low enzymatic activity indicates a high 515 recalcitrance of the applied biochar in HPS, as biochar has condensed aromatic 516 517 structures that make them less available to microbial degradation (Elzobair et al., 2016; Günal et al., 2018; Sohi et al., 2010). Despite of the increase of C content in soils 518 519 caused by biochar addition, respiration measurements showed that the application of 520 OPB or RHB to the Fluvisol did not modify CO<sub>2</sub> emissions rates. This is similar to the 521 findings previously reported by other authors (Sun et al., 2014; Phongthep et al., 2017). 522 In this study, no priming effect is found and a high stability of both sorts of biochar can 523 be predicted. Considering that soil metal pollution is a significant environmental issue, 524 the use of biochar is worthwhile for the remediation of trace element-polluted soils.

As expected, MPS plots showed greater diversity and abundance of vegetation species than HPS (Fig. 3). Comparing both biochar treatments, the application of OPB enhanced not only plant diversity but also the primary productivity in HPS.

528 The combination of digital image analysis, for measuring the total area of soil covered by the vegetation canopy, and the plant fresh weight approach, which provides 529 530 information on the plant yield, gave a rather good presentation of the effect of biochar addition on the vegetation production (Fig. 3c and d). It is interesting to note that the 531 532 application of OPB in HPS plots promoted a significant increase of fresh weight, reaching values similar to those observed for MPS (Fig. 3d), while the vegetation cover 533 534 of OPB-HPS was five times lower than the unamended and amended MPS plots (Fig. 535 3c). This is explained by the presence of different plant species in OPB-HPS and MPS 536 plots (Table S2). In MPS, the plant species greatly covered the soil surface (high 537 vegetation coverage area), but their stem diameter and height were much smaller than the species found in HPS plots. In OPB-treated HPS plots, few plants were found but 538 they displayed greater height and stem diameter, and less vegetation coverage. 539

540 The positive correlations obtained between plant data (number of species and 541 individuals, and fresh weight) with soil pH (Table S4), demonstrate that biochar is able 542 to enhance the properties of acidic soils, favouring the recovery of degraded polluted 543 soils due to the spill of heavy metals.

544 Changes in soil microbial community were also assessed in the biochar-amended and 545 untreated soils to inform about soil quality and biochar potential to restore soil 546 functionality. Monitoring microbial diversity by 16S rRNA gene NGS-based analyses 547 after 6, 12 and 20 months of biochar addition showed changes in the soil microbial 548 community structure, particularly in HPS plots after 6 months of soil amendment with

biochar. However, after 12 and 20 months, we did not find consistent phylum or orderlevel responses to biochar amendments (Fig. 4 and Fig. S1), as the treated plots showed higher similarity over control soils, as also reported by Song et al. (2017). Similarly, Shannon and Simpson index values indicated that the addition of RH and OP biochar solely promoted soil bacterial diversity in HPS at  $t_6$  (Table 3). These findings suggest that the type and dosages of biochar applied into HPS had a short-term effect on the distribution of microbial communities, which was dissipated over time.

- 556 From the microbial community structure displayed in Figure 4, we drew the conclusion 557 that biochar addition significantly decreased the relative abundance of members of the 558 *Chloroflexi* phylum in HPS-amended plots at t<sub>6</sub>, probably due to changes in soil pH and 559 elements immobilization as Chloroflexi have preference for extreme environments (Soo et al., 2009; Yabe et al., 2017). This phylum was mainly represented by the order 560 Ktedonobacterales (with 71% of relative abundance in C HPS) from the class 561 562 Ktedonobacteria, which are filamentous bacteria that inhabit forest and garden soils at 563 low abundances, as well as extreme environments such as geothermal areas and 564 caves (Yabe et al., 2017). The relative abundance of the Ktedonobacterial community (80%) in the HPS control plots abruptly declined (from 80% to 46%) with the 565 566 incorporation of biochar (Fig. S1a), probably due to changes in pH, which explains the negative correlation between pH and Chloroflexi in HPS (Fig. 6a). This decline in 567 Chloroflexi abundances after biochar application was also previously reported by 568 several authors (Nielsen et al., 2014; Xu et al., 2014; Ali et al., 2019; Li et al., 2019). 569 570 However, Chen et al. (2019) showed an increase in *Chloroflexi* with the application of 571 10% of biochar to calcareous soils.
- The relative abundance of *Firmicutes* was also reduced in HPS after 6 months of biochar application. *Firmicutes* can adapt to low nutrient environments and thrive in extreme conditions by forming spores (Bai et al., 2017; Li et al., 2014). Cole et al. (2019) also found a decline in relative abundance of *Firmicutes* with biochar application. However, Ali et al. (2019) reported an increase when a biochar produced from sewage sludge was applied.
- 578 Conversely, the increase of *Proteobacteria* observed in HPS-amended samples at  $t_6$  is 579 probably explained by their heterotrophic nature, as biochar increases soil carbon 580 content and nutrient conditions of poor-nutrient soils as HPS. Ali et al. (2019), Cole et 581 al. (2019), Li et al. (2019) and Su et al. (2015) also reported greater abundances of 582 *Proteobacteria* in amended soils than in control soils, obtaining good correlation 583 between *Proteobacteria* and labile C content.
- In addition to *Proteobacteria*, the relative abundance of *Acidobacteria* slightly increased after 6 months of biochar addition into HPS, as also reported by Cole et al. (2019).

