

# Influence of agricultural amendments on arsenic biogeochemistry and phytotoxicity in a soil polluted by the destruction of arsenic-containing shells

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Revised manuscript with accepted corrections

1	Influence of agricultural amendments on arsenic biogeochemistry and							
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3	shells							
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15								
16	Abstract							
17	Agricultural soils can contain high arsenic (As) concentrations due to specific geological							
18	contexts or pollution. Fertilizer amendments could influence As speciation and mobility thus							

increasing its transfer to crops and its toxicity. In the present study, field-relevant amounts of

fertilizers were applied to soils from a cultivated field that was a former ammunition-burning

site. Potassium phosphate (KP), ammonium sulfate and organic matter (OM) were applied to

these soils in laboratory experiments to assess their impact on As leaching, bioavailability to

Lactuca sativa and microbial parameters. None of the fertilizers markedly influenced As

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speciation and mobility, although trends showed an increase of mobility with KP and a 24 decrease of mobility with ammonium sulfate. Moreover, KP induced a small increase of As in 25 Lactuca sativa, and the polluted soil amended with ammonium sulfate was significantly less 26 phytotoxic than the un-amended soil. Most probable numbers of AsIII-oxidizing microbes and 27 AsIII-oxidizing activity were strongly linked to As levels in water and soils. Ammonium 28 sulfate negatively affected AsIII-oxidizing activity in the un-polluted soil. Whereas no 29 significant effect on As speciation in water could be detected, amendments may have an 30 impact in the long term. 31

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33 Key words: arsenic, soil, water, fertilizers, microorganisms, omega 3 index

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#### 36 **1. Introduction**

High concentrations of the toxic element arsenic (As) in soils generally originate from mining and industrial activities, long-term applications of As-containing pesticides or the geochemical background. Among industrial activities, storage or destruction of As-bearing molecules used in chemical weapons during the wars has locally resulted in high As concentrations in soils (Bausinger et al., 2007; Thouin et al., 2016; Hube 2017).

When soils affected by As pollution are submitted to agricultural practices, arsenic speciation, bio-availability for plants and mobility towards the water phase may be changed. Major phenomena influencing As mobility (Smith et al., 1998) include; (1) pH which influences AsIII and AsV oxy-anions charge, (2) redox conditions, which influences As speciation and the stability of iron oxides that are essential As-bearing phases, and (3) competing substances,

that may favour As desorption from solid phases. In particular, phosphate, a structural
analogue of AsV can compete with As for sorption on iron oxides (Smith and Naidu, 2009).

Thus, in agricultural soils, fertilizing practices involving phosphate amendments may affect 49 50 As speciation and mobility. Brackhage et al. (2014) observed an increase of As mobility and uptake by wheat associated to P-fertilization in soil flooding conditions. Conversely, N-51 fertilization seemed to attenuate As mobility and plant uptake (Brackhage et al., 2014, Van 52 53 Oort et al., 2017). In addition, agricultural soils are often fertilized with organic matter (OM). Many studies have described geochemical interactions between As and organic matter: 54 modification of As speciation (Redman et al., 2002), formation of soluble complexes (Saada 55 56 et al., 2003; Redman et al., 2002), competition for sorption sites (Bauer and Blodau, 2006), and influence of OM on microbial AsIII-oxidizing activity (Lescure et al., 2016). 57

Finally, all types of amendments may impact the structure of soil microbial communities 58 which exert a major influence on As speciation (Yamamura et al., 2009). Bacteria isolated 59 60 from soils have been shown to oxidize AsIII and/or reduce AsV (Macur et al., 2004; Inskeep 61 et al., 2007; Bachate et al., 2012), or to methylate this toxic metalloid (Huang et al., 2012). Filamentous fungi isolated from contaminated soils are able to reduce AsV and methylate As 62 (Su et al., 2011). Microbial transformations of As in soil have important implications because 63 mobility, toxicity and bioavailability of this metalloid are closely related to its speciation 64 (Smedley and Kinniburgh, 2002). The global AsIII-oxidizing activity of the microflora should 65 tend to reduce the risk of As transfer from soil to surface water or groundwater. This global 66 activity is the result of AsIII-oxidation and simultaneous AsV-reduction, that can occur in 67 aerobic conditions through the activity of As resistance genes. All the modifications due to the 68 69 amendments can lead to changes in the bioavailability and the toxicity of As for crops.

This study aimed to measure the impact of fertilizing practices on As mobility, speciation and
transfer to crops in soils from a former chemical-ammunition-destruction facility dating from

the interwar period (1918-1939) and subsequently converted into agricultural land near 72 Verdun, France (Hube, 2017). It is one of the most important historical areas of chemical 73 ammunition destruction of WW I, containing arsenical chemical warfare agents, located in a 74 sensitive zone for agriculture and groundwater. Fertilizers frequently applied for common 75 crops such as barley, corn and wheat (potassium phosphate (KP) fertilizer, ammonium sulfate 76 and OM) were applied to soils in a laboratory-scale experiment to assess their impact on the 77 speciation, mobility and phytotoxicity of As as well as on As transforming microbial 78 79 communities.

