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1 **Physiological and Molecular responses of flax (*Linum usitatissimum* L.) cultivars under a**
2 **multicontaminated technosol amended with biochar**

3
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12 **Abstract**

13 Soil pollution is a worldwide issue and has a strong impact on ecosystems. Metal(loid)s have toxic effects
14 on plants and affect various plant life traits. That is why metal(loid) polluted soils need to be remediated.
15 As a remediation solution, phytoremediation, which uses plants to reduce the toxicity and risk of polluted
16 soils, has been proposed. Moreover, flax (*Linum usitatissimum* L.) has been suggested as a potential
17 phytoremediation plant, due to its antioxidant systems, which can lower the production of reactive oxygen
18 species and can also chelate metal(loid)s. However, the high metal(loid) toxicity associated with the low
19 fertility of the polluted soils render vegetation difficult to establish. Therefore, amendments, such as
20 biochar, need to be applied to improve soil conditions and immobilize metal(loid)s. Here, we analyzed the
21 growth parameters and oxidative stress biomarkers (ROS production, membrane lipid peroxidation,
22 protein carbonylation and 8-oxoGuanine formation) of five different flax cultivars when grown on a real
23 contaminated soil condition, and in the presence of a biochar amendment. Significant correlations were

24 observed between plant growth, tolerance to oxidative stress, and reprogramming of phytochemical
25 accumulation. A clear genotype-dependent response to metal(loid) stress was observed. It was
26 demonstrated that some phenylpropanoids such as benzoic acid, caffeic acid, lariciresinol, and kaempferol
27 played a key role in the tolerance to the metal(loid)-induced oxidative stress. According to these results, it
28 appeared that some flax genotypes, *i.e.* Angora and Baikal, could be well adapted for the
29 phytoremediation of metal(loid) polluted soils as a consequence of their adaptation to oxidative stress.

30 **Highlights**

31 Flax cultivars were used in a biochar assisted phytoremediation process
32 Flax plants were able to grow on the amended contaminated mine soil
33 Cultivars showed different responses to metal(loid)-induced oxidative stress
34 The cultivar Eurodor was sensitive whereas Angora and Baikal were tolerant

35

36 **Keywords**

37 Flax; Soil pollution; Metal/Metalloid; Technosol; Biochar; Oxidative stress makers

38

39 1. Introduction

40 Nowadays, soil pollution is a worldwide issue. The discharge of wastes into the environment by
41 industries, as well as mining activities, and the use of fertilizers in agriculture, have led to multiple and
42 highly contaminated areas (Ahmad et al., 2015). In Europe, there are 2.8 million sites where polluting
43 activities took place or are taking place (Paya-Perez and Rodriguez-Eugenio 2018). Among pollutants
44 encountered, metal(loid)s are the most abundant ones. In addition to their abundance, metal(loid)s cannot
45 be degraded, and therefore accumulate in the environment, and the majority are toxic to plants and
46 humans (Ahmad et al., 2015).

47 Metal(loid)s have toxic effects on plants, they can affect the permeability of the cell membrane, the
48 biochemical activities and also reduce the growth and reproduction of the cells (Taran et al. 2020). In
49 particular, metal(loid)s can shift the balance of free-radical metabolism, which can result in oxidative
50 stress (Bajguz, 2011). The formation of reactive oxygen species (ROS), which is one of the first signs of
51 toxicity under stress conditions, can be counteracted by both enzymatic and non-enzymatic antioxidant
52 defense systems in plants (Raja et al., 2017; Jaskulak et al., 2018). Both antioxidant defense systems may
53 be outpaced by the ROS production, leading to cellular damage such as DNA mutation, protein oxidation
54 or lipid membrane peroxidation (Briffa et al. 2020), which are considered to be markers of oxidative
55 stress-induced toxicity in non-tolerant species (Posmyk et al., 2009; Wang et al., 2008). In addition, other
56 markers such as chlorophyll, flavonoid and/or phenolic contents may also provide information on the plant
57 adaptation to this abiotic stress (Esteban et al., 2008). To illustrate this point, Singh et al. (2020) grew *S.*
58 *polyrhiza* plants in a hydroponic experiment and showed that it had an important biochemical strategy to
59 cope with the toxicity induced by copper (Cu) and mercury (Hg). As antioxidant, phenolic compounds
60 (*e.g.*, flavonoids, lignans or hydroxycinnamic acids) may be involved in both ROS scavenging and
61 metal(loid) chelation (Dresler et al., 2017b). For example, chamomile roots under Cu and cadmium (Cd)
62 stress have shown increased caffeic, ferulic and *p*-coumaric acid content (Kováčik et al., 2009; Kováčik
63 and Klejdus, 2008). Interestingly, *p*-coumaric acid is a known precursor to flavonoids, which are

64 antioxidants, while ferulic acid is a precursor to lignans, known as metal chelators. Indeed, Fucassi et al.,
65 (2014) have shown that the lignan from flax, secoisolariresinol diglucoside, is capable of chelating many
66 metals, such as Cu^{2+} , Pb^{2+} , Fe^{2+} , Ni^{2+} and Ag^+ , with different affinities. However, few studies exist on the
67 interaction between flax and metal(loid)s in real soil conditions.

68 Therefore, the ability of a plant to cope with oxidative stress is of primary importance for its adaptation
69 and survival during a stress period (Verma and Dubey, 2003). Indeed, Dazy et al., (2008) studied three
70 plant species growing on both non-contaminated and contaminated areas, and concluded that a plant
71 species which exhibited an efficient defense system could cope with pollution, and therefore grow on
72 polluted soils. Similarly, Dresler et al., (2017a) compared two ecotypes of *Echium vulgare*, metallicolous
73 (M) and non-metallicolous (NM), and found that the M population presented a higher ability to respond to
74 the contamination than the NM population through a more efficient up-regulation of secondary
75 metabolites. Such studies show that plant metabolism has an important role in stress tolerance.

76 Moreover, pollution of soils induces a loss of biodiversity and its ecosystemic functions. These bare
77 contaminated soils thus present a risk of wind erosion as well as water leaching and run-off. Therefore, the
78 contamination can spread to non-contaminated environments, and possibly agricultural areas. Toth et al.,
79 (2016) analyzed soil samples all over the European Union and found that 6.24 % of agricultural lands
80 needed local assessment and may require remediation, which corresponded to 137 000 km² of agricultural
81 land. Consequently, crop culture on polluted soils can induce the transfer of the pollution into the food
82 chain (Kabata-Pendias, 2004), thereby impacting human health. For this reason, it is important to prevent
83 the spreading of contamination, and to remediate and valorize such contaminated soils. One remediation
84 solution which has been gathering attention over the last decades is phytoremediation, which uses plants
85 and their associated microbiota to reduce the toxic potential of contaminants (Gómez et al., 2019). In the
86 phytoremediation process, it is important to select the right plant species that can tolerate metal(loid)s
87 present on the site and produce a sufficient biomass. Flax (*Linum usitatissimum* L.) was demonstrated to
88 possess a tolerance towards diverse metal(loid)s, and more particularly Cd (Douchiche et al., 2010;

89 Smykalova et al., 2010). Moreover, flax plants produce a lignan, secoisolariciresinol diglucoside, that can
90 chelate metal(loid)s (Fucassi et al., 2014). Therefore, flax could have potential in the phytoremediation of
91 metal(loid) polluted soils (Angelova et al., 2004). In addition, the plantation of flax on polluted soils will
92 allow the vegetation and valorization of such areas, and preserve agricultural soils. Flax is well known for
93 producing fiber, which can be integrated into the production of biomaterials. Therefore, this study tested
94 several flax cultivars, representative of the existing flax diversity.

95 However, plant establishment on contaminated soils is often difficult due to the unfavorable conditions
96 (*i.e.* acidic pH, low nutrient quantity and availability) (Alvarenga et al., 2014; Lebrun et al., 2017).
97 Therefore, amendments have to be applied. In terms of the choice of amendment, biochar has gathered
98 attention in the last few years for its use in both uncontaminated agricultural soils and contaminated sites
99 (Barrow, 2012). Biochar is a carbon-rich, porous product obtained from the pyrolysis of organic materials
100 under low oxygen conditions and at relatively high temperatures (Barrow, 2012; Paz-Ferreiro et al., 2014).
101 Biochar is usually characterized by an alkaline pH, a high cation exchange capacity (CEC), an elevated
102 porosity, and a large specific surface area (Paz-Ferreiro et al., 2014; Tan et al., 2017). Several studies have
103 shown the positive effects of biochar application on soil properties: increase in pH, electrical conductivity
104 (EC), CEC, organic carbon content, water holding capacity (WHC) and nutrient concentrations and
105 availabilities (Forján et al., 2016; Herath et al., 2015; Janus et al., 2015), associated with a decrease in the
106 soil bulk density (Janus et al., 2015). Biochar can also be efficient in decreasing metal(loid) concentration
107 and mobility (Houben et al., 2013), due to its sorption capacity (Wiszniewska et al., 2016; Zhang et al.,
108 2017). These soil condition ameliorations were shown to improve the general plant growth (Wiszniewska
109 et al., 2016; Lebrun et al., 2017, 2018a, 2020; Herath et al., 2015).

110 To the best of our knowledge, no studies have analyzed the stress biomarkers in a real contaminated soil
111 condition and in presence of amendments. Moreover, no studies compared metal(loid) tolerance and
112 accumulation of different flax cultivars when grown on a biochar amended contaminated soil. Therefore,
113 the aim of this study was firstly to evaluate the effect of a biochar amendment on (i) the soil physico-

114 chemical properties, (ii) flax growth and metal(loid) accumulation and (iii) biochemical profiles (oxidative
115 stress biomarkers). Secondly, the study goal was to compare five flax cultivars in terms of their growth,
116 metal(loid) accumulation and biochemical profiles. Finally, the ultimate goal was to understand and
117 explain the difference in metal(loid) tolerance and accumulation of the flax cultivar by their ability to cope
118 with oxidative stress.