However, Li et al. (2019) and Fan et al. (2020) reported a decrease of *Acidobacteria* after biochar application, but Jenkins et al. (2017) found an increase of *Acidobacteria* even in control soils without biochar treatment, indicating variations by weather conditions. This is in accordance with our results for control HPS over time.

590 In this study, the relative abundance of *Planctomycetes*, *Bacteroidetes*, Gemmatimonadetes and Verrucomicrobia did not depend on biochar application but on 591 soil type and seasonal changes. Planctomycetes were more abundant in MPS plots 592 593 than in HPS and their relative abundance varied with different seasons. Rice husk 594 biochar only slightly reduced *Planctomycetes* after 12 months of application into HPS, 595 which is in accordance with the findings of Noyce et al. (2016) when low pH soil was 596 amended with wood chips biochar. However, Ali et al. (2019) showed an increase in 597 Planctomycetes abundance in a contaminated-agricultural soil after the application of rice straw biochar. 598

599 Chen et al. (2019) observed that the relative abundance of Bacteroidetes was higher in 600 the control soil than in the biochar-amended soil, attributing these changes to the initial 601 high pH and nutrient levels in the studied calcareous soils. However, Hu et al. (2014) 602 solely detected Bacteroidetes in the biochar amended soil. In this study, Bacteroidetes were found in control and amended MPS plots, but not in HPS. It could be due to their 603 604 copiotrophic nature and capability for living in rhizosphere conditions (Shen et al., 2018), as plant growth was solely observed in MPS plots at  $t_{6}$ . Khodadad et al. (2011) 605 reported an increase of Gemmatimonadetes in soils with natural or added pyrogenic 606 carbon, suggesting an active role of these microorganisms in soil pyrogenic C 607 608 metabolism. We observed a small increase when RHB was applied, which could 609 indicate that this biochar could be more accessible than OPB for these group of 610 bacteria.

611 Verrucomicrobia was only found in MPS plots, suggesting that its presence was 612 dependent on the soil type, instead of biochar application. In fact, Chen et al. (2019) 613 observed that Verrucomicrobia was greater in control than in biochar-amended soils. 614 Nevertheless, Fan et al. (2020) reported an increase in Verrucomicrobia phylum in soils 615 amended with biochar.

Actinobacteria are possibly involved in the redistribution of consumed C or in the degradation of more recalcitrant compounds (Blagodatskaya and Kuzyakov, 2008). Cole et al. (2019) and Khodadad et al. (2011) reported an increase in Actinobacteria in soils with natural or added pyrogenic carbon. However, Li et al. (2019) reported a decrease in the relative abundance of Actinobacteria after biochar addition to soil. Our results are more in accordance with this decline, particularly in RHB-amended MPS plots at t<sub>6</sub> and in HPS plots at t<sub>12</sub>. Jenkins et al. (2017) found an increase of

623 *Actinobacteria* even in control soils without biochar, indicating variations due to weather 624 conditions. In this study, the relative abundance of *Actinobacteria* also seemed to be 625 related to seasonal changes particularly in the case of HPS plots (Fig. 4).

In summary, the effects of biochar on soil bacterial communities are not unanimously 626 explained, as numerous other factors, such as soil type, pH, moisture and biochar 627 feedstock are likely to structure microbial communities (Chen et al., 2019; Jenkins et 628 629 al., 2017). In addition, environmental conditions and long-term biochar application may 630 have more influence in soil microbial communities than biochar types. It is worth 631 mentioning that this variability in soil microbial communities is mostly find in field 632 experiments, whereas in pot incubation experiments parameters are constrained (Hu et 633 al., 2014; Xu et al., 2014). Overall, the changes in the soil bacterial richness and diversity after soil amendment application were correlated with changes in soil pH (Fig. 634 6), as the incorporation of biochar increased pH, and bacterial diversity, as well as 635 636 plant growth.