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#### 81 **2.** Material and methods

#### 82 2.1. Origin and characterization of soil samples

Soils were sampled from a field near Verdun, France, where there was previously a chemical
ammunition destruction facility (Hube, 2017). The field was used as pasture in 2012 and was
cultivated from 2012 to 2015 with wheat, barley and corn. Since 2015, it is fallow ground as
farming was forbidden when the pollution was detected.

Surface soils (0-20 cm) were sampled with a spade in a highly polluted zone, and in a reference zone 25 m away. Each sample was taken as a composite of 5 points from 3 m × 3 m squares according to the GEMAS protocol (EGS 2008) but adapted to the small surfaces available. The soils were characterized (Table 1) according to the following methods: pH in water (NF ISO 10390), organic carbon (NF ISO 10694), total nitrogen (NF ISO 13878), Phosphorus (NF ISO 11263), total elements, major and trace elements (extraction with HF + HCIO<sub>4</sub>, NF ISO 14689-1).

As species in soils were analysed by HPLC-ICP-MS after extraction with 10 mL H<sub>3</sub>PO<sub>4</sub> 1 M added to 0.4 g of freeze-dried sample, ground and homogenized by sieving (2 mm) and microwave heating (Vergara Gallardo et al., 2001) in a closed system at 120°C during 20 min 97 (analyse performed by UT2A laboratory, Pau, France). The remaining solution was diluted to 98 50 mL with ultrapure water and then As species were analysed with HPLC-ICP-MS using 99 quantification by standard additions to avoid matrix effects. As species separation was 100 performed using an anion exchange column (Hamilton PRPX-100) and a mobile phase made 101 of ammonium hydrogen phosphate 15 mM at pH 8.5. These analytical conditions enable the 102 determination of AsIII, AsV and methylated As species (MMA and DMA). Details of the 103 analytical methods are provided in SM1.

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**Table 1.** Main characteristics of the two studied soils. (\*) total element.

Parameter	pН	$P_2O_5$ mg.kg <sup>-1</sup>	$K_2O mg.kg^{-1}$	Cd* mg.kg <sup>-1</sup>	Cr* mg.kg <sup>-1</sup>	Cu* mg.kg <sup>-1</sup>	Hg* mg.kg <sup>-1</sup>
Verdun Reference soil	8.3	19.7 (+/-6.7)	306 (+/-9.7)	0.4 (+/-0.01)	30.9 (+/-16)	25.7 (+/-1.5)	0.07 (+/-0.006)
Verdun Polluted soil	8.2	182 (+/-14)	678 (+/-23)	0.4 (+/-0.02)	41.4 (+/-2.9)	74.6 (+/-3)	1.1 (+/-0.2)
Parameter	Ni* mg.kg <sup>-1</sup>	Pb* mg.kg <sup>-1</sup>	Zn* mg.kg <sup>1</sup>	As* mg.kg <sup>-1</sup>	Mn* mg.kg <sup>-1</sup>	С %	N %
Verdun Reference soil	25.2 (+/-0.32)	21.9 (+/-0.05)	111.1 (+/-2.6)	21.8 (+/-0.9)	884.5 (+/-47.4)	1.9 (+/-0.7)	0.21 (+/-0.006)
Verdun Polluted soil	24.4 (+/-0.99)	45.8 (+/-10)	180.3 (+/-9.9)	983 (+/-130)	791.9 (+/-18.7)	2.8 (+/-0.17)	0.32 (+/-0.03)

Diphenylarsinic acid (DPAA) was analysed by HPLC-DAD and clark I, clark II, Clark oxide,
triphenylarsine (TPA), and 9-phenylarsafluorene were analysed by GC–MS (Envilytix,
Wiesbaden, Germany, details in SM1). They were detected only in the polluted soil that
contained 0.2% DPAA and 0.1% TPA. These two last species were not quantified in the next
steps of the study that focused on total As and bio-transformations of AsIII and AsV.

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113 2.2. Microcosm experiments

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115 2.2.1 Preparation

116 Microcosms were setup in 200 mL polystyrene pots, 50 mm diameter, whose bottoms were

117 perforated with a 0.9 mm needle, to make 13 holes in each pot. In order to retain soil particles



were calculated based on the real quantities applied on site (SM2) for KP fertilizer and for ammonium sulfate. A third amendment, organic manure, was chosen as the site was used as a pasture for many years (Table 2). The amount of manure added to the soils in the microcosms was based on real quantities usually applied in cultivated fields, i.e. 10 tons.ha<sup>-1</sup>.

**Table 2.** Amendments for microcosms. Hypothesis: Depth of amended soil 0.3 m, soil density

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1.3.