119 **2. Materials and Methods**

120 *2.1. Soil collection site*

121 A former mine extraction site was studied. This mine, located at Roure-les-Rosiers (St Pierre le Chastel),
122 belonged to the Pontgibaud mining district (Puy-de-Dôme, France), one of the most important mining
123 districts in Europe during the nineteenth century. The mining activities, which ended in 1897, led to
124 hundreds of tons of sandy technosol highly contaminated by arsenic (539 mg.kg^{-1}) and lead (11453 mg.kg^{-1}).
125 The soil has been characterized in previous studies, which showed that it had an acidic pH (pH 4.6)
126 (Lebrun et al. 2017), low organic matter content (2.6 %) (Lebrun et al. 2019) and low nutrient content and
127 availability (465 mg.kg^{-1} organic phosphorus, 6 mg.kg^{-1} available phosphorus and 75 mg.kg^{-1} nitrogen)
128 (Lebrun et al. 2020). An analysis of ten of the most found metal(loid)s in soil revealed that As and Pb
129 were the two pollutants above permissible limits (Lebrun et al. 2020).

130 *2.2. Compost and biochar used*

131 A commercial compost was used as a control. The biochar used as an amendment was provided by La
132 Carbonerie (Crissey, France). It was obtained from the slow pyrolysis of lightwood biomass (birch
133 biomass) under the following parameters: heating rate of $2.5 \text{ }^{\circ}\text{C.min}^{-1}$, pyrolysis temperature of $500 \text{ }^{\circ}\text{C}$
134 and residence time of 3 h. The pyrolysis product was then sieved to obtain a particle size between 0.2 and
135 0.4 mm. Biochar characteristics and methods used were described in a previous paper (Lebrun et al.,
136 2018a). All amendment properties are presented in Table 1.

137 *2.3. Plant*

138 Different cultivars of flax (*L. usitatissimum*) were used in this study: Angora, Baladin, Baikal, Drakkar
139 and Eurodor. Seeds were provided by Laboulet Semences (Airaines, France), Coopérative Linière Terre
140 de Lin (Saint-Pierre-le-Viger, France), and Arvalis-Institut Technique du Lin (Boigneville, France). These
141 cultivars were selected because they represent the flax diversity found in France and could thus give
142 information on the mechanisms of tolerance, through the oxidative stress response, and of how different
143 cultivars have contrasting toxicity responses towards metal(loid) contaminated soils.

144 *2.4. Treatments and growth conditions*

145 Three different treatments were prepared: (i) a commercial compost used as a “non-contaminated control”,
146 named C, (ii) Pontgibaud technosol, collected between 0 and 20 cm depth and sieved at 2 mm, named
147 P0% and (iii) Pontgibaud technosol amended with 5 % biochar, named PB5%.

148 Mixtures were put into 16 plastic pots (13*13*12.5 cm) per treatment. Twenty-five seeds of each flax
149 cultivar were sown in three pots per treatment, and one pot was left unvegetated. Flax growth was
150 conducted for 21 days under greenhouse conditions (temperature 22 ± 2 °C, light intensity $800 \text{ mmol.m}^{-2}.\text{s}^{-1}$,
151 photoperiod 16 h), and irrigation was provided every two days based on the water lost through
152 evapotranspiration. The 21 days of growing was chosen in order to evaluate, in the middle of the growing
153 phase, the roots and shoots answer of flax plants to metal(loid) pollution and amendments, before the
154 remobilization of resources during the flowering phase comes to disturb this partition.

155 *2.5. Soil pore water (SPW) sampling and analysis*

156 The soil pore water sampling was performed in three pots from each treatment before the seed sowing
157 (T0), using soil moisture samplers (Rhizon) (model MOM, Rhizosphere Research Products, Wageningen,
158 The Netherlands) as described in Lebrun et al., (2017). SPW was used directly (i) to measure pH using a
159 pHmeter (FE20/EL20, Metler-Toledo AG 2007) and electrical conductivity (EC) using a conductivity

160 meter (CDM210, MeterLab), as well as (ii) for performing a toxicity test using *Photobacterium*
161 *phosphoreum* (Qu et al., 2013). The toxicity test was performed in microplates, by adding 50 μ L of
162 phosphate buffer (100 mM, pH 7.0) to 50 μ L of SPW sample, followed by an addition of 50 μ L of
163 bacterial suspension (absorbance at 600 nm adjusted to 1). The plate was then covered and incubated at
164 room temperature. After 30 min, luminescence was measured with a luminometer (PolarStar Omega,
165 BMG Labtech). The relative toxicity was calculated using the control (C) as a reference and as follows:
166 $\text{relative luminescence (\%)} = \frac{\text{luminescence}_{\text{sample}}}{\text{luminescence}_{\text{control}}} \times 100$.

167 In addition, As and Pb concentrations were measured after sample acidification, using Inductively
168 Coupled Plasma Atomic Emission Spectroscopy (ICP-AES) (ULTIMA 2, HORIBA, Labcompare, San
169 Francisco, USA).

170 2.6. Soil collection and analysis

171 At harvest time, soil was collected from both the vegetated and non-vegetated pots. Soil from the three
172 vegetated pots of each cultivar (rhizosphere soil) was collected by shaking the roots inside a sterile bag
173 (Lebrun et al., 2019). The soils were air-dried prior to a microtoxicity test (adapted from Kołtowski et al.,
174 2017): 3 g of dried soil was mixed with 30 mL NaCl 2% and agitated at ambient temperature. After 10
175 minutes, the mixtures were centrifuged (10 minutes, 500 rpm) and the supernatants were collected and
176 used for the toxicity test, following the same procedure as for the SPW toxicity assessment. The relative
177 toxicity was calculated using non-vegetated compost (C) as a reference.

178 2.7. Plant harvest

179 After 21 days of growth, almost no growth was observed on the P0% substrate, as shown in figures 1 and
180 S1, and consequently, no further analyses were performed. On the other two treatments (C and PB5%), all
181 seedlings were harvested and counted. The seedlings were rinsed twice in tap water and once in distilled
182 water to remove any soil particles. Five seedlings were separated into above- and belowground parts and

183 immediately frozen and stored at -80 °C until further analysis. The other seedlings were grouped into lots
184 (4-6 plants).

185 2.8. *Physiological analyses*

186 Above- and belowground parts of each lot were separated, and then dried at 60 °C for 72 h in order to
187 determine dry weight (Lebrun et al., 2017). As and Pb concentrations were determined using ICP-AES,
188 and As and Pb mineralomass was calculated according to Lebrun et al. (2017): 0.2 g of sample was mixed
189 with 6 mL HNO₃ 65% and 3 mL HCl 35%, and digested in a microwave (program: 15 min temperature
190 increase up to 180 °C, 15 min at 180 °C and 15 min cool down); digested solution samples were recovered
191 and diluted to 30 mL with distilled water.

192 Finally, three indices were calculated based on the SPW and plant metal(loid) concentrations.
193 Translocation factor (TF) (equation 1) was the ratio of root metal(loid) concentrations to shoot metal(loid)
194 concentration.

$$195 \quad (1) \quad TF = \frac{\text{shoot metal(loid) concentration}}{\text{root metal(loid) concentration}}$$

196 Bioconcentration factor (BF) (equation 2) was the ratio of root metal(loid) concentration to SPW
197 metal(loid) concentration, while the biological accumulation factor (BAF) (equation 3) was the ratio of
198 shoot metal(loid) concentration to SPW metal(loid) concentration.

$$(2) \quad BF = \frac{\text{root metal(loid) concentration}}{\text{SPW metal(loid) concentration}}$$

$$(3) \quad BAF = \frac{\text{shoot metal(loid) concentration}}{\text{SPW metal(loid) concentration}}$$

199 2.9. *Phytochemical analysis*

200 2.9.1. *Extraction*

201 The frozen-dried vegetal material was used for these analyses. *L. usitatissimum* above and belowground
202 extracts were obtained by grinding 5 mg of material (using a balance precise at 0.1 mg) in 1 mL of 50 %
203 (v/v) aqueous ethanol (HPLC grade solvents, Thermo Scientific) using ultrasound-assisted extraction in a
204 USC1200TH (Prolabo) ultrasonic bath operating at a 30 kHz frequency, and a temperature set at 45 °C for
205 60 min. After extraction, the extract was centrifuged for 15 min at 5000 x g and the supernatant extract
206 was filtered through 0.45 µm of nylon syringe membranes.

207

208 2.9.2. Determination of total phenolic and total flavonoid content

209 The total phenolic content was analyzed using the Folin-Ciocalteu (Sigma) reagent method and the total
210 flavonoid content was determined using the AlCl₃ colorimetric method as described by Lopez et al.
211 (2016).

212 For total phenolic content (TPC), 20 µL of flax extract, 90 µL Na₂CO₃ and 90 µL Folin-Ciocalteu reagent
213 (Sigma-Aldrich) were mixed and incubated for 15 min at room temperature (25 ± 2 °C). The absorbance
214 at 630 nm was measured and the TPC was expressed in mg.g⁻¹ dry weight (DW) of gallic acid equivalent,
215 using a 5-point calibration curve (0–40 µg/mL; gallic acid; R² = 0.998).

216 For total flavonoid content (TFC), 20 µL of flax extract and 180 µL AlCl₃ (5% w/v prepared in methanol)
217 were mixed and incubated for 30 min at room temperature (25 ± 2 °C). The absorbance at 415 nm was
218 measured and the TFC was determined in mg.g⁻¹ DW quercetin equivalent using a 5-point calibration
219 curve (0–40 µg/mL; quercetin; R² = 0.998).

220 2.9.3. HPLC analysis

221 The quantification of compounds was carried out using a Varian high-performance liquid
222 chromatographic (HPLC) system as described by Corbin et al. (2015). The HPLC system is composed of:
223 a Varian Prostar 230 pump, a Metachem Degasit, Varian Prostar 410 autosampler, and a Varian Prostar

224 335 Photodiode Array Detector (PAD). Analysis was driven using Galaxie version 1.9.3.2 software. The
225 separation was performed at 35 °C using a Purospher (Merck) RP-18 column (250 x 4.0 mm i.d.; 5 µm)
226 with the detection set at 280 nm. The mobile phase was composed of solvent A (0.2% (v/v) acetic acid in
227 HPLC grade water) and solvent B (HPLC grade methanol). A nonlinear gradient was applied for the
228 mobile phase variation with a flow rate of 0.8 ml/min as follows: from 0 to 40 min of A–B: 90:10 (v/v) to
229 30:70 (v/v), from 41 to 50 min of A–B: 30:70 (v/v) to 0:100 (v/v), and A–B: 0:100 (v/v) from 51 to 60
230 min. Each compound was identified by comparison of its retention times and UV spectrum to those of
231 authentic standard. Quantification was performed using calibration curves of each standard ranging from
232 0.0125 to 0.5 mg/ml, with a correlation coefficient of at least 0.999, and using *o*-coumaric acid as an
233 internal standard (add at a 0.05 mg/mL final concentration in the extract prior extraction).