637

## 638 **5. Conclusions**

This field study conducted on polluted acidic soils has shown that the addition of 639 biochar allowed the recovery of plant cover and increased plant biodiversity, 640 641 particularly in moderately contaminated soils (MPS). Biochar application did not modify soil CO<sub>2</sub> emissions, nor significantly increase enzymatic activity beyond the first six 642 months of biochar application, which points to a great stability of the tested olive pit and 643 644 rice husk biochar (OPB and RHB) and their ability to be used for carbon sequestration 645 in degraded soils. Findings from 16S rRNA gene next-generation sequencing revealed 646 that the incorporation of biochar modified the soil microbial community in the highly 647 polluted soil (HPS). Bacterial diversity was found to be site-specific as the properties 648 differed among the studied soils. We conclude that the application of biochar from crop residues to trace-element polluted soils participated in soil conditioning, promoting 649 plant development, increasing bacterial diversity and soil carbon stabilization. This 650 651 suggested that the application of biochar is important in the ecological restoration of these degraded soils. Our results showed that long-term experiments under field 652 653 conditions are essential in the quest to investigate the performance of biochar without 654 constraining environmental parameters, as seasonal changes were remarkable in this study. This knowledge could help to fully understand the impact of biochar on global 655 656 nutrient cycles and on the recovery of soil ecological functions.

657

### 658 **Declaration of competing interest**

The authors declare that they have no conflict of interest to this work.

660

# 661 Acknowledgments

The Spanish Ministry of Economy, Industry and Competitiveness (MINEICO) and 662 AEI/FEDER UE are thanked for funding the project CGL2016-76498-R. P. Campos 663 thanks the "Fundación Tatiana Pérez de Guzmán el Bueno" for funding her PhD. 664 MINEICO is also acknowledged for funding the "Ramón y Cajal" post-doctoral contract 665 of José M. De la Rosa [grant number RYC2014-16338]. Ana Z. Miller acknowledges 666 667 the support from the Portuguese "Fundação para a Ciência e a Tecnologia" (FCT) 668 [grant number CEECIND/01147/2017]. S. Prats thanks the Portuguese FCT for his 669 research contract [CDL-CTTRI-88-ARH/2018 REF.-138-88-ARH/2018], funded in the 670 scope of Law 57/2017 and for the financial support to CESAM (UID/AMB/50017/2019).

671

# 672 **References**

AEMET. 2020. Available at: https://datosclima.es/Aemet2013/Tempestad2013.php(accessed on 20 May 2020).

- Ali, N., Khan, S., Li, Y., Zheng, N., Yao, H., 2019. Influence of biochars on the
  accessibility of organochlorine pesticides and microbial community in contaminated
  soils. Science of the Total Environment 647, 551–560.
- Arias, M.E., González-Pérez, J.A., González-Vila, F.J., Ball A.S., 2005. Soil health-a
  new challenge for microbiologists and chemists. International Microbiology 8, 13–21.
  Bai, R., Wang, J.-T., Deng, Y., He, J.-Z., Feng, K., Zhang, L.-M., 2017. Microbial
  community and functional structure significantly varied among distinct types of
  paddy soils but responded differently along gradients of soil depth layers. Frontiers
  in Microbiology 8, 945, 1–16.
- Bamminger, C., Marschner, B., Jüschke, E., 2014. An incubation study on the stability
  and biological effects of pyrogenic and hydrothermal biochar in two soils. European
  Journal of Soil Science 65, 72–82.
- Beesley, L., Moreno-Jiménez, E., Gomez-Eyles, J.L., 2010. Effects of biochar and
  greenwaste compost amendments on mobility, bioavailability and toxicity of
  inorganic and organic contaminants in a multi-element polluted soil. Environmental
  Pollution 158, 2282–2287.
- Blagodatskaya, E., Kuzyakov, Y., 2008. Mechanisms of real and apparent priming
  effects and their dependence on soil microbial biomass and community structure:
  critical review. Biology and Fertility of Soils 45, 115–131.
- Bolyen, E., Rideout, J.R., Dillon, M.R. et al. 2019. Reproducible, interactive, scalable
  and extensible microbiome data science using QIIME 2. Nature Biotechnology 37,
  852–857.

Cabrera, F., Clemente, L., Díaz Barrientos, E., López, R., Murillo, J.M., 1999. Heavy
metal pollution of soils affected by the Guadiamar toxic flood. Science of the Total
Environment 242, 117–129.

Campos, P., De la Rosa, J.M., 2020. Assessing the effects of biochar on the
immobilization of trace elements and plant development in a naturally contaminated
soil. Sustainability, 12, 6025–6044.