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kg.ton<sup>-1</sup> of soil Amendments Real average Mass of amendment in the quantity applied per microcosm, for **150 g** of hectare (kg) soil (mg) KP 0.07 10.5 273 95 3.6 Ammonium 0.024 sulfate 10,000 Organic 2.56 384 amendment

130

131 KP fertilizer (00 18 18) was provided by Soufflet Agri Service (Neuville-aux-Bois, France). It 132 contained 0.32 % organic C, 22.4 % K<sub>2</sub>O, 18.9 % P<sub>2</sub>O<sub>5</sub>, and less than 0.1 g. kg<sup>-1</sup> of total N. 133 Fertilizer granules were crushed to a powder before application (< 100  $\mu$ m). Biomarine 134 organic amendment (NF U 44-051 provided by Truffaut, France) was used as an organic 135 amendment. It is composed a mixture of vegetal and animal wastes (horse manure, sheep 136 manure, poultry manure, algae, grape marc), containing 33.9 % organic C, 31.8 g.kg<sup>-1</sup> total N,

- 137 4 % K and 0.6 % P. The two solid amendments (KP and manure) were added as powders to
- the dry soil and mixed for 24 h by rotation to homogenize. Ammonium sulfate was added as a
- 139  $2.4 \text{ g.L}^{-1}$  concentrated solution, 1.5 mL in each microcosm during the first watering.
- 140
- 141 2.2.2. Watering and incubation

Watering was always performed with Mont Roucous mineral water (pH 5.85; 3.1 mg.L<sup>-1</sup> Na<sup>2+</sup>; 2.4 mg.L<sup>-1</sup> Ca<sup>2+</sup>; 0.5 mg.L<sup>-1</sup> Mg<sup>2+</sup>; 2.0 mg.L<sup>-1</sup> SO<sub>4</sub><sup>2-</sup>; 6.3 mg.L<sup>-1</sup> HCO<sub>3</sub><sup>-</sup>; 3 mg.L<sup>-1</sup> NO<sub>3</sub><sup>-</sup>), used to simulate rain water.

For the first watering, 53 mL of water was carefully poured on the soil surface. Then, 24 h 145 after the first watering, the microcosms were watered again, sufficiently to recover 20 mL of 146 percolated water in the underlying pot. The quantity of inlet and outlet water was recorded by 147 weighing the recipients. Global aerobic non-saturated conditions were maintained. 148 149 Microcosms were incubated at 25°C in the dark, with 80 % atmospheric humidity. Soils never dried out during the incubation. Percolated water was filtered at 0.45 µm. Determination of 150 As speciation was performed using an ion exchange method (Kim, 2001), the separated As 151 species and the total As in water were determined by oven-AAS (details in SM1). Soils were 152 watered at the beginning of experiment then after 1 week, 1 month and 3 months. 153

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155 2.2.3. Final determination of biological parameters

At the end of the experiment (3 months), soils were sampled to determine biologicalparameters. All measurements were performed in triplicate.

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159 2.2.3.1. Most Probable Number (MPN) determinations

Active As-transforming microorganisms present in microcosms were enumerated by thefollowing MPN methods. To enumerate the active AsIII-oxidizing microorganisms (Thouin et

al., 2016), wet soil (equivalent to 2.5 g dry soil) was placed in a sterile glass erlenmeyer flask 162 with 10 ml of sterile physiological water (9 g.l<sup>-1</sup> NaCl in demineralized water), agitated for 30 163 min at 25°C, then sonicated 2 x 20 s at 45 kHz. Triplicate suspensions were prepared for each 164 soil. Soil suspensions were serially diluted in sterile physiological water up to dilution 10<sup>-7</sup>. 165 CAsO1 mineral medium (Battaglia-Brunet et al., 2002) containing 100 mg.L<sup>-1</sup> AsIII was 166 distributed in Microtest TM Tissue culture plates (96 wells), 250 µL by well. Each well was 167 inoculated with 25 µL of each soil suspension dilution. Five wells were inoculated for each 168 dilution. Culture plates were incubated at 25°C for 10 days. Presence of AsIII in the wells was 169 revealed by the formation of insoluble white complex AsIII-PyrrolidineDithioCarbamate 170 (PDC): in each well 150 µL 0.1 M acetate buffer (pH 5) and 100 µL PDC solution (5 g.L<sup>-1</sup>) 171 were added. A white precipitate appeared when AsIII was present, i.e. when AsIII-oxidizing 172 bacteria were absent (negative well). Un-inoculated wells served as negative blanks, and wells 173 containing CAsO1 medium with 100 mg.L<sup>-1</sup> AsV served as a positive reference. The number 174 of positive wells for each dilution was determined, and the most probable number of bacteria 175 176 in dilutions was given by the Mc Grady table for 5 tubes.

To enumerate Active AsV-reducing microorganisms (Thouin et al., 2018), soil suspensions 177 were prepared as described for AsIII-oxidizing microorganisms then diluted in sterile 178 physiological saline solution to a dilution of 10<sup>-6</sup>. CAsO1 basal mineral medium (Battaglia-179 Brunet et al., 2002) was complemented with 20 mM lactic acid and AsV (100 mg.L<sup>-1</sup>). The 180 medium was distributed in Microtest TM Tissue culture plates (96 wells), 250 µL per well. 181 Each well was inoculated with 25 µL of diluted soil suspension. Five wells were inoculated 182 with for each dilution. Culture plates were incubated at 25°C for 10 days in anaerobic jars 183 with Anaerocult packs (Merck). Presence of AsIII formed in the wells during incubation was 184 revealed as described above. Positive well numbers were determined for each dilution, and the 185

most probable number of AsV-reducing microorganisms was given by the Mc Grady table forfive tubes.