234 *2.10. Antioxidant free radical scavenging activity*

235 The antioxidant free radical scavenging activity (FRSA) of flax extracts was evaluated by their ability to
236 scavenge the DPPH (2,2-diphenyl-1-picrylhydrazyl) radical as described by Lopez et al. (2016). For this,
237 20 µL of extract sample were mixed with 180 µL of 6.10⁻⁵ M DPPH solution in a 96-well microplate. The
238 plate was then placed in the dark for 1 h at room temperature (25 ± 2 °C). The absorbance at 515 nm was
239 determined using a microplate reader. Extraction solvent (HPLC grade 50 % (v/v) aqueous ethanol
240 solution) was used as a blank. The formula used for calculating the FRSA was: FRSA (%) = 100 × (1 –
241 AE/AB), with AE the mixture absorbance at 515 nm, while AB represents the absorbance of the blank.

242 *2.11. Oxidative stress markers analysis*

243 *2.11.1. ROS quantification*

244 ROS (hydrogen peroxide) content was estimated as described in Hano et al. (2008) by using fluorescent
245 dihydrofluorescein diacetate. The fluorescence intensity was measured with VersaFluor fluorimeter
246 (Biorad) using λ_{ex} = 490 nm and λ_{em} = 514 nm.

247 2.11.2. Membrane lipid peroxidation

248 Membrane lipid peroxidation was conducted as described in Hano et al. (2008). A thiobarbituric acid
249 (TBA) reactive substances (TBARS) assay was employed to determine the membrane lipid peroxidation
250 level. A sample fraction (75 μ L) was then mixed with 25 μ L of SDS (3 % (w/v)), 50 μ L of TBA (3 %
251 (w/v) in a 50 mM NaOH solution) and 50 μ L of HCl (23 % (v/v)) with vigorous mixing after each
252 addition. The mixture was incubated at 80 °C for 20 min and then cooled on ice. The TBARS value was
253 determined by measuring absorbance at 532 nm, and subtracting non-specific absorbance at 600 nm using
254 UV-Vis spectrophotometer (Varian).

255 2.11.3. Protein carbonyl content

256 Total proteins were extracted as described in Hano et al. (2008), and their carbonylation content was
257 determined using the ELISA method (OxiSelect™ Protein Carbonyl ELISA Kit, Cell Biolabs). Total
258 protein content was determined using the Quant-iT Protein Assay Kit (Invitrogen) using the Qubit
259 fluorometer (Invitrogen). The protein carbonylation level was determined as described by the
260 manufacturer's instructions, by measuring the absorbance value at 405 nm, and the relative protein
261 carbonylation level was expressed as a percentage relative to the A_{405} value measured for the control.

262 2.11.4. DNA 8-oxo-Guanine content

263 DNA was extracted as described in Hano et al. (2008) using cetyl-trimethyl-ammonium bromide (CTAB),
264 and the 8-oxoGuanine level was determined using the ELISA method (Oxiselect oxidative DNA damage
265 ELISA kit, Cell Biolabs). DNA content was determined using the Quant-iT DNA BR Assay Kit
266 (Invitrogen) using the Qubit fluorometer (Invitrogen). The 8-oxoGuanine level was determined using the
267 ELISA method (Oxiselect oxidative DNA damage ELISA kit, Cell Biolabs, San Diego, CA, USA) as
268 described by the manufacturer's instructions, by measuring the absorbance value at 405 nm, and the
269 relative protein carbonylation level was expressed as a percentage relative to the A_{405} value measured for
270 the control.

271 2.12. Statistical analyses

272 The data regarding soil, soil pore water and plant physiological parameters were analyzed using the R
273 software (R Development Core Team, 2009). Shapiro followed by Bartlett tests were used to assess the
274 normality and homoscedasticity of the data. Then, for each treatment, the cultivar effect was analyzed by
275 comparing the means using an Anova or Kruskal test, depending on the result from the previous tests,
276 followed by a post-hoc test (TukeyHSD test and Pairwise-Wilcox test, respectively). The results of the
277 Anova and Kruskal tests to assess cultivar and treatment effects are shown in Table S1. Finally, for each
278 cultivar, the treatment effect was studied using the Student test, for normal data, or Wilcox test, for non-
279 normal data. Differences were considered significant when $p < 0.05$. For the phytochemical analysis, data
280 were the results of two biological and two technical replicates, therefore only a comparison between the
281 two treatments (C and PB5%) was performed for each cultivar using Student tests (XL-STAT 2019). For a
282 better discrimination of the cultivars and the treatments, Pearson correlations were performed using R
283 software version 3.0.2 and a PCA was performed with PAST software, with each parameter considered as
284 a discrete variable. The initial dataset was then converted into principal components (PCs), and it was
285 possible to graphically display the relationships among the considered parameters. HCA was visualized
286 with MeV4 software.

287 3. Results and Discussion

288 3.1. Physico-chemical characterization of the soil pore water (SPW) properties

289 Soil pore water was sampled at the beginning of the experiment in the three conditions. The pH
290 measurements showed that control SPW presented a pH around neutrality (6.25) (Table 2). However,
291 SPW of the contaminated soil P0% presented an acidic pH (4.23), which increased by 2.41 units when 5
292 % biochar was added. Similar results were found by Oustriere et al., (2016), with two commercial
293 biochars (prepared from pine bark and chicken manure) applied to a Cu-contaminated topsoil. Similarly,
294 Beesley et al., (2010) amended a multi-contaminated soil (As, Cd, Cu, Pb, Zn) with hardwood biochar and

295 observed a 2.1 unit pH increase. Finally, 5% *Miscanthus* biochar application increased the pH of an acidic
296 mine soil (Novak et al., 2018). This acidity correction can be attributed to the alkaline nature of the
297 biochar (pH 9.56) (Masulili et al., 2010) and to a decrease in competition between H⁺ ions and metal ions
298 for cation exchange sites (Lomaglio et al., 2017).

299 The SPW electrical conductivity followed the same trend as the pH (Table 2): the control condition had a
300 high EC while the contaminated soil showed a low EC, at 274 $\mu\text{S}\cdot\text{cm}^{-1}$. Biochar addition increased SPW
301 EC by almost five times, which is consistent with previous works (Albuquerque et al., 2014; Nigussie et
302 al., 2011). However, SPW EC was lower for PB5% than the control. Three mechanisms could explain
303 such an EC rise with biochar: (i) the ashes contained in the biochar (Fellet et al., 2014), (ii) the
304 enhancement of nutrient leaching into the soil solution (Lomaglio et al., 2017), and (iii) the dissolution of
305 soluble salts and cations present on the biochar's surface (Chintala et al., 2014). SPW As concentrations
306 were really low for the three conditions, below 0.03 $\text{mg}\cdot\text{L}^{-1}$ (Table 2). No difference in As concentrations
307 was found between the three conditions. A non-biochar effect on SPW As concentration has already been
308 observed when applying 10 % orchard biochar to a multi-contaminated mine soil (Beesley et al., 2014),
309 and could be due to the limited sorption ability of biochar towards As anions (Wang et al., 2015).
310 Regarding SPW Pb concentrations, a high concentration was found in the non-amended condition (9.32
311 $\text{mg}\cdot\text{L}^{-1}$) (Table 2). Biochar application to P0% induced an 86 % decrease in Pb concentration, up to levels
312 similar to the ones observed in the control. Such results have been observed previously and explained by:
313 (i) a pH increase (Houben et al., 2013; Mahar et al., 2015), and (ii) Pb sorption onto the biochar surface
314 (Lebrun et al., 2018b; Shen et al., 2015).

315 *3.2. Soil and soil pore water toxicity*

316 At the beginning of the experiment, substrate toxicity was assessed for the soil pore water samples. The
317 results were expressed in relative toxicity compared to the control: samples showing a value inferior to 1
318 (value of the non-vegetated compost) were considered more toxic than the control while samples with a
319 value above 1 were less toxic. The results showed that P0% was more toxic than C (Table 3). However,

320 biochar application decreased the toxicity. Indeed, SPW toxicity values of PB5% and C were not
321 statistically different. This toxicity reduction was in accordance with the study of Rees et al., (2017),
322 which showed a decrease in the potential genotoxicity of contaminated soils after biochar addition. In this
323 case, the toxicity reduction can be associated with the decrease in SPW Pb concentrations (Sáez et al.,
324 2016).

325 At the end of the experiment time course, substrate toxicity was measured for the soil samples, as SPW
326 could not be sampled. Relative toxicity was expressed using the non-vegetated compost as a reference
327 (value 1). On the soil C, Baikal and Eurodor growth increased the soil toxicity, by 2-fold on average
328 (Table 3), compared to the non-vegetated condition. On the PB5% substrate, all five cultivars reduced the
329 soil toxicity compared to the non-vegetated condition, with Angora having the highest effect. When
330 Baikal, Baladin and Eurodor seedlings were grown, there was no significant difference in soil toxicity
331 between the compost and the amended-contaminated soil. However, after Angora and Drakkar growth, the
332 two soil conditions were significantly different. These differences observed among the treatments and the
333 cultivars can be attributed to a plant root effect. Indeed, Punz and Sieghart, (1993) demonstrated that in
334 response to metal stress, plants were inducing: (i) biochemical and enzymatic changes on the root surface,
335 (ii) extracellular deposition, and (iii) binding to the cell wall components. The difference between
336 cultivars at the phytochemical level will be explored in the next sections. In addition, plants can exude
337 low molecular weight organic acids that can chelate metals, thus reducing their toxicity (Dong et al., 2008;
338 Montiel-Rozas et al., 2016). Those root excretions have been shown to depend on species and growth
339 conditions (Kidd et al., 2009). Finally, in the non-vegetated condition, both metal(loid) contaminated soils,
340 P0% and PB5%, presented a higher toxicity than the control, due to the soil contamination. However,
341 biochar had no effect on the soil toxicity.