- Campos, P., Miller, A.Z., Knicker, H., Costa-Pereira, M.F., Merino, A., De la Rosa, J.M.,
  2020. Chemical, physical and morphological properties of biochars produced from
  agricultural residues: Implications for their use as soil amendment. Waste
  Management 105, 256–267.
- Chen, J., Lee, X., Tang, Y., Zhang, Q., 2019. Long-term effects of biochar amendment
  on rhizosphere and bulk soil microbial communities in a karst region, southwest
  China. Applied Soil Ecology 140, 126–134.
- Cole, E.J., Zandvakili, O.R., Blanchard, J., Xing, B., Hashemi, M., Etemadi, F., 2019.
  Investigating responses of soil bacterial community composition to hardwood
  biochar amendment using high-throughput PCR sequencing. Applied Soil Ecology
  136, 80–85.
- Dick, R.P., Breakqill, D., Turco, R., 1996. Soil enzyme activities and biodiversity
  measurements as integrating biological indicators. In Doran, J.W., Jones, A.J.,
  Handbook of Methods for Assessment of Soil Quality. Soil Science Society of
  America Specific Publications, Madison, WI, 242–272.
- De la Rosa, J.M., Paneque, M., Miller, A.Z., Knicker, H., 2014. Relating physical and
   chemical properties of four different biochars and their application rate to biomass
   production of Lolium perenne on a Calcic Cambisol during a pot experiment of 79
   days. Science of the Total Environment 499, 175–184.
- De la Rosa, J.M., Rosado, M., Paneque, M., Miller, A.Z., Knicker, H., 2018. Effects of
  aging under field conditions on biochar structure and composition: Implications for
  biochar stability in soils. Science of the Total Environment 613-614, 969–976.

De la Rosa, J.M., Sánchez-Martín, A.M., Campos, P., Miller, A.Z., 2019. Effect of pyrolysis conditions on the total contents of polycyclic aromatic hydrocarbons in biochars produced from organic residues: Assessment of their hazard potential.

- biochars produced from organic residues: Assessment of their hazard potential.
  Science of the Total Environment 667, 578–585.
- EEA; 2007. Progress in Management of Contaminated Sites. CSI 015. Copenhagen,
   Denmark: European Environmental Agency.
- Fivazi, F., Tabatai, M., 1990. Factors affecting glucosidase and galactosidase in soils.
  Soil Biology and Biochemistry 22, 145–152.

- Elzobair, K.A., Stromberger, M.E., Ippolito, J.A., Lentz, R.D., 2016. Contrasting effects
   of biochar versus manure on soil microbial communities and enzyme activities in an
- 735 Aridisol. Chemosphere 142, 145–152.
- Fan, S., Zuo, J., Fond, H., 2020. Changes in Soil Properties and Bacterial Community
   composition with Biochar Amendment after Six Years. Agronomy 10, 746, 1–15.
- Ferraz de Almeida, R., Rezende Naves, E., Pinheiro da Mota, R., 2015. Soil quality:
- 739 Enzymatic activity of soil β-glucosidase. Global Journal of Agricultural Research and
  740 Reviews 3, 146–150.
- Frankenberger, W., Johanson, J., 1982. Effect of pH On Enzyme Stability in Soils. Soil
  Biology and Biochemistry 14, 433–437.
- Gascó, G., Paz-Ferreiro, J., Cely, P., Plaza, C., Méndez, A., 2016. Influence of pig
  manure and its biochar on soil CO<sub>2</sub> emissions and soil enzymes. Ecological
  Engineering 95, 19–24.
- Günal, E., Erdem, H., Demirbaş, A., 2018. Effects of three biochar types on activity of
   β-glucosidase enzyme in two agricultural soils of different textures. Archives of
   Agronomy and Soil Science, 64, 14, 1963–1974.
- Hagemann, N., Schmidt, H.-P., Kagi, R., Bohler, M., Sigmund, G., Maccagnan, A.,
  McArdell, C.S., Bucheli, T.D., 2020. Wood-based activated biochar to eliminate
  organic micropollutants from biologically treated wastewater. Science of the Total
  Environment 730, 138417, 1–11.
- Hagemann, N., Spokas, K., Schmidt, H.-P., Kägi, R., Böhler, M.A., Bucheli, T.D., 2018.
  Activated Carbon, Biochar and Charcoal: Linkages and Synergies across Pyrogenic
  Carbon's ABCs. Water 10, 182, 1–19.
- Han, G., Lan, J., Chen, Q., Yu, C., Bie, S., 2017. Response of soil microbial community
  to application of biochar in cotton soils with different continuous cropping years.
  Scientific Reports 7, 10184, 1–11.
- Hu, L., Cao, L., Zhang, R., 2014. Bacterial and fungal taxon changes in soil microbial
  community composition induced by short-term biochar amendment in red oxidized
  loam soil. World Journal of Microbiology and Biotechnology 30, 1085–1092.
- 762 IUSS Working Group WRB, 2015. World Reference Base for Soil Resources 2014,
- update 2015. International soil classification system for naming soils and creating
  legends for soil maps. World Soil Resources Reports 106. FAO, Rome, 1–203.
- Jenkins, J. R., Viger, M., Arnold, E.C., Harris, Z.M., Ventura, M., Miglietta, F., Girardin,
  C., Edwards, R.J., Rumpel, C., Fornasier, F., Zavalloni, C., Tonon, G., Alberti, G.,
  Taylor, G., 2017. Biochar alters the soil microbiome and soil function: results of nextgeneration amplicon sequencing across Europe. Global Change Biology Bioenergy
  9, 591–612.