188 2.2.3.2. Activity tests

AsIII-oxidizing tests were performed in 250 mL Erlenmeyer flasks filled with 100 mL of 189 CAsO1 medium (Battaglia-Brunet et al., 2002) supplemented with 100 mg.L<sup>-1</sup>/ AsIII and 190 inoculated with soil (equivalent to 0.2 g dry weight). Flasks were incubated at 25°C in 191 192 oxidizing conditions under agitation (100 rpm). AsV-reducing tests were performed in 250 mL serum flasks filled with 100 mL of CAsO1 medium supplemented with 20 mM lactic 193 acid, 0.2 g.L<sup>-1</sup> yeast extract and AsV (100 mg.L<sup>-1</sup>). Flasks were inoculated with soil 194 (equivalent to 0.2 g of dry weight). Flasks were hermetically closed, flushed with N<sub>2</sub>, and 195 incubated at 25°C in static conditions. Flasks were sampled every day in order to monitor the 196 197 evolution of AsV concentrations: 5 mL of culture were filtered at 45 µm with cellulose acetate filters and frozen at -20°C until AsIII/AsV separation with the PDC/MIBK method (Battaglia-198 199 Brunet et al., 2002, details in SM1), As in the AsV-containing aqueous phase was quantified 200 by flame AAS (Varian, Palo Alto, CA, USA). AsIII-oxidation and AsV-reduction rates were calculated between each point of analyse (evolution of AsV concentration divided by the time 201 between sampling events). 202

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#### 204 2.3. Toxicity and transfer to plants

Impact of fertilizers on As bioavailability and phytoxicity was evaluated with the ecotoxicological test AFNOR XP X 31-233 (2012), using the Omega-3 Index based on the analysis of leaf fatty acid composition of *Lactuca sativa* grown under controlled conditions and by measuring the As leaf content in the same plants. Fifteen *Lactuca sativa* seeds were sown in plastic pots filled with 430 g of dry soil, with or without amendment, and control pots containing a standard soil (loam). Each condition was tested in triplicate using the same

quantities of amendments as for the microcosm experiment. Amendments were added to the 211 soil at the beginning of the experiment and seedlings were grown for 17 days in a growth 212 chamber under a 16 h/8 h photoperiod (8000 lx or 10000 lx white light intensity) and a 213 22°C/16°C day/night temperature. Pot location inside the growth chamber was randomly 214 changed every 2 to 3 days to homogenize light exposure and watering. One week after 215 germination, germination rates were determined and the number of young seedlings per pot 216 was reduced. 14 days after germination, the aerial parts of seedlings were weighed, and the 217 first leaf of some plants (or a section of it: 20 to 200 mg of fresh tissue) used to determine the 218 leaf fatty acid composition, was placed in a glass tube containing 1 mL of a methanol/H<sub>2</sub>SO<sub>4</sub> 219 solution. Determination of leaf fatty acid composition was performed as described in Le 220 Guédard et al. (2008). Determination of As concentrations was carried out by harvesting all 221 the plants in each pot used for measuring the leaf fatty acid content. Plants were thoroughly 222 223 washed in tap water and rinsed three times with deionised water. Plant biomass was then dried at 40 °C to constant weight and ground in a plastic bag to approximately 2 mm, to facilitate 224 225 the digestion step. Samples (1 g Dry Weight) were digested in clean, dry PTFE screw cap 226 vessels in hot concentrated HNO<sub>3</sub>, according to Zarcinas et al. (1987) and As concentrations in the extracts were measured by ICP-MS (details in SM1). 227

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#### 229 **2.4. Statistical analysis**

Statistical analysis were performed with XLSTAT 2019.3.2.61397. Significance of differences between results of As concentrations, most probable number of bacteria and Omega-3 index were evaluated using the non-parametrical Kruskal and Wallis test, with multiple pairwise comparison using Dunn's procedure, two tailed test, at a significance level of 5% followed by a Dunn post hoc test. Correlations between parameters were calculated with XLSTAT 2019.3.2.61397, Pearson (n) correlation matrix, (p < 0.05). Data of maximum

AsIII oxidation and AsV reduction rates were tested for homogeneity of variance and normal distribution. One-way analysis of variance (ANOVA) and Tukey HSD (Honestly Significantly Different) tests were carried out to test for any significant differences between the means. Differences between means at the 5% level (P < 0.05) were considered significant.