342 *3.3. Plant growth parameters*

343 On the contaminated condition P0% and for all the five cultivars, the seedling growth was not sufficient to
344 allow harvest and further analysis (Figures 1 and S1). The application of 5 % biochar to the contaminated

345 Pontgibaud soil increased seedling growth for all five cultivars (Figures 1 and S1): on this treatment, flax
346 plants produced between 5 and 11 mg of aerial tissues and between 2 and 3 mg of roots. Likewise, Yue et
347 al., (2017) observed an increase in dry matter production when using sewage sludge biochar and a mix of
348 grass seeds. Such growth amelioration can be attributed to: (i) an increase in WHC, organic matter content
349 and nutrient content and bioavailability (Agegnehu et al., 2015; Lebrun et al., 2019); (ii) a decrease in
350 metal(loid) availability (Puga et al., 2015), and (iii) a general improvement in the soil conditions
351 (Agegnehu et al., 2016), *i.e* increase in pH, EC and decrease in metal(loid) SPW concentrations, as shown
352 in Table 1.

353 When comparing the control and the amended-contaminated soil PB5%, two contrasting behaviors were
354 observed, depending on the organs (Figure 2). Firstly, the shoot dry weights were two to three times lower
355 when plants were grown on PB5% compared to C, for the five cultivars. This decline in shoot DW on
356 PB5% could be due to soil toxicity, as demonstrated by the toxicity test performed on the substrates, as
357 well as metal(loid) toxicity, when metal(loid)s are transported into the aerial parts. Secondly, contrary to
358 the shoot, the root DW was increased, by two to five-folds, in the amended Pontgibaud soil compared to
359 the compost. This is consistent with the rhizobox experiment performed by Rees et al., (2016). Several
360 mechanisms can explain such an improvement. Firstly, the biochar-induced decrease in bulk density
361 lowers soil resistance, which promotes root development and growth (Albuquerque et al., 2014; Rees et
362 al., 2016). Secondly, biochar amendment can decrease or increase nutrient availability and increase water
363 availability (Rees et al., 2016; Xiang et al., 2017). Such amelioration of the soil conditions improved root
364 growth, due to the direct contact with the soil, whereas the aerial parts were less positively affected,
365 probably due to a lower translocation of nutrients associated with a translocation of metal(loid)s.

366 *3.4. As and Pb above- and belowground concentrations*

367 For the five cultivars and both elements, As and Pb, higher above- and belowground concentrations were
368 found when seedlings were grown on the amended contaminated soil compared to the control, which was
369 consistent with the metal(loid) concentrations found in the polluted soil (Figure 3).

370 Additionally, on the contaminated soil PB5%, above- and belowground concentrations were different
371 depending on the flax cultivar: the lowest aerial and root As and Pb concentrations were found in the
372 Drakkar cultivar, while the Baladin cultivar presented the highest concentrations. In more details, on
373 PB5% Drakkar accumulated 129 mg.kg^{-1} As and 962 mg.kg^{-1} Pb in its aerial parts, whereas root
374 concentrations were 918 mg.kg^{-1} As and $7,949 \text{ mg.kg}^{-1}$; Baladin plants presented 241 mg.kg^{-1} As and
375 $2,096 \text{ mg.kg}^{-1}$ Pb in its aerial parts and $3,308 \text{ mg.kg}^{-1}$ and $26,138 \text{ mg.kg}^{-1}$ Pb in its roots (Figure 3). These
376 results demonstrated the diverse accumulation ability among the flax cultivars. As and Pb quantities
377 (mineralomasses) were calculated based on DW and metal(loid) concentrations. It showed that Baladin
378 was the cultivar with the highest As quantity in both shoots ($1.7 \mu\text{g}$) and roots ($8.1 \mu\text{g}$) (Figure 4A), while
379 Drakkar had the lowest (0.9 and $2.2 \mu\text{g}$, respectively). Regarding Pb accumulation, aerial quantity was the
380 highest in Baikal ($20.2 \mu\text{g}$), followed by Baladin ($14.8 \mu\text{g}$), Eurodor ($10.6 \mu\text{g}$) and Angora ($7.8 \mu\text{g}$), and
381 Drakkar ($6.9 \mu\text{g}$). However, Baladin harbored the highest root Pb quantity ($64.2 \mu\text{g}$), followed by Baikal
382 ($39.0 \mu\text{g}$), Eurodor ($36.0 \mu\text{g}$) and Angora ($32.4 \mu\text{g}$), and finally Drakkar ($18.8 \mu\text{g}$) (Figure 4B). It should
383 be noted that Drakkar presented the lowest As and Pb quantities in both plant parts, mainly due to its
384 reduced As and Pb concentrations, as DW production was similar to other cultivars.

385 A higher metal(loid) accumulation in the roots compared to the shoots has been observed in several
386 studies for diverse plant species (Beesley et al., 2013; Marchiol et al., 2007). The present study also
387 showed higher As and Pb concentrations and quantities in the roots compared to the shoots, for all the five
388 cultivars. Indeed, all the TFs were low and below 1 (Table 4), showing the poor translocation of As and
389 Pb to the aerial parts (Fayiga and Ma, 2006). Among the five cultivars, only Baladin and Eurodor differed
390 from the others, showing the lowest TF values. Moreover, all five cultivars showed a similar translocation
391 of As and Pb, except for Baikal, which presented a higher translocation of Pb than As. This testified for a
392 limited metal(loid) movement along the plant conductive system (Angelova et al., 2004). Such results can
393 be explained by the high toxicity of the metal(loid)s and their low solubility once inside the root cells
394 (Tanhan et al., 2007). In addition, in general, plants can limit the translocation of toxic elements in order

395 to reduce the adverse effects on the biological processes. Indeed, the Casparian strip in the endodermis
396 acts as a barrier (Pourrut et al., 2011), and therefore metal(loid)s are compartmentalized in the roots
397 (Douchiche et al., 2010). Moreover, the main accumulation of metal(loid)s in the roots, associated to their
398 reduced translocation in the aerial part, can indicate the suitability of all five cultivar for phytostabilization
399 over phytoextraction process.

400 BF values (Table 4) were the highest in the case of Baladin for both metal(loid)s (Table 4). BF_{As} was
401 similar for Baikal and Eurodor, which presented higher values than Angora and Drakkar. However, BF_{Pb}
402 values were similar for the four other cultivars and were all lower than Baladin.

403 BAF values (Table 4) showed a different pattern. In the case of arsenic, two groups can be formed:
404 Drakkar and Eurodor presented lower BAF_{As} values than Angora, Baikal and Baladin. For Pb, again
405 Drakkar and Eurodor showed the lowest BAF_{Pb} values, followed by Baladin and Angora, and finally
406 Baikal, which presented the highest value. Moreover, BF and BAF values were very high, due to the fact
407 that these indices were calculated using SPW values on PB5%, and not soil, and that biochar greatly
408 decreased metal(loid) mobility. Therefore, even though BF and BAF values are very high, in this case,
409 flax cannot be considered as a hyperaccumulator due to its low TF values and low concentrations in plant
410 tissues compared to soil concentrations usually found in Pontgibaud soil, around $11,000 \text{ mg.kg}^{-1}$ Pb and
411 500 mg.kg^{-1} As (Lebrun et al., 2019, 2017). However, such indices showed that plants still accumulated
412 As and Pb in concentrations much higher than SPW concentrations. There are two possible explanations
413 for these results: (i) Pontgibaud soil was still able to re-supply As and Pb to the soil, which was directly
414 taken up by the plants, and/or (ii) flax plants were able to uptake metal(loid)s, even when bound to soil
415 particles. Finally, based on these indices, Baladin seems a better choice for phytostabilization as it showed
416 the highest As and Pb root accumulation (BF values) associated to the lowest translocation to upper parts
417 (TF values).

418 *3.5. Oxidative stress and its consequences on membrane lipids, proteins and DNA integrity*

419 The relative quantification of ROS (hydrogen peroxide) revealed an induction of oxidative stress observed
420 as a result of the metal(loid) pollution of each flax cultivar (Figure 5B). Measured oxidative stress was not
421 only localized to the root part, which was directly in contact with the metal(loid)s, but also affected the
422 aerial part of the plants, most likely due to As and Pb translocations from the roots to the aerial parts. The
423 production of hydrogen peroxide, for all flax cultivars, was two- to seven-times higher in the presence of
424 metal(loid)s than under control conditions. No major difference was observed between accumulations in
425 root vs aerial tissue, except for Eurodor and Drakkar. The highest level of ROS production was noted in
426 Eurodor and Drakkar cultivars. Additionally, for these two cultivars, the production of ROS in aerial
427 tissue was more intense than in root tissue. Interestingly, these two cultivars also exhibited a low
428 accumulation capacity of As and Pb. The toxicity of metal(loid)s has been related to their ability to induce
429 oxidative stress in many plant species (Dutta et al., 2018; Štolfa et al., 2015). The higher oxidative stress
430 measured for these two flax cultivars could reflect their lower tolerance to the metal(loid) stress. As a
431 result of this increased ROS production, increases in membrane lipid peroxidation, protein carbonylation
432 and 8-oxo-guanine DNA levels were observed (Figure 5C, D, E). These different oxidative stress markers
433 confirmed that the oxidative stress induced by metal(loid)s was more severe for Eurodor and Drakkar
434 cultivars.