- Jian, L.-L., Han, G.-.M., Lan, Y., Liu, S.-N., Gao, J.-P., Yang, X., Meng, J., Chen, W.-F.,
  2017. Corn cob biochar increases soil culturable bacterial abundance without
  enhancind their capacities in uitlizing carbon sources in Biolog Eco-plates. Journal
  of Integrative Agriculture 16, 712–724.
- Kammann, C., Ratering, S., Eckhard, C., Müller, C., 2012. Biochar and hydrochar
  effects on greenhouse gas (carbon dioxide, nitrous oxide, methane) fluxes from
  soils. Journal of Environmental Quality 41, 1052–1066.
- Karer, J., Wawra, A., Zehetner, F., Dunst, G., Wagner, M., Pavel, P.-B., Puschenreiter,
   M., Friesl-Hanl, W., Soja, G., 2015. Effects of Biochars and Compost Mixtures and
   Inorganic Additives on Immobilisation of Heavy Metals in Contaminated Soils.
- 780 Water, Air, & Soil Pollution 226, 342, 1–12.
- Khodadad, C.L.M., Zimmerman, A.R., Green, S.J., Uthandi, S., Foster, J.S., 2011.
  Taxa-specific changes in soil microbial community composition induced by
  pyrogenic carbon amendments. Soil Biology and Biochemistry 43, 385–392.
- Knicker, H., Hilscher, A., De la Rosa, J.M., González-Pérez, J.A., González-Vila, F.J.,
  2013. Modification of biomarkers in pyrogenic organic matter during the initial phase
  of charcoal biodegradation in soils. Geoderma 197–198, 43–50.
- Kolb, S.E., Fermanich, K.J., Dornbush, M.E., 2009. Effect of charcoal quantity on
  microbial biomass and activity in temperate soils. Soil Science Society of America
  Journal 73, 1173–1181.
- Kumar A, Joseph S, Tsechansky L, Privat, K., Schreiter, I.J., Schüth, C., Graber, E.R.,
  2018. Biochar aging in contaminated soil promotes Zn immobilization due to
  changes in biochar surface structural and chemical properties. Science of the Total
  Environment 626, 953–961.
- Kuzyakov, Y., Bogomolova, I., Glaser, B., 2014. Biochar stability in soil: Decomposition
   during eight years and transformation as assessed by compound-specific <sup>14</sup>C
   analysis. Soil Biology and Biochemistry 70, 229–236.
- Kuzyakov, Y., Subbotina, I., Chen, H., Bogomolova, I., Xu, X., 2009. Black carbon
   decomposition and incorporation into soil microbial biomass estimated by <sup>14</sup>C
   labelling. Soil Biology and Biochemistry 41, 210–219.
- Lehmann, J., Rillig, M.C., Thies, J., Masiello, C.A., Hockaday, W.C., Crowley, D., 2011.
  Biochar effects on soil biota a review. Soil Biology and Biochemistry 43, 1812–
  1836.
- Li, C., Yan, K., Tang, L., Jia, Z., Li, Y., 2014. Change in deep soil microbial
  communities due to long-term fertilization. Soil Biology and Biochemistry 75, 264272.