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#### 241 **3. Results**

Initial soil analyses showed that phosphate concentration was ten times higher in the polluted 242 243 soil than in the reference soil, and the potassium concentration was two times higher in the polluted soil than in the reference soil (Table 1). Average concentrations of total As were 244 close to 1000 ppm in the polluted soil and to 20 ppm in the reference soil. Concentrations of 245 246 total As in the percolated water were significantly lower for the reference soil microcosms (1 to 6  $\mu$ g.L<sup>-1</sup>, Figure 1A) compared to polluted soil microcosms (2000 to 5000  $\mu$ g.L<sup>-1</sup>, Figure 247 1B). Although the ratio polluted/reference for total As was close to 50X in the solids, it was in 248 the range of 1000X in the percolated water. This indicated that As was much more mobile in 249 the polluted soil than in the reference soil. An increase of leached As was linked to the 250 addition of KP fertilizer and a decrease of this leached As was linked to the addition of 251 ammonium sulfate amendment in the polluted soil, at the first watering event (day 0, Figure 252 1B). 253

Cumulated amounts of total As leached from the soils (Figure 2) indicated that total As leached from the polluted soil was 1000X higher than the cumulated As leached from the reference soil. None of the amendments significantly increased or reduced As leaching compared with the blank experiment. A slight increase could be linked to the addition of KP fertilizer in the polluted soil and the lowest values of leached As were obtained with the ammonium sulfate amendment (Figure 2A and 2B).

Speciation (quantification of AsIII and AsV) was monitored in water from microcosms 260 containing the polluted soils only, as As concentrations were too low in the reference soil 261 percolation water. AsIII concentrations of AsIII varied between 5 and 15 µg.L<sup>-1</sup> in these 262 leachates, representing 0.1 to 0.5 % of total As. Total AsIII leached during the experiment 263 (Figure 2C) ranged from 700 to 1000 ng. These amounts of AsIII represent less than 1 % of 264 the leached As, however they are higher than the total amounts of As leached from the 265 reference soil. The amount of total leached AsIII was slightly higher with the KP amendment. 266 However, the difference with other conditions is not statistically significant. Globally, the 267 amendments had no significant influence on As speciation in the leached water during this 268 experiment. 269

Final pH values (SM3) were close to the initial values, the general tendency being a small decrease during the experiment. None of the amendments induced an increase in pH sufficient to mobilize As.

Numbers of AsIII-oxidizing microorganisms were in the range of 10<sup>4</sup> cells.g<sup>-1</sup> for the 273 reference soil, and 10<sup>6</sup> cells.g<sup>-1</sup> for the polluted soil (Figure 3A), thus 100 times higher in the 274 polluted soil. Numbers of AsV-reducing microorganisms were in the range of 10<sup>6</sup> cells.g<sup>-1</sup> 275 whatever the soil type (Figure 3B). Globally cell numbers were slightly lower in the reference 276 soil compared to the polluted soil. However, this difference was less marked than it was for 277 AsIII-oxidizing microorganisms. Non-parametric statistical tests (XLSTAT 2018.2.50583 -278 Kruskal-Wallis test Two-sample t-test and z-test), performed as the comparison of all values 279 with reference soil and all values with polluted soil indicates that the two groups (a and b on 280 Figure 3) are significantly different for both AsIII-oxidizing and AsV-reducing bacteria. 281 However, no significant influence of the different amendments on the MPN of As-282 transforming microorganisms could be detected. 283

Results of the AsIII-oxidizing activity tests (Figure 4A) clearly showed two trends: one 284 observed with the group of polluted soil conditions, whose microbial communities oxidized 285 AsIII very rapidly, and a second trend observed with the reference soils where kinetics were 286 slower and clearly affected by the amendments. The maximum AsIII-oxidation rate was 287 significantly higher with the polluted than with the reference soil. Considering exclusively the 288 reference soil, the ammonium-amended soil gave a value of maximum AsIII-oxidation rate 289 significantly lower than the non-amended soil, whereas the KP and organic amendments 290 tended to decrease the AsIII-oxidizing rate but not significantly. 291

Results of the AsV-reducing activity tests (Figure 4B) indicated a rapid complete reduction ofAsV after two days, for all soils. Thus, the AsV-reduction rate was similar for all conditions.

As contents in the lettuce leaves are shown in Figures 5A and 5B. As contents in leaves are significantly higher in lettuces grown on polluted soils (7000 to 9000  $\mu$ g.kg<sup>-1</sup> DW) than in those grown on reference soils (60 to 120  $\mu$ g.kg<sup>-1</sup> DW). As transfer factor from soil to shoot (TF<sub>shoot/soil</sub>) for polluted soil is 2.7-fold higher than for the TF<sub>shoot/soil</sub> from the reference soil. Thus, as observed with microcosm experiments, As from the polluted soil appeared more mobile and more phytoavailable than As from the reference soil.

While the phosphate concentration in the soil is 10X higher in the polluted soil compared with the reference soil, the phosphorus content in the lettuce leaves was similar for both soils (SM4). Concerning the amendment intake, results showed a significant (except for As in the leaves of plants grown on the reference soil) increase of As and P contents in lettuce leaves in both soils amended with KP (Figure 5 and SM4). The other amendments did not show any differences with the un-amended soil, regardless of the soil type (reference or polluted).

The Omega-3 Index is a standardized biomarker to evaluate the possible toxic effects of contaminants on plants (AFNOR XP X31-233, 2012). This biomarker measures the degradation of polyunsaturated fatty acids and decreases when lipid peroxidation, caused by

an excess of reactive oxygen species (ROS) in the presence of contaminants, increases. 309 Results showed that the Omega-3 Index was significantly lower for lettuces grown on the 310 polluted soil compared to the reference soil (Figure 6). This indicated that lipid peroxidation 311 was higher in lettuces grown on the polluted soil, which was therefore more phytotoxic than 312 the reference soil. This phytotoxicity of As in the polluted soil caused a decrease of the lettuce 313 seedling growth, as shown in SM5 (69.02 mg of FW.plant<sup>-1</sup>±13.73 and 30.34 mg of FW.plant<sup>-1</sup> 314  $^{1}\pm4.97$  for reference soil and the polluted soil, respectively). Moreover, while there was no 315 difference between the un-amended and amended reference soils, the Omega-3 Index 316 increased significantly for lettuces grown on polluted soil amended with ammonium sulfate. 317 318 Thus, addition of ammonium sulfate in the polluted soil seemed to reduce As induced lipid peroxidation. 319

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#### 321 **4. Discussion**

Among the parameters controlling the behaviour of As in soils, redox conditions play a major 322 role. Experiments performed with microcosm systems allowing control of redox conditions 323 showed that low Eh values induced mobilization of As from a flood plain soil (Frohne et al., 324 2011), a freshwater marsh delta soil (Shaheen et al., 2016) and an historically contaminated 325 coastal soil (LeMonte et al., 2017). Here, non-saturated conditions were maintained in 326 327 microcosms, so the mobility of As should mainly be controlled by other factors. Although AsV adsorption on iron oxides is known to decrease with pH (Dixit and Hering, 2003), As 328 mobility in the polluted soil compared to the reference soil was probably not exclusively 329 330 linked to soil pH (8.2 and 8.3 in the reference and the polluted soil, respectively, Table 1). As mobility could also be related to the higher concentration of phosphate in the polluted soil. 331 Indeed, phosphate concentration was ten times higher (and the potassium concentration two 332 times higher) in the polluted soil than in the reference soil. One hypothesis to explain this is 333

that the farmer might have provided higher quantities of KP fertilizer in this precise area in an 334 attempt to improve crop yields. Indeed plant growth was strongly affected by the undetected 335 pollution in this area of the field. Phosphate is an analogue of arsenate AsV (Smith and Naidu, 336 2009) which competes with As for adsorption sites on iron oxides. In the percolation water, 337 the highest total As concentrations were always observed with the KP fertilizer and could be 338 linked to the addition of phosphate. Consequently increased As mobility may have increased 339 the toxic impact on plant growth. In addition, as the polluted soil contained both more 340 phosphate and more As than the reference soil, the ratio between the density of adsorption 341 sites and concentrations of both elements was lower in the polluted than in the reference soil. 342 This has previously been observed in soils polluted with As, but with higher doses of 343 phosphate fertilizers (Davenport and Peryea, 1991; Peryea and Kammereck, 1997). In terms 344 of concentration, As in the microcosm' percolation water was in the range 2-5 mg.L<sup>-1</sup>, in the 345 346 same range as concentrations already reported in water phases in contact with soils polluted by different sources of As. Thus, Cao et al. (2003) found 5-6 mg.L<sup>-1</sup> of As in water leached 347 348 from soil polluted with a chromium-copper-arsenic (CCA) pesticide, presenting a total As content of 135 mg.kg<sup>-1</sup>, and Qi and Danahoe (2008) found 1 mg.L<sup>-1</sup> of As when they used 349 acid rainwater to leach a soil polluted by historical herbicide application and containing 300 350 mg.kg<sup>-1</sup> As. 351

MPN of As-transforming microbes suggested that the pollution induced an increase of the abundance of active microorganisms able to modify As speciation in the soil (Figure 3). The difference observed between the reference soil and the polluted soil was more marked for AsIII-oxidizing than for AsV-reducing microbes, and that was true both for MPN and activity tests: AsIII-oxidizing microbes were roughly 100 times more abundant in the polluted soil (Figure 3A), and this soil presented a higher AsIII-oxidizing activity than the reference soil. Thouin et al. (2016) showed that the AsIII-oxidizing rate increased with the As concentration

in samples of polluted technosoils, and suggested that the level of As concentration exerted a 359 selective pressure on the microbial community, increasing its global efficiency to oxidize 360 AsIII. Here, whereas amendments did not influence the AsIII-oxidizing activity of the 361 polluted soil (Figure 4A), the fertilizers exerted a clear influence on this activity in the 362 reference soil. AsIII-oxidizing activity was diminished by the ammonium sulfate amendment 363 (Figure 4A). Yet, this fertilizer tended to decrease As mobility from soils towards percolation 364 water, in particular at the beginning of the experiment with the polluted soil (Figure 1B). Our 365 results are in agreement with those of a long-term (1929 to 2018) experiment studying the 366 effects of agricultural amendments on the behaviour of trace elements. In that experiment, 367 authors showed that ammonium-based fertilization induces an increase of immobilized As in 368 the soil, compared with other types of amendments, due to reduced As leaching (van Oort et 369 al., 2017). This phenomenon might be linked to H<sup>+</sup> production during ammonium oxidation 370 371 (nitrification). Even if no macroscopic pH decrease was observed here, protonation of the surface hydroxyl groups of iron oxides could increase AsV adsorption on these minerals in 372 373 the soil (Dixit and Hering, 2003). Reduced As mobility from solids to the water phase might 374 decrease As bioavailability, consequently reducing the selective pressure on the global microbial community and decreasing its efficiency to oxidize AsIII. 375

The organic fertilizer did not significantly influence As behaviour nor the activity and 376 abundance of As-transforming microbes (Figures 1 to 4). In previous studies, OMwas 377 reported to induce higher As mobility in soils (Beesley et al., 2014). However, contradictory 378 results were reported about the impact of OM on As mobility (Kumpiene et al., 2008). In 379 terms of effects on microbial transformation of As, Lescure et al. (2016) showed that OM 380 exerted a positive effect on AsIII-oxidation rates from 0 to 0.08 g.L<sup>-1</sup> of organic carbon, then 381 tended to decrease the AsIII-oxidation activity at higher concentrations. Here, the dose of 382 organic fertilizer (0.26 %) was probably not sufficient to modify microbial activities. 383

The solid and liquid phases of the soils contained AsV as the main As form, whereas the 384 MPN and activity of AsV-reducing microbes was important according to Figures 3 and 4. 385 Even when As is mainly in the form of AsV, it is not a static distribution of AsV between 386 solids and water phases, but rather a dynamic equilibrium, involving both microorganisms 387 oxidizing AsIII and microorganisms reducing AsV. In this system, bioreduction of AsV might 388 play a role in As mobilisation. Turpeinen et al. (1999) have already observed that microbes 389 increased the As mobility, mainly as AsV, from soils incubated in aerated conditions. We 390 found AsV as the main As form leached from the polluted soil, with no significant effect of 391 the amendments on this speciation, in soils that were not saturated with water, i.e. with no 392 limitation of oxygen availability. Our results must be confirmed with further experiments 393 performed with diverse polluted sites, and comparing biotic to abiotic conditions, and non-394 saturated to saturated conditions. However, they suggest that microbial parameters would be 395 396 very sensitive bio-indicators of the dynamics of As concentrations and speciation in the water phase. 397

Considering the pot experiments with plants, As transfer was significantly more important in 398 lettuces grown on the polluted soil. These results showed that As was highly mobile and as a 399 400 consequence highly phytoavailable in the polluted soil. This correlated with results from the microcosms where As was shown to be more mobile from soil to water within the polluted 401 soil compared to the reference soil. In lettuces, amendment with KP induced an increase of 402 both As and P contents in lettuce leaves, as already observed by Cao and Ma (2004). 403 Phosphate-induced plant As uptake may have been related to the slight increase of As 404 concentration in the leachate. The lettuces' Omega-3 Index showed that the high As content in 405 the polluted soil increased lipid peroxidation in lettuce leaves compared to the reference soil 406 (Figure 6). Bustingorri et al. (2017) observed an increase of lipid peroxidation in soybean 407 plants linked to the presence of As. Here, the increase of lipid peroxidation induced by the 408

polluted soil, corresponding to nearly 50 % decrease of the Omega3 index (Figure 6) was coupled with a decrease of the lettuce seedling growth. Thus, in agreement with findings obtained from previous studies, the excessive production of ROS induced by As exposure promotes lipid peroxidation and causes damage in thylakoid membranes that may lower photosynthetic efficiency (Abbas et al., 2018). In agreement with previous results of As transfer from soil to lettuces (Figure 5), these results confirm that As is highly phytoavailable and as a consequence highly phytotoxic in the polluted soil.

The amounts of agricultural amendments applied to fields can influence the mobility, 416 bioavailability and toxicity of As. Addition of ammonium sulfate to polluted soil was 417 associated with an increase of the Omega-3 Index (Figure 6) and as a consequence a reduction 418 of lipid peroxidation in lettuce leaves while no decrease was observed concerning the As 419 uptake. This was not observed in the reference un-polluted soil, suggesting that the decrease 420 421 of lipid peroxidation is possibly influenced by the contribution of ammonium sulfate to As detoxification. These results are in agreement with several studies showing that, under heavy 422 423 metal stress, an application of sulfur stimulates plants (Mishra et al, 2008; Duan et al, 2013). In fact, sulfur (S) in higher plants is a vital component for the synthesis of some amino acids 424 (Cysteine and Methionine) and metabolites such as gluthatione (GSH) and phytochelatins 425 (PCs) involved in the detoxification of heavy metals. Some studies relate that As induces ROS 426 production in plants leading to lipid peroxidation (Shukla et al., 2018) and the synthesis of 427 PCs enzymatically synthesized from GSH. Then, PCs form complexes with As before its 428 sequestration into the vacuoles through ABCC transporters (Song et al., 2010). This induction 429 of the biochemical pathways of plant S metabolism increases the S requirement under As 430 stress (Leao et al., 2014; Khare et al., 2017). Therefore, the decrease of lipid peroxidation in 431 lettuces grown on the polluted soil amended with ammonium sulfate may be explained by an 432 improvement of the intracellular As detoxification processes due to an increase of the 433

synthesis of GSH and PCs linked to an increase of S assimilation by plants. Similarly, 434 nitrogen (N) could also play an important role in the detoxification of heavy metals in plants. 435 Indeed, many studies related that, under abiotic stress, nitrate supply stimulates root nitric 436 oxide production (Sun et al, 2010; Simontacchi et al., 2015) which is a bioactive signalling 437 molecule involved in plants' response to heavy metal stress by detoxifying ROS (Hassan et 438 al., 2005; Kaur et al., 2015). Thus, according to these results, it seems that a better N and S 439 nutrition status may protect the plants from As and as a consequence lead to a decrease of 440 oxidative stress. However, more research is needed to fully understand the mechanisms of 441 interactions between ammonium sulfate and As in plants and the importance of ammonium 442 sulfate in detoxification. 443

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#### 446 5. Conclusions

The addition of fertilizing amendments, at the real average dose applied on the sites, did not 447 strongly influence the speciation and the quantity of mobile As. Observed trends were an 448 increase of As mobility with KP fertilizer, a decrease of mobility with ammonium sulfate 449 amendment, and no effect of organic amendments. The quantity of As mobilised in the 450 percolation water was bioavailable for plants and soil microorganisms. Results of an original 451 combination of active As-transforming bacteria enumeration and As-related microbial activity 452 tests showed that microbial parameters were strongly linked to As levels in water and in soils. 453 In particular, microbial AsIII-oxidizing activity proved to be very sensitive to low doses of 454 ammonium sulfate. Although the impacts of KP and ammonium sulfate on As speciation were 455 insignificant, the small effects observed on plant As-uptake and bacterial activities may have 456 an impact in the long term. Consistent results were observed with microbial and plant 457 parameters, in particular concerning the effect of ammonium sulfate fertilization. These tests 458

459 could be used as sensitive indicators of As bioavailability and toxicity in soils. Whereas 460 microbial parameters can be quicker to obtain than plant indicators, As bioavailability for 461 plants will always be very important to determine, particularly when these plants are 462 cultivated and consumed.

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#### 472 **Figure captions**

**Figure 1.** Evolution of total As concentrations in the leachates. A: reference soil; B: polluted soil. B: blank experiment without amendment; KP: KP fertilizer; N: ammonium sulfate; O: organic amendment. P for polluted. Significance of differences between amendment conditions was evaluated separately for each incubation time using the non-parametric statistical method (details given in material and methods section), T0 (A, B, C); T7 days (a, b, c); T 29 days (i); T 87 days ( $\alpha$ ). Error bars represent the standard deviation of the means (3 replicates).

**Figure 2.** Cumulated leached total As and AsIII. A: reference soil total As; B: polluted soil total As; C: polluted soil AsIII. KP: KP fertilizer; N: ammonium sulfate; O: organic amendment. P for polluted. Significance of differences between amendment conditions was evaluated separately for each incubation time using a non-parametric statistical method (details in material and methods section). Error bars represent the standard deviation of the means (3 replicates).

Figure 3. Most probable numbers of As-transforming microorganisms. A: AsIII-oxidizing
microorganisms; B: AsV-reducing microorganisms. KP: KP fertilizer; N: ammonium sulfate;
O: organic amendment. P for polluted. Error bars represent the standard deviation of the
means (3 replicates). Groups A and B were statistically different according to the KruskalWallis test.

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**Figure 4.** Activity tests. A: AsIII-oxidizing activities; B: AsV-reducing activities. Error bars represent the standard deviation of the means (3 replicates). KP: KP fertilizer; N: ammonium sulfate; O: organic amendment. P for polluted. AsIII-oxidation and AsV-reduction rates were calculated between each point of analyse (evolution of AsV concentration divided by the time between sampling events). Max R: Maximum rates. Values are the means (n = 3). Values with different letters are significantly different (P < 0.05, ANOVA, Tukey-HSD).

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Figure 5. As content in the lettuce leaves grown on A: the reference soil; B: the polluted soil
(P) un-amended or amended with different amendments. KP: KP fertilizer; N: ammonium
sulfate; O: organic amendment. P for polluted. Different letters (A or B) indicate significant
differences between the different amended soils. Each value is the mean of 3 samples.

<sup>487</sup> 

**Figure 6.** Omega-3 Index measured in lettuces grown on A: the reference soil; B: the polluted soil (P), un-amended or amended with different amendments. KP: KP fertilizer; N: ammonium sulfate; O: organic amendment. P for polluted. Different letters (A, B or C) indicate significant differences between the different soils un-amended or amended. Each value is the mean of 3 samples.

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--B --KP --N --O --PB --PKP --PN --PO