435 Increased membrane lipid peroxidation has been reported in different species of plants as a consequence
436 of metal(loid) stress (Kumari et al., 2018; Mubarak et al., 2016). The reduction in membrane lipid
437 peroxidation appears to be an important parameter for plant adaptation to polluted soils, and also for plant
438 metal extraction capacity. In fact, this feature has been associated with higher plant tolerance and growth
439 in polluted soils as well as higher metal accumulation capacity in plant tissues (Kumari et al., 2018;
440 Mubarak et al., 2016). For example, maintenance of membrane integrity has been reported as one of the
441 specific characteristics of the Cd hyper-accumulator tolerant species *Brassica juncea* compared to the non-
442 tolerant non-hyper-accumulator *Brassica napus* (Mourato et al., 2015). Membrane integrity maintenance
443 also allows the tolerant species to maintain ion homeostasis through ion efflux mechanisms that prevent

444 and/or reduce the entry of toxic metal ions into the cells (Kumar and Trivedi, 2016). Several efficient
445 efflux transporter proteins have been identified for some tolerant plant species (Kumar and Trivedi, 2016).
446 Oxidative damage at protein level could therefore hinder this detoxification process and reduce plant
447 tolerance to metal(loid) stress (Bhagyawant et al., 2019; Kumar and Trivedi, 2016). Translocation from
448 the root to the aerial part is another aspect that could also be altered by protein oxidative damages,
449 affecting important carriers and/or channels. Plant exposure to heavy metal(loid)s also resulted in DNA
450 damage, indicating the possible genotoxic effect of metal stress that could lead to cell death (Bhagyawant
451 et al., 2019; Imtiaz et al., 2016). In addition to the classical DNA breaks already reported, here, we
452 reported the formation of 8-oxo-guanine, one of the major mutagenic DNA damages caused by
453 metal(loid)-induced oxidative stress. The highest accumulations of As and Pb were observed in Baikal and
454 Baladin flax cultivars, which were the most efficient in circumventing ROS production and ROS damage.
455 These characteristics may be interpreted as a sign of tolerance of these cultivars (Baikal and Baladin). On
456 the contrary, the highest ROS production and oxidative damages were observed with Eurodor and Drakkar
457 cultivars, which failed to accumulate high amounts of metal(loid)s, and can therefore be considered as
458 sensitive. The third strategy of the Angora cultivar could be considered stress avoidance, possibly relying
459 on efficient ion efflux mechanisms to prevent and/or reduce the entry of toxic metal ions into the cells,
460 thus limiting the production of ROS and related oxidative damage. Antioxidant response leading to ROS
461 scavenging is a way of avoiding the toxic effects of metal(loid)s in the cells. To do so, plants have
462 developed a set of antioxidant defense systems relying on enzymatic and/or non-enzymatic mechanisms
463 (Dutta et al., 2018; Štolfa et al., 2015). Our next goal was therefore to evaluate the non-enzymatic
464 antioxidant system, which has been less studied in most plant species and has not been studied in flax to
465 date.

466 *3.6. Accumulation of phenolic compounds and antioxidant response*

467 Among antioxidant compounds, the low molecular weight antioxidants such as the phenolic substances
468 are considered water-soluble. In particular, flavonoids accumulate in the vacuole and are direct scavengers

469 of H₂O₂, singlet O₂ and radical OH (Kolupaev et al., 2020). Flavonoids can trap free radicals and chelate
470 metal ions; all flavonoids have been found to be more or less involved in the antioxidant defense system
471 (Kolupaev et al., 2020). The trends of total phenolic contents (TPC), total flavonoid contents (TFC) and
472 free radical scavenging activities (FRSA) were similar for the five flax cultivars (Figures 5A and 6A, B).
473 For each analyzed tissue, metal(loid) stress resulted in increases in TPCs (2- to 3-fold) and FRSA (2 to 5
474 folds) in the five cultivars. Angora and Baikal cultivars presented the highest increases in TPC and FRSA.
475 TPC and FRSA were higher in roots than in aerial parts of Eurodor and Drakkar cultivars, while no
476 significant differences were noted with the three other flax cultivars. TFCs also increased in response to
477 metal(loid) stress but showed some variations in response intensity (Figure 6B). Angora and Baikal
478 showed the highest increase in TFC, but with a distinct pattern of tissue accumulation. TFC showed a
479 preferential accumulation in root parts for all cultivars, with the exception of Baikal for which TFC was
480 higher in aerial parts.

481 Targeted HPLC analysis focusing on the major compounds accumulated in flax confirmed the effect of
482 metal stress on the activation of the phenylpropanoid pathway. Global increases, from the most abundant
483 to the least abundant, were observed for ferulic acid, lariciresinol, secoisolariciresinol, pinoresinol, *p*-
484 coumaric acid, caffeic acid, benzoic acid, quercetin and kaempferol (Figure 6).

485 Heatmap representation (Figure 7), normalized with the average accumulation of each compound in the
486 five cultivars was used to depict natural genetic differences *vs* metal stress variations among cultivars. To
487 do this, we used a hierarchical clustering analysis (HCA) using Spearman correlation coefficients to show
488 pairwise comparisons of the effect of metal stress on the phenylpropanoid accumulations of flax cultivars
489 with different genetic backgrounds. A similar ranking was observed for the two analyzed tissues (*i.e.*,
490 roots *vs* aerials). Two distinct clusters were clearly observed. The first cluster, divided into two separate
491 groups, brought together the five flax cultivars grown under control conditions as well as Eurodor under
492 metal stress. This cluster accounted for the natural genetic variations of these cultivars grown under
493 control conditions, with Angora, Eurodor and Drakkar presenting a lower accumulation of

494 phenylpropanoids, whereas Baladin and Baikal accumulated slightly higher amounts. In particular,
495 comparatively, Baladin was naturally rich in kaempferol and benzoic acid in its roots as well as in
496 lariciresinol and kaempferol in its aerial parts. Roots and aerial parts of Baikal presented high TPC under
497 normal conditions, and in particular had relative high amounts of pinoresinol and ferulic acid in its roots.
498 Eurodor grown under metal(loid) stress conditions was linked with Baladin and Baikal grown under
499 control conditions. This could be explained by the low increase in phenylpropanoid accumulation in
500 response to metal(loid) stress observed for this cultivar, particularly in aerial parts. The other cultivars
501 grown under metal(loid) stress conditions were grouped in the second cluster with 2 subgroups: Angora
502 and Baikal on the one hand, and Baladin and Drakkar on the other hand. In particular, in response to
503 metal(loid) exposure, Angora and Baikal roots accumulated higher amounts of benzoic acid, caffeic acid,
504 ferulic acid, lariciresinol and secoisolariciresinol. Under these metal(loid) stress conditions, Angora also
505 accumulated high levels of *p*-coumaric acid in its aerial parts. A marked difference in tissue distribution
506 was also observed for this cultivar in response to metal(loid) stress with a preferential accumulation of
507 ferulic acid and benzoic acid in root tissue compared to aerial parts, whereas the opposite repartition was
508 observed for *p*-coumaric acid and pinoresinol.

509 Phenolic compounds are of particular interest in the context of metal stress, not only because of their
510 inherent antioxidant capacity, but also because of their ability to chelate metals (Kumar et al., 2017).
511 Some studies have reported an induction of the phenylpropanoid pathway in response to metal stress
512 (Sharma et al., 2019). In particular following exposure to Pb, an increase in TPC in *Brassica juncea* (Kaur
513 Kohli et al., 2018; Kohli et al., 2018), as well as in ferulic acid and caffeic acid in *Prosopis farcta* (Zafari
514 et al., 2016) were reported. The response was more complex in *Zea mays*, for which the increase in TPC
515 was associated with a decrease in caffeic acid and ferulic acid (Kısa et al., 2016). Exposure to As also
516 induced an increase in TPC in *Azolla filiculoides* (Sánchez-Viveros et al., 2011). Flax is of particular
517 interest as a rich source of various classes of antioxidant phenylpropanoid-derived compounds (Garros et
518 al., 2018; Oomah, 2001), which may be involved in chelating processes, such as secoisolariciresinol,

519 ferulic acid, *p*-coumaric acid or quercetin (Fucassi et al., 2014). Here, the differential induction of
520 different classes of phenylpropanoid-derived compounds in response to metal(loid) stress in five flax
521 cultivars with a contrasting response is therefore of particular interest. Correlation analyses were used to
522 evidence possible molecular mechanisms involving these phytochemicals which could contribute to
523 metal(loid) tolerance of flax.

524 *2.7. Correlation analyses*

525 Clear discrimination between the control and metal conditions with principal component analysis (PCA)
526 accounting for 99.37% (F1 + F2) of the initial variability of the data was obtained (Figure 8, S2, S3). The
527 discrimination between the control and metal conditions occurred principally in the first dimension (F1
528 axis). This F1 axis explained 99.19% of the initial variability, with BF(As), BF(Pb), BAF(As), [As], and
529 [Pb] as the main discriminating contributor for this axis. The second dimension (F2 axis), accounting for
530 only 0.18% of the initial variability, nevertheless gave a clear separation of the different genotypes
531 according to their phytochemical accumulation and antioxidant response to metal stress. The main
532 discriminating contributors of this F2 axis were benzoic acid, caffeic acid, lariciresinol and kaempferol
533 contents as well as the FRSA. The flax genotypes grown under control conditions were circumscribed in
534 the lower left part of the ellipse whereas the genotypes under metal(loid) stress conditions were more
535 largely distributed with Drakkar and Baladin located in the bottom upper part, and Angora and Baikal in
536 the lower right part, whereas Eurodor was the only genotype located in the left part of the biplot. These
537 results confirmed the contrasting response of the 5 flax genotypes to the stress of metal(loid)s with similar
538 discrimination to the one observed on the HCA.

539 The Pearson correlation analysis study supported this pattern by connecting BF(As), BF(Pb), BAF(As),
540 [As] and [Pb], on the one hand, with the content of benzoic acid, caffeic acid, lariciresinol and
541 kaempferol, and on the other, with RSA.

542 These results demonstrated the genotype-dependent nature of flax response to metal(loid) stress and the
543 strong impact of resistance to induced oxidative stress, in particular RRSA and accumulation in stress
544 tolerance of certain essential phenylpropanoids such as benzoic acid, caffeic acid, lariciresinol and
545 kaempferol. Based on these results, the different behavior observed for the five flax cultivars indicates
546 that, as a function of this specific phytochemical accumulation capacity, certain genotypes could be well
547 adapted to the phytoremediation of metal(loid) contaminated soils: the cultivar Eurodor was the one
548 having the best growth, but was a sensitive cultivar, while Angora and Baikal were the two most tolerant
549 cultivars. Therefore, Angora and Baikal could be used in associated to biochar for the remediation of an
550 As and Pb contaminated soils.

551

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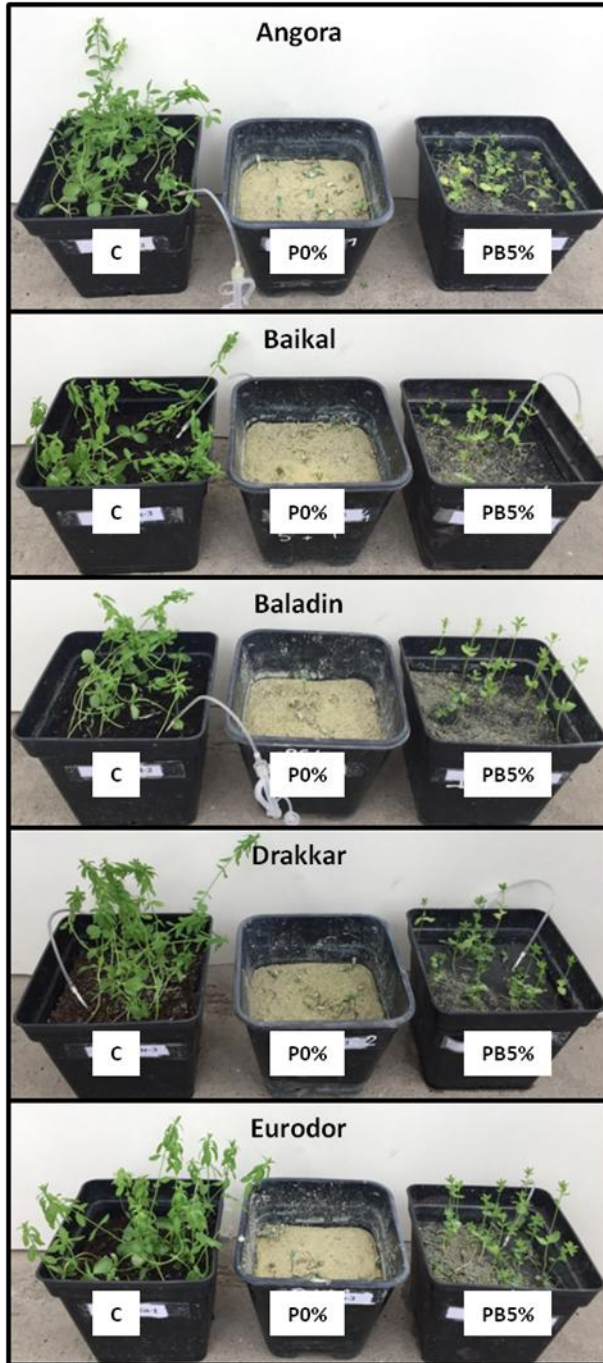
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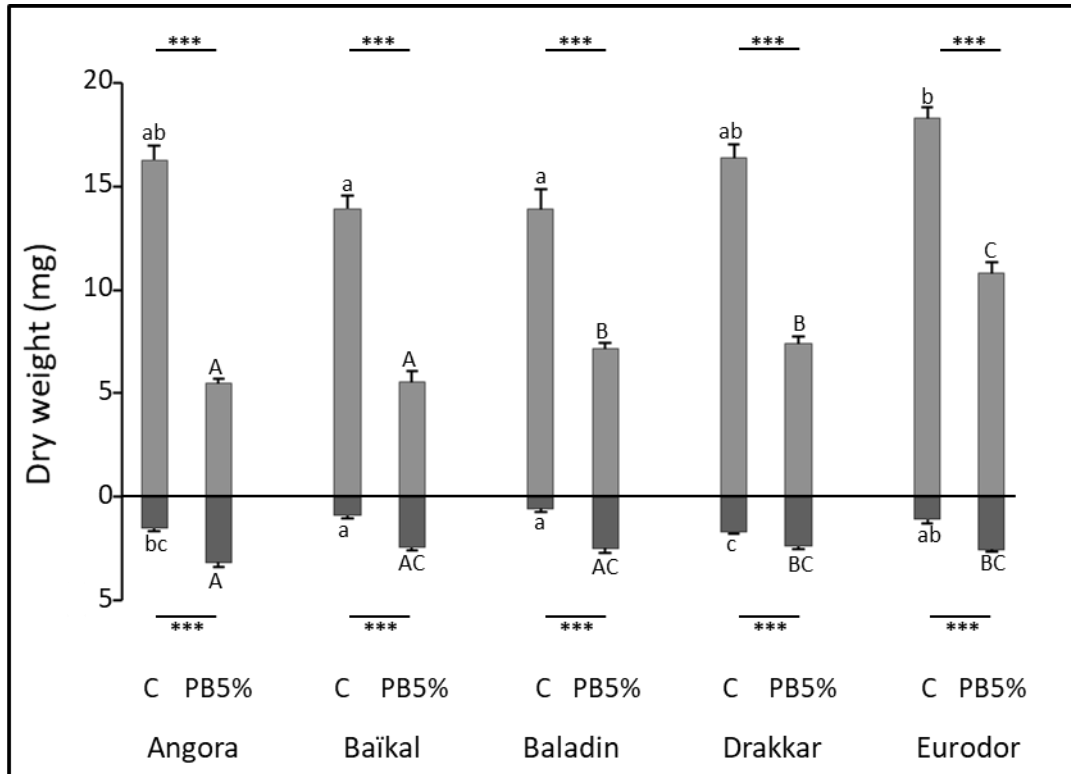
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856 Figure 1: Photographies of *Linum usitatissimum* plants in the pots (13*13*12.5 cm) after 21 days of
 857 growth on the different substrates. C = compost (non contaminated control); P0% = Pontgibaud technosol;
 858 PB5% = Pontgibaud technosol amended with 5% biochar.

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861 Figure 2: Dry weight (mg) of the aerial (light grey) and root (dark grey) parts of the 5 *Linum usitatissimum*
 862 cultivars (Angora, Baikal, Baladin, Drakkar and Eurodor) after 21 days of growth under 2 conditions. C =
 863 compost (non contaminated control); PB5% = Pontgibaud technosol amended with 5% biochar. Capital
 864 letters indicate significant difference between the cultivars in the PB5% condition whereas the small
 865 letters indicate a significant difference between the cultivars in the C condition (n = 5-13 ± SD). *** (p <
 866 0.001) indicates significant difference between the 2 conditions for one cultivar.

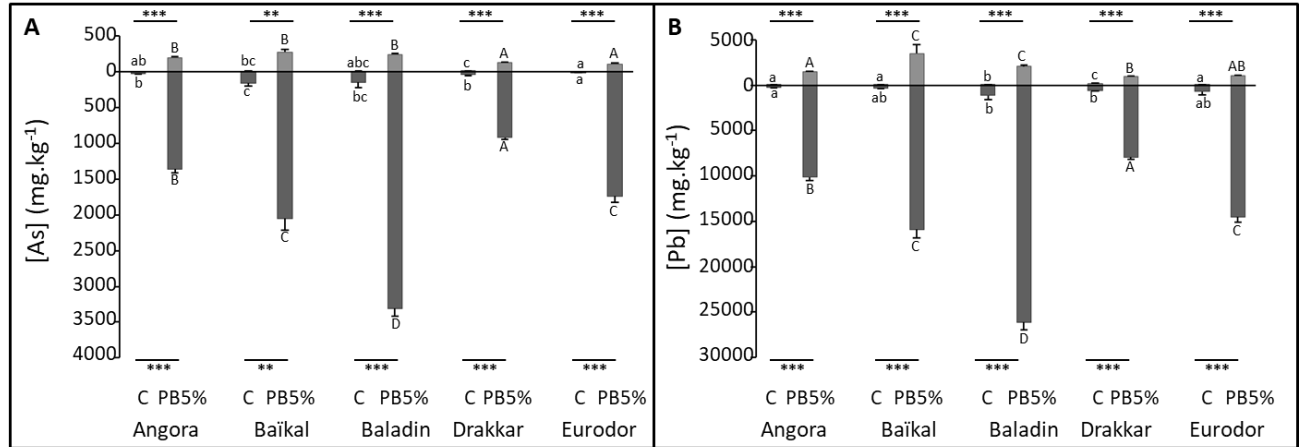
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873 Figure 3: Arsenic (A) and lead (B) concentrations (mg.kg⁻¹) of the aerial (light grey) and root (dark grey)
874 parts of the 5 *Linum usitatissimum* cultivars (Angora, Baikal, Baladin, Drakkar and Eurodor) after 21 days
875 of growth under 2 conditions. C = compost (non contaminated control); PB5% = Pontgibaud technosol
876 amended with 5% biochar. Capital letters indicate significant difference between the cultivars in the
877 PB5% condition whereas the small letters indicate a significant difference between the cultivars in the C
878 condition (n = 5-13 ± SD). ** (p < 0.01) and *** (p < 0.001) indicate significant difference between the 2
879 conditions for one cultivar.

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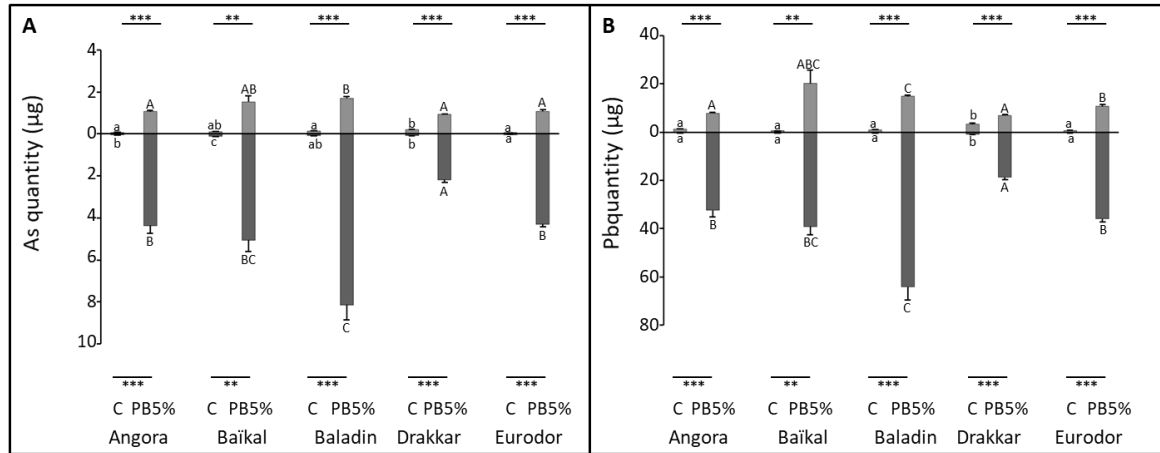
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889 Figure 4: Arsenic (A) and lead (B) quantities ($\text{mg}\cdot\text{kg}^{-1}$) of the aerial (light grey) and root (dark grey) parts
890 of the 5 *Linum usitatissimum* cultivars (Angora, Baikal, Baladin, Drakkar and Eurodor) after 21 days of
891 growth under 2 conditions. C = compost (non contaminated control); PB5% = Pontgibaud technosol
892 amended with 5% biochar. Capital letters indicate significant difference between the cultivars in the
893 PB5% condition whereas the small letters indicate a significant difference between the cultivars in the C
894 condition ($n = 5-13 \pm \text{SD}$). ** ($p < 0.01$) and *** ($p < 0.001$) indicate significant difference between the 2
895 conditions for one cultivar.

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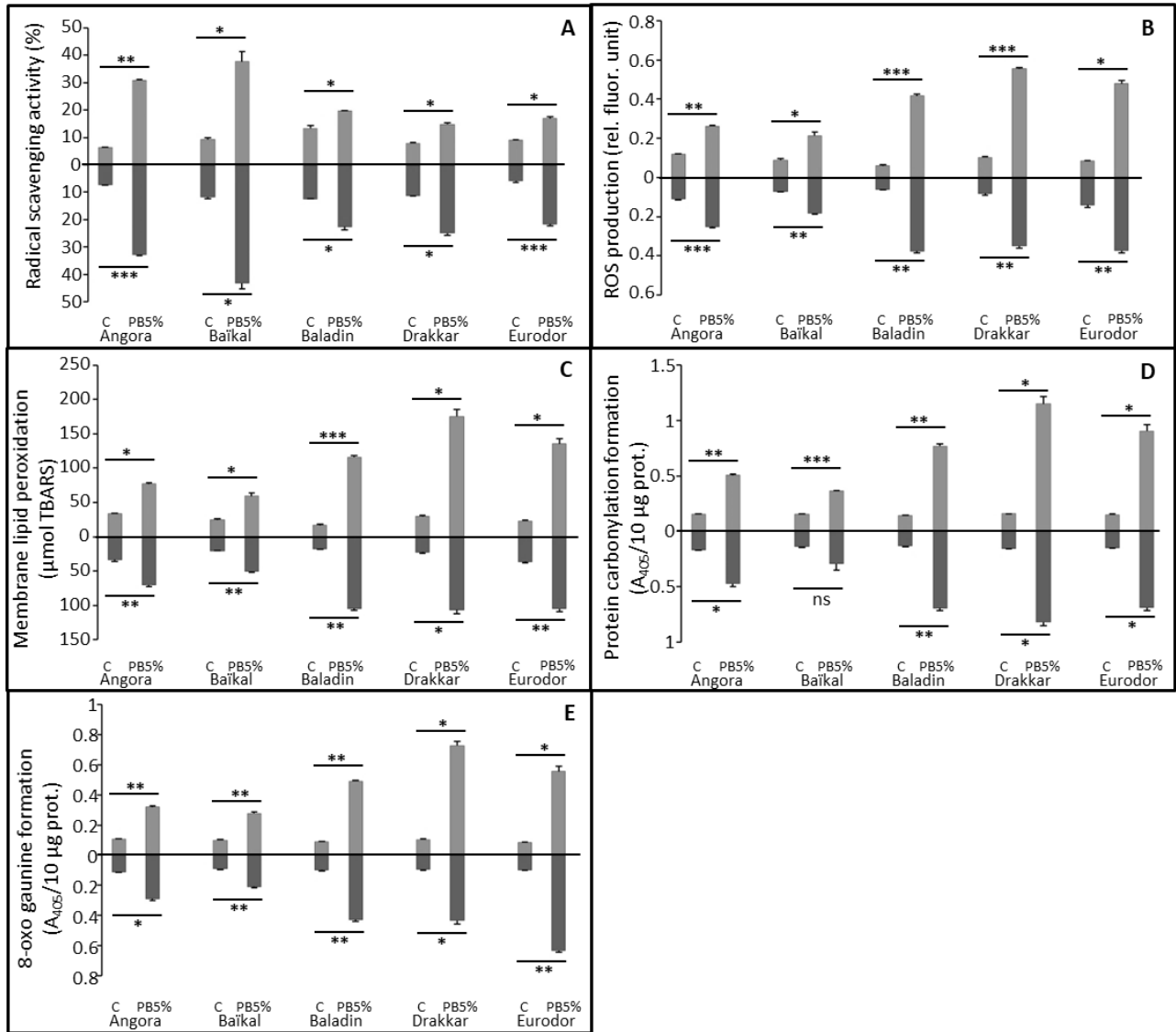
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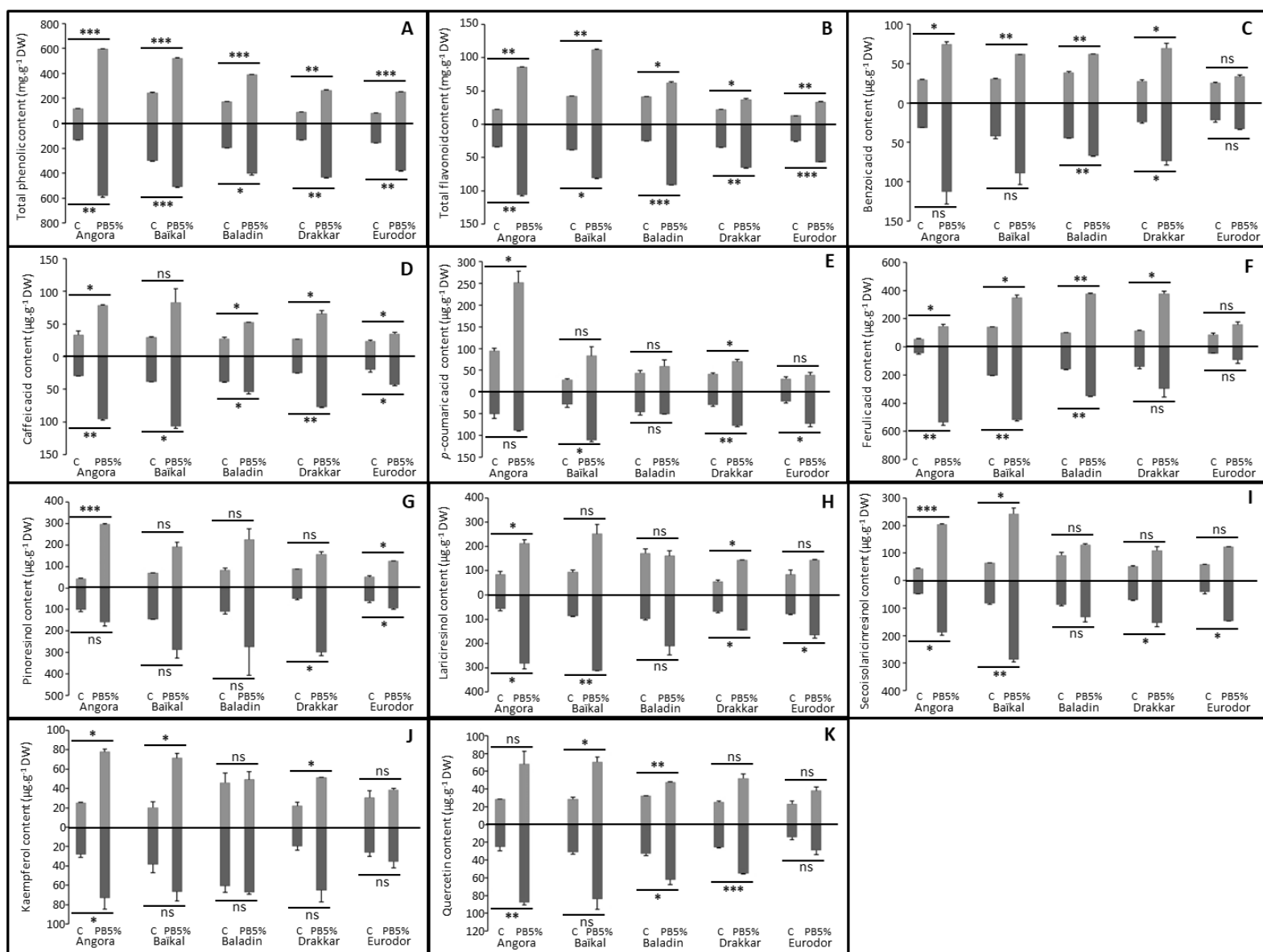
905 Figure 5: Radical scavenging activity (%) (A), reactive oxygen species (ROS) formation (relative

906 fluorescence unit) (B), membrane lipid peroxidation (µmol TBARS) (C), protein carbonylation formation

907 (A₄₀₅/10 µg protein) (D) and 8-oxo guanine formation (A₄₀₅/10 µg protein) (E) measured in the aerial

908 (light grey) and root (dark grey) parts of the 5 *Linum usitatissimum* cultivars (Angora, Baikal, Baladin,

909 Drakkar and Eurodor) after 21 days of growth under 2 conditions. C = compost (non contaminated
910 control); PB5% = Pontgibaud technosol amended with 5% biochar. Values are means \pm SD of 2
911 independent experiments (2 biological and 2 technical replicates); * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$
912 (comparison with the corresponding compost condition).



914 Figure 6: Content (mg.g⁻¹ or µg.g⁻¹ dry weight) in phytochemical compounds (total phenolic (A), total
915 flavonoid (B), benzoic acid (C), caffeic acid (D), p-coumaric acid (E), ferulic acid (F), pinoresinol (G),
916 lariciresinol (H), secoisolariciresinol (I), kaempferol (J) and quescetin (K)) of the aerial (light grey) and
917 root (dark grey) parts of the 5 *Linum usitatissimum* cultivars (Angora, Baikal, Baladin, Drakkar and
918 Eurodor) after 21 days of growth under 2 conditions. C = compost (non contaminated control); PB5% =
919 Pontgibaud technosol amended with 5% biochar. Values are means ± SD of 2 independent experiments (2
920 biological and 2 technical replicates); * $p<0.05$, ** $p<0.01$, *** $p<0.001$, ns: not significant at $p<0.05$
921 (comparison with the corresponding compost condition).

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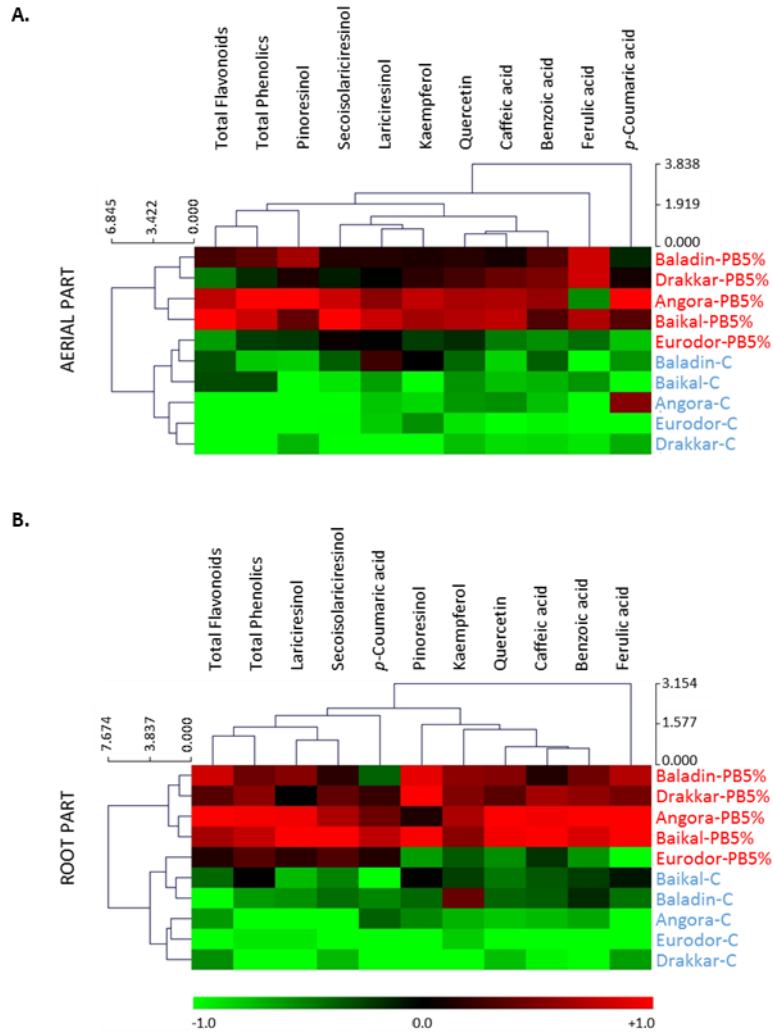
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936 Figure 7: Hierarchical clustering analysis of the phytochemical compounds measured in the aerial (A) and

937 root (B) parts of the 5 *Linum usitatissimum* cultivars (Angora, Baikal, Baladin, Drakkar and Eurodor) after

938 21 days of growth under 2 conditions. C = compost (non contaminated control); PB5% = Pontgibaud

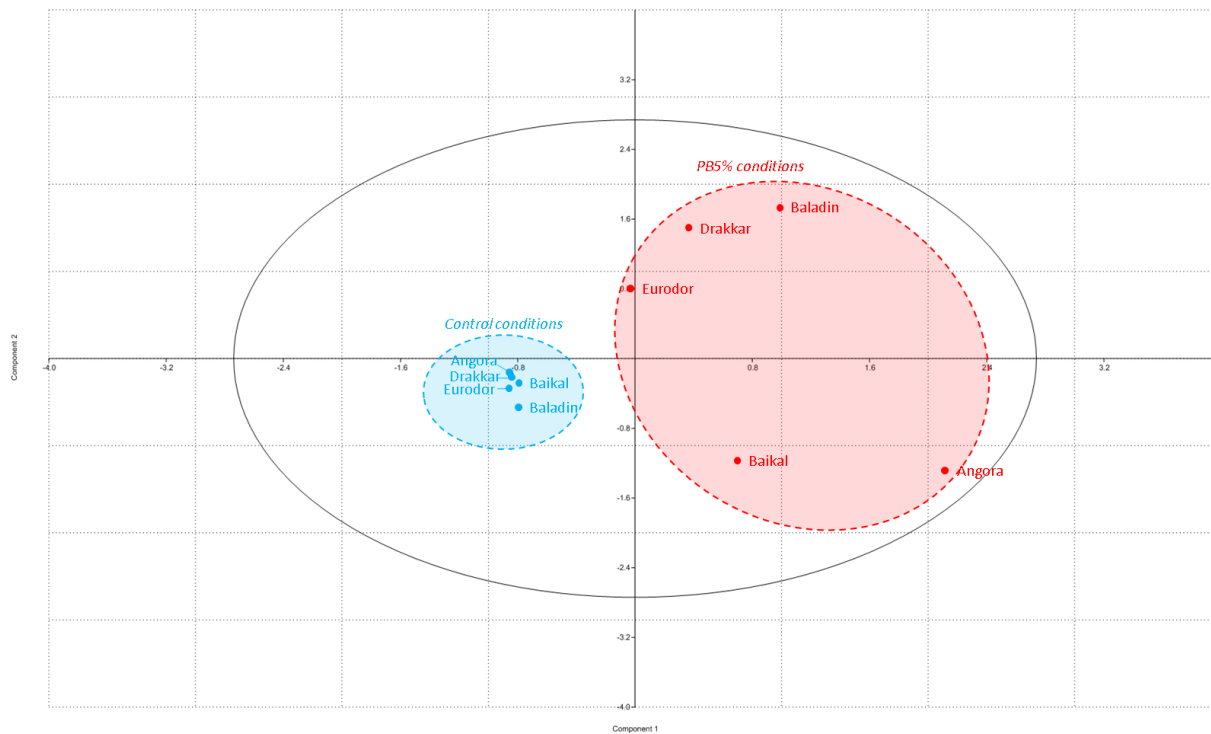
939 technosol amended with 5% biochar.

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945 Figure 8: Principal component analysis of the different parameters measured the 5 *Linum usitatissimum*
 946 cultivars (Angora, Baikal, Baladin, Drakkar and Eurodor) after 21 days of growth under 2 conditions (non
 947 amended control = blue, PB5% = pink). The Hotelling's ellipse confines the confidence region (95%) of
 948 the score plot.

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Table 1: Biochar and compost characteristics.

	Compost	Biochar
pH	6.5 ⁽¹⁾	9.56 ± 0.01 ⁽²⁾
Electrical conductivity ($\mu\text{S}\cdot\text{cm}^{-1}$)	520 ⁽¹⁾	849 ± 4 ⁽²⁾
Water holding capacity (% mass)	60 ⁽¹⁾	194 ± 2
Organic matter (% dry)	68 ⁽¹⁾	nd
Particle size (mm)	nd	0.2-0.4 ⁽¹⁾
Specific surface area ($\text{m}^2\cdot\text{g}^{-1}$)	251.42	nd
Total pore volume ($\text{m}^3\cdot\text{g}^{-1}$)	0.1486	nd
Pore diameter	2.3639	nd

(1) Data provided by the supplier

(2) From Lebrun et al. (2018a)

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Table 2: Soil pore water physico-chemical properties (pH and electrical conductivity (EC) (mg.L^{-1}) and metal(loid)s (As and Pb) concentrations (mg.L^{-1}), at the beginning of the experiment for the 3 conditions. C = compost (control); P0% = Pontgibaud technosol; PB5% = Pontgibaud technosol amended with 5% biochar. Letters indicate significant difference between the 3 treatments ($n=3 \pm \text{SD}$).

	pH	EC ($\mu\text{S.cm}^{-1}$)	[As] (mg.L^{-1})	[Pb] (mg.L^{-1})
C	6.25 ± 0.05 b	5263 ± 446 c	0.023 ± 0.014 a	0.131 ± 0.043 a
P0%	4.23 ± 0.19 a	274 ± 23 a	0.025 ± 0.002 a	9.322 ± 2.150 b
PB5%	6.64 ± 0.07 b	1302 ± 73 b	0.011 ± 0.007 a	1.304 ± 0.014 a

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Table 3: Relative toxicity of the soil pore water, determined at the beginning of the experiment (T0), and of the soil, determined at the end of the experiment (T21) . C = compost (control); P0% = Pontgibaud technosol; PB5% = Pontgibaud technosol amended with 5% biochar. NV = non-vegetated pots. Minuscule letters indicate significant difference between the 3 treatments (n=3), while capital letters indicate significant difference between cultivar (n = 6 ± SD).

Soil pore water		Soil					
T0		T21					
		NV	Angora	Baikal	Baladin	Drakkar	Eurodor
C	1 b	1 bB	1.21 ± 0.07 aB	0.53 ± 0.04 aA	0.95 ± 0.05 aB	1.50 ± 0.06 bB	0.50 ± 0.06 aA
P0%	0.79 ± 0.04 a	0.29 ± 0.02 a	-	-	-	-	-
PB5%	0.96 ± 0.05 b	0.25 ± 0.02 aA	1.46 ± 0.09 bE	0.45 ± 0.02 aBC	0.87 ± 0.02 aD	0.42 ± 0.02 aB	0.51 ± 0.02 aC

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Table 4: Translocation factor, bioconcentration factor and biological accumulation factor determined for each cultivar in the PB5% condition. Letters indicate significant difference ($p < 0.05$) ($n = 5-13 \pm SD$).

	Translocation factor (TF)		Bioconcentration factor (BF)		Biological accumulation factor (BAF)	
	As	Pb	As	Pb	As	Pb
Angora	0.15 ± 0.01 b	0.15 ± 0.01 b	135872 ± 6138 b	7809 ± 312 a	20055 ± 1768 b	1123 ± 100 ab
Baikal	0.14 ± 0.03 b	0.23 ± 0.07 b	204984 ± 18461 c	12218 ± 834 a	27367 ± 4051 b	2712 ± 855 c
Baladin	0.07 ± 0.00 a	0.08 ± 0.00 a	330842 ± 11818 d	20106 ± 703 b	24108 ± 1607 b	1612 ± 107 b
Drakkar	0.14 ± 0.01 b	0.12 ± 0.01 b	91828 ± 3443 a	6114 ± 177 a	12946 ± 886 a	740 ± 50 a
Eurodor	0.06 ± 0.01 a	0.07 ± 0.01 a	173667 ± 9088 c	11160 ± 477 a	10681 ± 1226 a	802 ± 86 a