- Li, Y., Yang, Y., Shen, F., Tian, D., Zeng, Y., Yang, G., Zhang, Y., Deng, S., 2019.
  Partitioning biochar properties to elucidate their contributions to bacterial and fungal
  community composition of purple soil. Science of the Total Environment 648, 1333–
  1341.
- Luo, Y., Durenkamp, M., De Nobili, M., Lin, Q., Devonshire, B.J., Brookes, P.C., 2013.
  Microbial biomass growth, following incorporation of biochars produced at 350 °C or
  700 °C, in a silty-clay loam soil of high and low pH. Soil Biology and Biochemistry
  57, 513–523.
- Martín-Peinado, F. J., Romero-Freire, A., García-Fernández, I., Sierra Aragón, M.,
  Ortiz-Bernad, I., Simón Torres, M., 2015. Long-term contamination in a recovered
  area affected by a mining spill. Science of the Total Environment 514, 219–223.
- Nielsen, S., Minchin, T., Kimber, S., van Zwieten, L., Gilbert, J., Munroe, P., Joseph,
  S., Thomas, T., 2014. Comparative analysis of the microbial communities in
  agricultural soil amended with enhanced biochars or traditional fertilisers.
  Agriculture, Ecosystems and Environment 191, 73–82.
- Noyce, G.L., Winsborough, C., Fulthorpe, R., Basiliko, N., 2016. The microbiomes and
  metagenomes of forest biochars. Scientific Reports 6, 26425, 1–12.
- Oustriere, N., Marchand, L., Lottier, N., Motelica, M., Mench, M., 2017. Long-term Cu
  stabilization and biomass yields of Giant reed and poplar after adding a biochar,
  alone or with iron grit, into a contaminated soil from a wood preservation site.
  Science of the Total Environment 579, 620–627.
- Paz-Ferreiro, J., Gascó, G., Gutiérrez, B., Méndez, A., 2012. Soil biochemical activities
  and the geometric mean of enzyme activities after application of sewage sludge and
  sewage sludge biochar to soil. Biology and Fertility of Soils 48, 511–517.
- Phongthep, H., Jiranut, W., Tanakit, S., Sathaporn, J., Sukanya, T., 2017. Soil
  respiration in rubber tree plantation applied with biochar. Research Journal of
  Chemistry and Environment 21 (10), 27–34.
- Shen, G., Zhang, S., Liu, X., Jiang, Q., Ding, W., 2018. Soil acidification amendments
  change the rhizosphere bacterial community of tobacco in a bacterial wilt affected
  field. Applied Microbiology and Biotechnology 102, 9781–9791.
- Shen, Z., Hou, D., Jin, F., Shi, J., Fan, X., Tsang, D.C.W., Alessi, D.S., 2019. Effect of
  production temperature on lead removal mechanisms by rice straw biochars.
  Science of the Total Environment 655, 751–758.
- Sohi, S.P., Krull, E., Lopez-Capel, E., Bol, R., 2010. A review of biochar and its use
  and function in soil. Advances in Agronomy 105, 47–82.
- Song, Y., Bian, Y., Wang, F., Xu, M., Ni, N., Yang, X., Gu, C., Jiang, X., 2017. Dynamic
  Effects of Biochar on the Bacterial Community Structure in Soil Contaminated with

- Polyciclic Aromatic Hydrocarbons. Journal of Agricultural and Food Chemistry 65,6789–6796.
- Soo, R.M., Wood, S.A., Grzymski, J.J., McDonald, I.R., Cary, S.C., 2009. Microbial
  biodiversity of thermophilic communities in hot mineral soils of Tramway Ridge,
  Mount Erebus, Antarctica. Environmental Microbiology 11, 715–728.
- Su, P., Lou, J., Brookes, P.C., Luo, Y., He, Y., Xu, J., 2015. Taxon-specific responses
  of soil microbial communities to different soil priming effects induced by addition of
- plant residues and their biochars. Journal of Soils and Sediments 17 (3), 674–684.
- Sun, Z., Bruun, E.W., Arthur, E., Wollesen de Jonge, L., Moldrup, P., HauggaardNielsen, H., Elsgaard, L., 2014. Effect of biochar on aerobic processes, enzyme
  activity and crop yields in two sandy loam soils. Biology and Fertility of Soils 50,
  1087–1097.
- Tabatabai, M.A., 1982. Soil enzymes. In: Page, A.L., Miller, R.H., Keeney, D.R. (Eds.),
  Method of Soil Analysis, Part 2. Chemical and Microbiological Properties. American
  Society of Agronomy, Madison, 903–948.
- Tack, F., Rinklebe, J., Ok, Y.S., 2018. Interactions between biochar and trace elements
  in the environment. Science of The Total Environment 649, 792–793.
- Trevors, J.T., 1984. Dehydrogenase activity in soil: a comparison between the INT and TTC assay. Soil Biology and Biochemistry 16 (6), 673–674.
- Uchimiya, M., Bannon, D.H., Wartelle, L.H., Lima, I.M., Klasson, K.T., 2012. Lead
  retention by broiler litter biochars in small arms range soil: impact of pyrolysis
  temperature. Journal of Agricultural and Food Chemistry 60, 5035–5044.
- Uchimiya, M., Klasson, K.T., Wartelle, L.H., Lima, I.M., 2011. Influence of soil
  properties on heavy metal sequestration by biochar amendment: 2. Copper
  desorption isotherms. Chemosphere 82, 1438–1447.
- Xu, H.J., Wang, X.H., Li, H., Yao, H.-Y.Y., Su, J.Q., Zhu, Y.G., 2014. Biochar impacts
  soil microbial community composition and nitrogen cycling in an acidic soil planted
  with rape. Environmental Science Technology 48, 9391–9399.
- Xu, N., Tan, G., Wang, H., Gai, X., 2016. Effect of biochar additions to soil on nitrogen
  leaching, microbial biomass and bacterial community structure. European Journal of
  Soil Biology 74, 1–8.
- Yabe, S., Sakai, Y., Abe, K., Yokota, A., 2017. Diversity of Ktedonobacteria with
  Actinomycetes-Like morphology in terrestrial environments. Microbes and
  Environment 32, 61–70.
- Zakrzewski, M., Proietti, C., Ellis, J.J., Hasan, S., Brion, M.J., Berger, B., Krause, L.,
  2017. Calypso: a user-friendly web-server for mining and visualizing microbiomeenvironment interactions. Bioinformatics 33, 782–783.

Journal Pre-proof

# 881 **FIGURE CAPTIONS:**

882

Fig. 1. Location of the field experiment, Aznalcóllar mine and Guadiamar GreenCorridor.

885

**Fig. 2.** Enzymatic activities in control and biochar amended soils. a)  $\beta$ -Glucosidase activity in Highly Polluted Soil, b)  $\beta$ -Glucosidase activity in Moderately Polluted Soil, c) Dehydrogenase activity in Highly Polluted Soil and d) Dehydrogenase activity in Moderately Polluted Soil. Different letters for each sampling period indicate significant differences between treatments (*p* < 0.05) based on one-way ANOVA test followed by the Tukey HSD test.

892

**Fig. 3.** a) Number of different vegetation species per plot at  $t_{12}$ . b) Number of plants per m<sup>2</sup> in control and biochar amended plots at  $t_{12}$ . c) Average of vegetation cover (%) per plot. d) Average fresh weight of plants per plot. Different letters indicate significant differences between treatments (p < 0.05) based on one-way ANOVA test followed by the Tukey HSD test.

898

Fig. 4. Relative abundance of the OTUs at the phylum level in the control (C\_HPS and C\_MPS) and biochar-amended soils (RHB\_HPS, OPB\_HPS, RHB\_MPS and OPB\_MPS) at: a)  $t_6$  (6 months after biochar application into soils); b)  $t_{12}$  (12 months), and c)  $t_{20}$  (20 months after biochar application).

Fig. 5. Assessment of bacterial diversity using principal component analysis (PCA) of
 highly polluted soil (HPS) and moderately polluted soil (MPS) at: a) t<sub>6</sub>; b) t<sub>12</sub>, and c) t<sub>20</sub>.
 RHB and OPB represent biochars derived from rice husk and olive pit, respectively,
 whereas C\_HPS and C\_MPS correspond to control soils.

907

908Fig. 6. Correlation heatmaps between soil physicochemical properties (pH, WHC, TC,909moisture, soil respiration, β-Glucosidase, Dehydrogenase) and bacterial abundance for910a)  $t_6$ ; b)  $t_{12}$ , and c)  $t_{20}$ .

		$\mathbf{p_r}$		

**Table 1**. pH, total carbon (TC), total nitrogen (TN) and trace elements contents of rice husk biochar (RHB), olive pit biochar (OPB), highly polluted soil (HPS) and moderately polluted soil (MPS). Values represent means (n = 3) and standard deviation.

	(g kg⁻¹	) (g kg⁻¹)	(ma ka <sup>-1</sup> )	(mag. 1/m <sup>-1</sup> )	<1x	C ,				
			(33)	(mg kg )	(mg kg <sup>-</sup> ')	(mg kg⁻¹)	(mg kg <sup>-1</sup> )	(mg kg⁻¹)	(mg kg⁻¹)	(mg kg⁻¹)
Biochars RHB 10.17	′± 0.34 537±1.	0 1.6± 0.9	7.3	0.05	35.0	1224.2	8.5	1.7	11.6	42.6
<b>OPB</b> 9.34	± 0.19 927±2.	0 5.1± 2.4	<loq*< td=""><td><loq< td=""><td>5.9</td><td><loq< td=""><td><loq< td=""><td>0.4</td><td>4.4</td><td><loq< td=""></loq<></td></loq<></td></loq<></td></loq<></td></loq*<>	<loq< td=""><td>5.9</td><td><loq< td=""><td><loq< td=""><td>0.4</td><td>4.4</td><td><loq< td=""></loq<></td></loq<></td></loq<></td></loq<>	5.9	<loq< td=""><td><loq< td=""><td>0.4</td><td>4.4</td><td><loq< td=""></loq<></td></loq<></td></loq<>	<loq< td=""><td>0.4</td><td>4.4</td><td><loq< td=""></loq<></td></loq<>	0.4	4.4	<loq< td=""></loq<>
Soils HPS 3.85	5± 0.14 7.2± 0	1 0.6± 0.1	47.1	1.28	240.6	53023.3	15.6	569.0	53.7	249.3
MPS 4.82	2± 0.13 9.0± 0	.6 0.8± 0.1	93.3	1.56	215.5	36945.7	15.6	156.5	38.6	293.5

\*<LOQ: below limit of quantification.

**Table 2**. Changes in soil characteristics, soil total carbon (TC) and soil respiration during the field experiment ( $t_1$ : 1 month-spring,  $t_6$ : 6 months-autumn,  $t_{12}$ : 12 months-spring,  $t_{20}$ : 20 months-autumn).

		WHC (%)				TC (g kg <sup>-1</sup> )		Soil respiration ( $\mu$ mol CO <sub>2</sub> m <sup>-2</sup> s <sup>-1</sup> )				
Sample	t <sub>6</sub>	t <sub>12</sub>	t <sub>20</sub>	t <sub>6</sub>	t <sub>12</sub>	t <sub>20</sub>	t <sub>6</sub>	t <sub>12</sub>	t <sub>20</sub>	t <sub>6</sub>	t <sub>12</sub>	t <sub>20</sub>
C_HPS	3.57±0.13a	3.59±0.08a	3.52±0.20a	20±7a	14±5a	32±2a	7.9±0.6a	9.6±1.3a	12.3±2.7a	1.0±0.2a	1.0±0.2a	0.9±0.2a
RHB_HPS	3.64±0.18a	3.68±0.21a	3.77±0.17a	23±1a	20±2ab	26±4a	12.9±4.8a	14.9±4.7ab	16.7±1.5a	1.0±0.3a	0.8±0.2a	0.6±0.3a
OPB_HPS	3.63±0.03a	3.64±0.04a	3.69±0.17a	23±5a	17±5ab	30±4a	13.6±8a	16.6±4.5b	18.4±6.1a	1.2±0.4a	0.9±0.2a	1.2±0.6a
C_MPS	4.18±0.21a	4.8±0.10b	4.87±0.20b	32±2b	23±5ab	47±6ab	12.4±3.5a	9.5±0.3a	21.7±1.3a	2.9±0.2b	1.2±0.8a	3.9±0.2b
RHB_MPS	4.75±0.01b	5.02±0.22b	5.07±0.18b	39±1b	25±8b	52±2b	12.2±1.7a	14.3±3.2ab	19.9±9.2a	2.7±0.3b	1.3±0.7a	3.3±0.4b
OPB_MPS	4.74±0.54b	4.86±0.50b	5.11±0.20b	38±7b	19±7ab	67±3c	17.0±7.9a	12.7±0.3ab	20.3±10.2a	3.1±0.3b	0.5±0.1a	3.7±1.0b

WHC: Water holding capacity. Different letters within a column indicate significant differences between treatments (*p* < 0.05) based on a one-way ANOVA test followed by the Tukey HSD test.

Journal Press

**Table 3**. Alpha-diversity indices of microbial community structure in the unamended and biochar-amended HPS and MPS samples. The diversity indices (Shannon and Simpson index) and richness index (Chao1 and OTUs) were determined at 97% sequence similarity.

		Alpha-diversity											
Sample	No. OTUs	Shannon				Simpson		Chao1					
ID		t <sub>6</sub>	t <sub>12</sub>	t <sub>20</sub>	t <sub>6</sub>	t <sub>12</sub>	t <sub>20</sub>	t <sub>6</sub>	t <sub>12</sub>	t <sub>20</sub>			
C_HPS	426	2.00	3.23	3.84	0.71	0.91	0.96	52	142	232			
RHB_HPS	468	2.76	3.42	3.33	0.86	0.91	0.91	76	193	199			
OPB_HPS	488	2.62	2.84	4.24	0.83	0.86	0.97	73	173	242			
C_MPS	1475	4.47	5.33	4.72	0.97	0.99	0.98	291	702	482			
RHB_MPS	1493	4.62	5.20	4.73	0.98	0.99	0.98	359	703	431			
OPB_MPS	1530	4.61	5.18	5.16	0.98	0.99	0.99	341	646	543			
					1			1					









Journal Prendicos



ounalprove



# **Highlights:**

- Biochar was applied in moderately and highly polluted soils under field • conditions.
- Biochar application increased bacterial and plant diversity in highly polluted soil. •
- Bacterial diversity slightly changed in moderately polluted soil with biochar. •
- Soil pH was key factor of bacterial community structure. •
- Seasonal changes affected microbial community, soil respiration and • vegetation.

#### **Declaration of interests**

 $\boxtimes$  The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: