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19 Abstract

Mining activities frequently leave a legacy of residues that remain in the area for long periods 20 21 causing the pollution of surroundings. We studied on a 100 year-old mine, the behaviour of 22 potentially toxic elements (PTEs) and their ecotoxicological impact on activity and diversity 23 of microorganisms. The PTEs contamination assessment allowed the classification of the materials as highly (reference- and contaminated-samples) and very highly polluted (illegal 24 25 spill of olive mill wastes (OMW), tailings and dumps). OMW presented the lowest enzymatic activities while tailings and dumps had low dehydrogenase and arylsulfatase activities. All the 26 27 alpha diversity indices studied were negatively impacted in dumps. Tailings had lower Chao1 28 and PD whole tree values as compared to those of reference-samples. Beta diversity analysis showed similar bacterial community composition for reference- and contaminated-samples, 29 significantly differing from that of tailings and dumps. The relative abundance of 30 Gemmatimonadetes, Bacteroidetes and Verrucomicrobia was lower in OMW, tailings and 31 dumps as compared to reference-samples. Fifty-seven OTUs were selected as responsible of 32 33 the changes observed between samples. This study highlights that assessing the relationship between physicochemical properties and microbial diversity and activity gives clues about 34 ongoing regulating processes that can be helpful for stakeholders to define an appropriate 35 36 management strategy.

Keywords: Potentially toxic elements, mining soil, enzyme activities, high-throughput
sequencing, microbial ecotoxicology,

40 **1. Introduction**

Mining activities are found all over the world because they provide access to mineral 41 resources that fuel various industrial activities in both developed and in developing countries. 42 43 Although recognized of great importance for the world gross domestic product, mining operations are often viewed as an important source of pollution with negative impact on the 44 environment. During the processes of mineral extraction and preparation, large amounts of 45 46 ore wastes and debris are commonly accumulated in the proximity of the mining operation site. These materials are essentially fractured rocks and soil devoid of vegetation, 47 characterized by high concentrations of heavy metals and metalloids. Consequently, the 48 49 environment is drastically transformed in highly polluted barren areas (Martín Duque et al., 2015; Sánchez-Donoso et al., 2019), which can be toxic to human health and other life, 50 including plants and microorganisms (Giller et al., 1998; Nagajyoti et al., 2010; Tchounwou 51 et al., 2012). Furthermore, potentially toxic elements (PTEs) from these polluted areas can 52 transfer to surrounding aquatic and terrestrial compartments via leaching or runoff (Elmayel 53 54 et al., 2020; Fernández-Martínez et al., 2019; Jung and Thornton, 1996; Kisková et al., 2018), disperse in the atmosphere (Esbrí et al., 2020; Naharro et al., 2020, 2018), and indirectly 55 contribute to pollutant dissemination. The recent interest in the reclamation of abandoned 56 57 mining sites in arid and semiarid regions for agricultural purposes highlights the need to understand the biogeochemical processes contributing to soil health and fertility (Favas et al., 58 2018; Higueras et al., 2019b; Krzaklewski and Pietrzykowski, 2002; Mendez et al., 2008) 59

Microorganisms such as bacteria and fungi are key players in soil ecosystem services. They are involved in multiple geochemical cycles, influence plant growth and contribute to climate regulation and soil restoration, among others (Ayangbenro et al., 2018; Cavicchioli et al., 2019; Domeignoz-Horta et al., 2018, 2015; Van Der Heijden et al., 2008). Toxicity from heavy metals and metalloids can compromise their growth and survival, with enormous

consequences on ecosystem functioning (Bell et al., 2005; Delgado-Baquerizo et al., 2020, 65 2017, 2016b, 2016a; Domeignoz-Horta et al., 2020; Griffiths and Philippot, 2013; Philippot et 66 al., 2013; Wagg et al., 2014). However, some microorganisms have developed several 67 resistance mechanisms to cope with their toxic effects such as extra and intracellular 68 sequestration, exclusion by permeability barriers, enzymatic detoxification, efflux-pumps and 69 specific reduction of cellular targets' sensibility (Hobman et al., 2007; Nies, 2003; Rouch et 70 71 al., 1995). In this frame, soil microorganisms constitute useful bio indicators of soil quality 72 (Alvarenga et al., 2018; Thiele-Bruhn et al., 2020). Measurement of microbial activity through analysis of soil extracellular enzymatic activities has long been used as indicator of 73 74 soil disturbance and can allow the assessment of the impact of environmental contaminates on soil microbial processes (Campos et al., 2018a, 2018b; Chu et al., 2007; Elmayel et al., 2020; 75 Higueras et al., 2019a; Hinojosa et al., 2004). Additionally, the implementation of state of the 76 77 art high throughput sequencing approaches could provide a high-resolution analysis of the ecotoxicological effects of PTEs on the composition and diversity of soil microbial 78 79 populations (Fernandes et al., 2018; Gallego et al., 2019; Jacquiod et al., 2018; Kisková et al., 80 2018).

81 The province of Ciudad Real, in the center of the Iberian Peninsula, is scattered with abandoned and decommissioned mines (Bravo et al., 2019; García-Lorenzo et al., 2019; 82 Higueras et al., 2017, 2012; Martín-Crespo et al., 2015; Rodríguez et al., 2009; Ruiz et al., 83 2009). In this regard, the San Quintín mine site (Cabezarados, Ciudad Real, Spain) offers an 84 unprecedented opportunity to shed light on this topic. This area covers about 100 Ha, with a 85 complex long-lasting history of mining and mineralurgical operations (Fig. 1). Between the 86 87 years 1888 and 1923, three mine shafts and a rudimentary froth flotation plant operated to obtain Ag-rich galena (PbS) concentrates, leaving a first legacy of dumps, very heterogeneous 88 89 in grain size and containing high concentrations of Pb, and Zn (García-Lorenzo et al., 2019).

In 1973, a second froth flotation plant aimed to recuperate the sphalerite (ZnS) as well as 90 91 residual Ag-rich galena was established. During this period, the froth flotation process favored the conversion of the dumps in wastes dams, which accumulated the residues from 92 this reprocessing. Additionally, cinnabar (HgS) from the adjacent Hg mines of Almadén 93 (Higueras et al., 2006; Tejero et al., 2015) was accumulated in the area during the last years of 94 activity of the froth flotation plant (1988-1990). In 2000, olive-oil milling wastes (OMW) 95 were illegally applied over the soils without any remediation interest. Currently, soil and 96 waste materials from the area are used in local road pavements, resulting in a redistribution of 97 the soil. Consequently, the area is deeply polluted with Pb, Zn and associated elements 98 99 (García-Lorenzo et al., 2019; Higueras et al., 2017; Martín-Crespo et al., 2015; Rodríguez et al., 2009; Ruiz et al., 2009). 100

101 The characterization of the behavior of PTEs from heavily degraded mining soils and their impact on the environment is the first step for planning restoration strategies in contaminated 102 areas. We hypothesized that decades of exposure to different levels of pollutants have exerted 103 104 a selective pressure on microbial communities in the San Quintín mine area. Within this context, this study aims to investigate the link between geochemical characteristics and 105 microbial communities in a long-term polluted mining site and assess the effects of PTEs on 106 107 the activity, composition and diversity of microbial communities through enzymatic analysis and high throughput sequencing. 108

109

110 2. Materials and Methods

111 **2.1 Samples collection**

112 Twenty samples of circa 800 g corresponding to the first 10-15 cm and relative to five 113 materials were collected (four independent replicates per material, each replicate was a

composite made out of three subsamples collected and homogenized in situ). The five 114 different materials included: non-directly polluted soil taken from the surrounding area and 115 used as local background (reference-samples); PTEs polluted soil from mine operations (via 116 leaching and runoff) and collected within the precincts of the mining site (contaminated-117 samples); olive-oil mill wastes residues spilled in the area (OMW); processed waste 118 accumulated in the dams (tailings); and mine wastes without any treatment (dumps). Samples 119 were immediately transported to the laboratory and sieved (<2 mm) within 24 hours. Samples 120 were divided in three aliquots and stored at -4°C (physicochemical and enzyme analysis) 121 (Peoples and Koide, 2012) or -20°C (DNA extraction) until use. Location and main 122 123 characteristics of sampling points are described in Fig.1 and Table 1.

124

125 **2.2 Physicochemical analysis**

Physicochemical analyses were conducted in the Laboratory of Soil Biogeochemistry at 126 the EIMIA-UCLM, Almadén (Spain). Electric conductivity and pH were determined using 127 ISO-10390 (ISO-10390, 2005) and ISO-7888 (ISO-11265, 1994) protocols, respectively. 128 Analyses were performed on 2 g dwt of sieved samples. Texture was determined on dried 129 samples treated with a solution of 3% (v/v) H_2O_2 for 48 h to remove organic matter. Samples 130 were then wet and sieved at 2 mm and analyzed with a Fritsch Analysette MicroTec Plus 22 131 (Fritsch, Germany). A textural classification triangle plot was used to identify the 132 granulometry of samples (Gerakis and Baer, 1999). Organic matter (OM) was determined 133 using the Walkley-Black method (Walkley and Black, 1934). 134

Elemental concentration analyses were performed from ground dried samples withparticle length minor to 100 μm determined by means of X-Ray Fluorescence spectroscopy,

using an Epsilon 1 device (Malvern Panalytical, England). Pollution Load Index (PLI) (Jorfi
et al., 2017) was calculated according to the equation:

139
$$PI = Cn/Bn$$

140
$$PLI = \sqrt{PI1\tilde{n}PI2\dots PIn}$$

141 Where PI is the single factor pollution index of each metal, C_n and B_n is the concentration of 142 metal in the sample and background, respectively (mg Kg⁻¹). Reference average values from 143 this study were used as local background level. The values of PLI allow the qualification of 144 samples as follows: PLI < 2: moderately to unpolluted; $2 \le PLI < 4$: moderately polluted; $4 \le$ 145 PLI < 6: highly polluted; PLI > 6: very highly polluted.

146

147 **2.3 Enzyme analysis**

Enzyme activities were measured through colorimetric techniques using a Biochrom Libra
S60 spectrophotometer (Biochrom, United Kingdom). Dehydrogenase (DHA) activity was
measured according to the method described by Casida (1977), with slight modifications.
Acid and alkaline phosphomonoesterase (PhA 6.5 and PhA 11, respectively) activities were
determined following Tabatabai and Bremner (1969) method. Arylsulfatase (ARS) activity
was measured as described by Tabatabai and Bremmer, (1970) and β-galactosidase (β-Gal)
activity was assayed according to Eivazi and Tabatabai (1988).

155 Functional diversity from selected enzymes was assessed with the geometric mean index156 (GMean) (Lessard et al., 2014) calculated as follow:

157
$$GMean = \left(\prod_{i=1}^{n} y_i\right)^{\frac{1}{n}}$$

158 Where y_i is the mean value for each enzyme activity, *n* is the total number of enzymes.

160 **2.4 Microbiological analysis**

DNA was extracted using the Power soil DNA isolation kit (Qiagen). The extracted DNA 161 was quantified by using fluorescent dye of a Quant-iTTM PicoGreen® dsDNA assay kit 162 163 (Invitrogen). The diversity of the bacterial community was determined by high throughput sequencing of 16S rRNA amplicons generated in a two-step PCR. First PCR reaction was 164 performed using the universal bacterial primers U341_F-805_R with overhang adapters 165 166 (forward adapter: TCGTCGGCAGCGTC AGATGTGTATAAGAGACAG, reverse adapter: GTCTCGTGGGCTCGGAGATGTGTATA AGAGACAG). The reaction mixture contained 2 167 ng of DNA as template, 7.5 µL of 2X Phusion High Fidelity PCR Mastermix (Thermo 168 169 Scientific, Waltham, MA, USA), 250 ng of T4 gene 32 protein (MP Biomedicals, Santa Ana, CA, USA), 0.375 µL of each primer (10 µM) and ultrapure sterile water to a total volume of 170 15 µL. Thermal conditions were 3 min at 98°C, 25 cycles at 98° C for 30 sec, 55° C for 30 sec 171 and 72° C for 30 sec followed by a final extension of 10 min at 72°C. Duplicates of each 172 amplicon were pooled and then 6 µL aliquot was used as template in a second PCR carried 173 174 out with multiplexed primers containing the universal overhang adaptors and specific barcodes and using a 384 Nextera XT index kit (Illumina, San Diego, CA, USA). The reaction 175 mixture was carried out in 30 µL reaction volumes containing 2.5 µL sterile water, 15 µL 2X 176 177 Phusion HF master mix (Thermo Scientific, Waltham, MA, USA), 250 ng of T4 gene 32 protein (MP Biomedicals, Santa Ana, CA, USA), 3 µL of each primer (10 µM) and 6 µL of 178 the step-one PCR product. The thermal cycling was 98°C for 3 min, followed by eight cycles 179 of 98° C for 30 sec, 55°C for 30 sec and 72° C for 30 sec, with a final extension of 72° C for 180 10 min. The size of the amplicons was verified by electrophoresis on a 2 % agarose gel. 181 Amplicons were purified (amplicon library purification, PicoGreen® quantification and 182 pooling) and sequenced (Illumina MiSeq 2 x300bp) by Microsynth (Balgach, Switzerland). 183 Amplicons were normalized (SequalPrepTM kit), purified (Pippin prep) and sequenced by 184

Microsynth (Switzerland). The sequence data was analyzed using an in-house developed 185 186 Phyton notebook pipeline together with different bioinformatics tools: 16S rDNA sequences were assembled using the PEAR software (Zhang et al., 2014) with the default settings; 187 further quality checks were conducted using the QIIME 1 pipeline (Caporaso et al., 2010a); 188 sequences shorter than 350 bp were removed; reference-based and de novo chimera detection, 189 as well as clustering in OTUs were performed with the VSEARCH software (Rognes et al., 190 191 2016) using appropriate reference databases (Greengenes' representative set of sequences for 192 16S rDNA) with a threshold placed at 94 % identity; representative sequences for each OTU were aligned using PyNAST (Caporaso et al., 2010b); phylogenetic trees were constructed 193 194 using FastTree (Price et al., 2009); taxonomy was assigned using UCLUST (Edgar, 2010) and the latest released Greengenes database (v.05/2013, McDonald et al., 2012) for 16S rDNA 195 sequences; sequences were deposited in the GenBank to the sequence read archive (SRA) 196 197 under the accession number PRJNA646888. A range of bacterial a-diversity indices pertaining to richness (Chao1, observed species), evenness (Simpson reciprocal, equitability, 198 199 dominance, Shannon) and relatedness (PD whole tree) were calculated based on rarefied 200 tables (13,000 sequences per sample). Bray Curtis distance matrices were also computed to detect changes in the composition of microbial communities. Canonical Analysis of Principal 201 Coordinates (CAP) of OTU Bray Curtis distance matrices were also performed and plotted. 202 203 The relative abundance of the different bacterial phyla was also determined using a comparative bar chart. 204

205

206 **2.5 Statistical analysis**

All statistical analyses were performed in R (http://www.r-project.org). The normality of the data and residuals was checked (Shapiro Wilk's with p>0.05) and the homogeneity of variances was verified (Levene's test with p>0.05). Root square, inverse and logtransformations on the data were performed when necessary. For parametric distributions,
ANOVA followed by Tukey's test and t-student were used to determine differences. For nonparametric distributions, data was compared using Kruskal Wallis test. To detect significant
differences in communities' structure among sample types, Permutational multivariate
analysis of variance (PermANOVA) was used on Bray Curtis dissimilarity matrix using
Adonis function from R package "vegan" (Oksanen et al., 2018).

Integration and visualization of amplicon sequencing and physicochemical and enzymatic analysis data sets were performed through the R package mixOmics (Rohart et al., 2017) using DIABLO (Data Integration Analysis for Biomarker discovery using a Latent component method for Omics studies) in order to identify correlated key omics variables in both datasets (Singh et al., 2019).

221

222 **3. Results**

223 **3.1** Physicochemical and geochemical characterization of samples

Table 2 shows the physicochemical parameters of the samples of the five selected 224 materials. According to their granulometry, samples were classified with some minor 225 variability in sandy loam (reference-, contaminated-samples and dumps), silt loam (tailings) 226 and loamy sand (OMW) (Table 2, Table S1 and Fig. S1). Reference-, contaminated-samples 227 and tailings presented a slightly acidic character while OMW and dumps had lower pH 228 values. EC varied from 62.4 ± 27.8 (reference-samples) to 2282.3 ± 1786.4 mV (dumps). 229 Overall, no significant differences were found in the pH and EC (p>0.055), except for the 230 dumps, which presented significant lower pH (p=0.00371) and significant higher EC values 231 (p=0.004) as compared to the reference-samples. OM ranged from $1.57\% \pm 0.54$ (dumps) to 232 $60.60 \% \pm 19.67$ (OMW). Similar organic content was found in reference-samples, tailings 233

and dumps, while higher values were found in contaminated-samples, although not significant (p>0.5006), probably due to their high heterogeneity. On the contrary, significant higher organic matter was found in OMW as compared to the rest of the samples (p=0.0000271).

237 The concentration of PTEs for the different samples is shown in Table 3. Among the different PTEs studied, those which were part of the ore (Pb, Zn and Hg) were found at very 238 high concentrations in all samples except reference-samples, with significantly higher 239 240 concentrations for Pb (p<0.0043) and Zn (p<0.00577) in dumps and OMW as compared to reference-samples. Similarly, Sb, Co and Cu concentrations were significantly higher in 241 242 dumps and OMW than in the three other samples (p<0.0407). Regarding Mn, Ni and Th, 243 similar concentrations were recorded in all samples and no significant differences were observed (p>0.05). A high variability was found for all the PTEs studied. The pollution index 244 for each PTE was calculated (Table S2) and yielded values higher than 100 for Hg (in all 245 samples except reference-samples), Pb (dumps and OMW) and Zn (OMW). Pollution load 246 index (PLI) allowed to classify the samples as follows: reference- and contaminated-samples 247 248 as highly polluted ($4 \le PLI \le 6$) and OMW, dumps and tailings (from most polluted to less polluted respectively) as very highly polluted (PLI > 6) (Table 3). 249

250

251 **3.2 Enzymatic analysis**

Five enzymatic activities including the dehydrogenase (DHA), arylsulfatase (ARS), the acid and alkaline phosphomonoesterase (PhA 6.5 and PhA 11, respectively) and β galactosidase (β -Gal) were measured in the samples collected from the five sites studied in mining area (Fig. 2 and Table S3). Reference- and contaminated-samples showed similar enzyme activities (p>0.05). The lowest enzymatic activities (ARS, PhA 6.5, PhA 11 and β -Gal) were recorded in the OMW, which were significantly different from the values recorded in reference- and contaminated-samples (p<0.0398351) Tailings and dumps had significantly lower DHA value (reduction of about 96 %) and ARS activities than reference-samples (p<0.0018687). Tailings had PhA 6.5 activities significantly lower than that of the referencesamples (p=0.0223). The geometric mean index was significantly lower in OMW samples (p=0.0000002) than that of all the other samples, which were similar (p>0.06922) (Fig. S2).

263

264 **3.3 Bacterial composition and diversity**

The composition and diversity of the bacterial community was assessed by high throughput sequencing of 16S rRNA amplicons amplified from DNA extracts. In total, after de-multiplexing and removal of low-quality raw sequence reads, high throughput sequencing generated 1,796,830 high quality sequences with an average sequence length of 461 ± 11bp. Using a threshold at 94% nucleotide sequence identity, these sequences were grouped in 16,299 different OTUs.

271 A range of α -diversity indices pertaining richness (Chao1), relatedness (PD whole tree) and evenness (Simpson reciprocal) were calculated using rarefied data (Fig. 3). Overall, 272 Chao1, PD whole tree and Simpson reciprocal α -diversity indices were higher in reference-273 samples than in contaminated-samples, OMW, tailings and dumps, ranked from the highest to 274 the lowest values. Chao1 values were significantly lower in OMW, tailings and dumps 275 (p<0.027) while significant differences were found for tailings and dumps in PD whole tree 276 277 values as compared to reference-samples values (p<0.019). Regarding Simpson reciprocal, no significant differences were found between reference-, contaminated-samples, OMW and 278 tailings, and only significant lower values were found in dumps (p=0.024). 279

Differences in the composition of bacterial communities measured in the different samples
were assessed by multivariate beta-diversity analysis using Canonical Analysis of Principal

Coordinates (CAP) based on Bray Curtis distances matrices (Fig. 4 & Fig. S3). A good 282 283 reproducibility between replicates of each sample type was observed regardless the dataset used as explanatory factors. CAP ordinations using a range of PTEs (Co, Cu, Hg, Mn, Ni, Sb, 284 285 Th, Pb, Zn) as explanatory factors, explained up to 78 % of the variance observed, of which 29% is explained in the first two axes (17% and 12%, respectively) (Fig. 4a). The sample type 286 was the most explanatory variable (36.96%) followed by the Hg concentration (6.41%), both 287 288 associated with the first axis. All the other variables (Mn, Co, Ni, Cu, Zn, Sb, Pb and Th) 289 analyzed were not significant (Table S4). Likewise, CAP ordinations using physicochemical parameters and enzymatic activities as variables explained 73% of the variance observed 290 (17% and 12% respectively for the first two axes) (Fig. 4b). None of the physicochemical 291 parameters nor the enzymatic activities were significant explanatory variables of the 292 composition of the bacterial community. For all CAP ordinations, reference- and 293 294 contaminated-samples showed similar bacterial community composition (p=0.734) that was differing from those of tailings and dumps which were similar (p=0.056). The composition of 295 296 the bacterial community of OMW was markedly different from that of all the other samples 297 analyzed no matter the variables considered (p < 0.036).

Taxonomic analysis of the 16S rRNA amplicon sequences at phylum level showed that 298 299 the relative abundance of the major bacterial phyla was very similar in reference- and contaminated- samples. On the contrary, it markedly differed between reference-samples and 300 dumps. All samples were dominated by bacteria belonging to Proteobacteria, Actinobacteria, 301 302 Chloroflexi, Acidobacteria, Gemmatimonadetes, Bacteroidetes, Firmicutes, TM7, Verrucomicrobia and Cyanobacteria (Fig. 5). Proteobacteria, Actinobacteria, Chloroflexi, 303 304 Acidobacteria, Firmicutes, TM7 and Cyanobacteria were observed in similar abundance in all the sample types (p>0.05). Gemmatimonadetes and Bacteroidetes were found in significantly 305 306 lower amount in dumps than in reference-samples (p<0.035). Verrucomicrobia had significantly lower relative abundances in dumps, tailings and OMW than in the referencesamples (p < 0.024).

309 To further identify the OTUs responsible of differences in the composition of the bacterial 310 community between samples, a correlation analysis was performed with OTUs represented at least once in all four replicates of a given sample type (a total of 76 different OTUs) using the 311 diablo R package. Among these 76 OTUs, 57 responsible of the differences observed between 312 313 substrate types (Pearson's correlation > 0.4) were plotted in a heatmap (Fig 6). All discriminant OTUs affiliated to 7 of the 10 most abundant bacterial phyla detected 314 315 (Actinobacteria, Chloroflexi, Acidobacteria, Bacteroidetes, Cyanobacteria, 316 Gemmatimonadetes and Proteobacteria).

317 Similar abundances were found in discriminant OTUs for reference- and contaminatedsamples, confirming the results from α -and β -diversity analyses. Microorganisms belonging 318 to the orders Actinomycetales, Sphingomonadales, Burkholderiales, Rhodospirillales, 319 320 Stramenopiles, RB41, Solibacterales, Ellin 329 and some microorganisms belonging to the 321 order Rhizobiales were highly represented in reference- and contaminated-samples. Microorganisms belonging to the families Acidobacteriaceae, Sphingobacteriaceae and 322 Xanthomonadaceae, the order Solirubrobacterales, N1423WL, some Actinomycetales and 323 324 some Rhizobiales; and the genera Sphingomonas and Segetibacter were highly represented in OMW but scarcely detected in reference- and contaminated-samples. Regarding the dumps, 325 326 five OTUs belonging to Acidomicrobiales, B12-WMSP1, Acidobacteriaceae and Enterobacteriaceae phylotypes were found in high abundance. Finally, a high heterogeneity 327 was observed for the discriminant OTUs in tailings. 328

329

330 4. Discussion

Mining soils are generally rich in PTEs that are known to have ecotoxicological effects on 331 332 soil microorganisms by disrupting their metabolism, affecting soil enzymatic activity and having consequence on the abundance, composition and diversity of soil microbial 333 334 communities (Zhao et al., 2020). The analysis of the PTEs concentrations in the five different sample types allowed the classification of the samples as highly polluted (reference- and 335 contaminated-samples) and very highly polluted (OMW, tailings and dumps). Although the 336 potential value of OMW as fertilizer or metal immobilization amendment in polluted soils 337 was shown in previous studies (Hmid et al., 2014; Paredes et al., 1999), here the application 338 of OMW showed no remediation activity. It is noteworthy that a number of recent studies 339 have shown the negative impact of OMW on aquatic and terrestrial ecosystems (Ntougias et 340 al., 2013). Soil enzymes are mainly produced by microorganisms, which play a crucial role in 341 nutrient cycling and consequently, in soil fertility (Adetunji et al., 2017). Microbial enzymes 342 343 are highly sensitive to metals, and their use as standard biochemical indicators has been widely proposed to evaluate the quality of polluted soils (Pajak et al., 2018; Tang et al., 2019). 344 345 However, several studies have showed no effect of PTEs contamination on some soil 346 enzymatic activities, such as dehydrogenase, catalase, acid phosphomonoesterase and amylase (Campos et al., 2018a; Liu et al., 2020; Yang et al., 2016). This phenomenon could be due to 347 three reasons: (I) the long-term presence of metals in the soil, which might promote the 348 development of heavy metal resistant microorganisms (Ciarkowska et al., 2014); (II) 349 variations in soil physicochemical properties, which may influence the activity of soil 350 microbial communities (Xian et al., 2015), the bioavailability and speciation of the pollutants 351 352 (Rieuwerts et al., 1998); and (III) antagonistic effects between various PTEs (Tang et al., 2019). Our results showed that enzyme activities were similar in reference- and contaminated-353 354 samples. This might be explained by the high organic matter content in these two sample types, which may on the one hand, provide nutrient source for the development of 355

microorganisms and the production of enzymes and on the other hand, reduce the 356 bioavailability of metals and metalloids, limiting their toxicity towards microbial enzymatic 357 activities (De Santiago-Martín et al., 2013; Lair et al., 2007). It is important to note, that 358 organic matter can sometimes increase the mobility and bioavailability of pollutants (Lindsay, 359 1991; Meunier et al., 2011; Wang and Mulligan, 2006). Consequently, further research is 360 needed to better understand the processes governing the complex interactions between organic 361 matter and pollutants. Additionally, all enzymatic activities were very low in OMW as 362 confirmed by the geometric mean index. The important amount of phenols with powerful 363 antimicrobial properties and high content of PTEs, might explained the low enzymatic 364 activities recorded in OMW samples (Siles et al., 2014). DHA activities measured in tailings 365 and dumps were significantly lower than those of reference-samples. Keeping in mind the fact 366 that the inhibition of DHA activity in response to PTEs such as Pb, Cu, Cd, Zn has been 367 368 previously described (De Santiago-Martín et al., 2013; Pająk et al., 2018; Parelho et al., 2016; Tang et al., 2019), one could hypothesize that the high concentration of PTEs in tailings and 369 370 dumps would explain the observed low enzyme activities. Similar results were found for ARS 371 activity, that is a key player in the sulfur cycle and known to be inhibited by PTEs (De Santiago-Martín et al., 2013; Gülser and Erdoğan, 2008; Stefanowicz et al., 2020). No-matter 372 the sample type, acid PhA activities were higher than those of alkaline PhA probably due to 373 374 the acidic nature of all the samples (Adetunji et al., 2017). Acid PhA activity was lower in tailings than in the reference-samples which is in accordance with known sensitivities of these 375 enzymes to heavy metals (Angelovičová et al., 2014; De Santiago-Martín et al., 2013; Gao et 376 377 al., 2010). Despite the fact that β -Gal has been reported as a sensitive bioindicator of metal contamination of soils (De Santiago-Martín et al., 2013; Martínez-Iñigo et al., 2009), all 378 379 samples showed similar β -Gal activities except in dumps where slightly lower activities were recorded. 380

As observed with microbial enzymatic activities, the exposure to PTEs can also affect key 381 382 microbial processes, which reflects on changes in microbial composition and diversity. On the one hand, PTEs such as Pb, Zn or Hg can exert toxic effects compromising the survival and 383 growth of specific microbial guilds (Abdu et al., 2017; Doelman and Haanstra, 1984). On the 384 other hand, PTEs can promote the emergence of specific resistant microorganisms able to 385 cope with the ecotoxicity of these contaminants (Gillard et al., 2019; Martinez et al., 2006; 386 Sobolev and Begonia, 2008). Here, the lowest values of all the α-diversity indices (Chao1, PD 387 whole tree and Simpson reciprocal) were recorded in dumps, probably as consequence of the 388 very high concentrations of Pb, Zn, Sb, Co and Cu present in these samples and low organic 389 390 matter. Indeed, the reduction in soil bacterial α-diversity as consequence of Pb, Zn, Sb, Co or Cu toxic effects has previously been reported (Luo et al., 2018; Nunes et al., 2016; 391 Stefanowicz et al., 2008; Xie et al., 2016). Similarly, Chao1 and PD whole tree values were 392 393 lower in tailings than in the reference-samples. It is noteworthy that PTEs concentrations in tailings were not significantly different from that of reference-samples, suggesting that 394 395 differences observed in a-diversity may result from other physicochemical parameters 396 governing the bioavailability of PTEs. Chao1 diversity indices in OMW samples were significantly lower than that of reference-samples. As observed for the enzymatic activities, 397 the high content of organic matter rich in phenols in OMW could have ecotoxicological 398 399 effects on survival and growth of phenol-sensitive microorganisms and therefore decreased the microbial diversity. In this regard, changes in the structure of ammonium oxidizers 400 bacteria and Actinobacteria have been previously reported (Karpouzas et al., 2010; Mekki et 401 402 al., 2006). All alpha diversity indices were similar in both reference- and contaminatedsamples. 403

404 Multivariate β -diversity analysis based on Bray Curtis distance matrix confirmed the 405 results observed from α -diversity: reference- and contaminated-samples clustered together no 406 matter the explanatory factors (PTEs, enzymatic activities or texture). These two samples 407 were clearly separated from tailings and dumps. OMW was discriminated from all others 408 whatever the factors considered. Besides the sample type, which explained 37% of the 409 variance observed, concentration of Hg was the only significant explanatory factor (6% of the 409 variance). In line with our work, the influence of soil properties and heavy metals on the 411 structure of bacterial communities has been previously showed in contaminated soils (Li et 412 al., 2017; Liu et al., 2018; Nunes et al., 2016; Tipayno et al., 2018)

Phylogenetic analysis revealed that the majority of bacteria from all samples affiliated to 413 414 the Proteobacteria, Actinobacteria, Chloroflexi, Acidobacteria, Gemmatimonadetes, 415 Bacteroidetes, Firmicutes, TM7, Verrucomicrobia and Cyanobacteria phylotypes, in line with previous studies performed in a range of PTEs polluted soils (Chen et al., 2018; Guo et al., 416 2017; Pacwa-Płociniczak et al., 2018; Tipayno et al., 2018; Zhen et al., 2019). Overall, their 417 relative abundances were similar in all samples, with the exception of the bacteria belonging 418 to Gemmatimonadetes, Bacteroidetes and Verrucomicrobia, whose relative abundance were 419 420 significantly lower in dumps as compared to reference-samples. A significant reduction in the relative abundance of Verrucomicrobia was also observed in tailings and OMW. The 421 significantly higher concentrations of PTEs found in in tailings and OMW may explain the 422 reduction of these phylotypes. This assumption is in line with the reduction of 423 Verrucomicrobia previously reported in a study performed on an agricultural paddy soil 424 polluted with cadmium (Luo et al., 2019) and Yellow river sediments heavily polluted with 425 426 cadmium, arsenic, lead and mercury (Chen et al., 2018). Similarly, the relative abundance of members of several phyla including Verrucomicrobia and Bacteroidetes was found to 427 428 significantly decrease in a long-term Cu polluted site (Berg et al., 2012). In this regard, our findings are consistent with a recent study that proposes that changes in the response of some 429 430 bacterial groups (the Verrucomicrobia/Chlamydiae ratio) can be used as bioindicators of heavy metal pollution (Schneider et al., 2017). Contrary to our results, studies from a metal
polluted forest soil showed an increase in the Gemmatimonadetes (Azarbad et al., 2015) but a
negative correlation between the relative abundance of Verrucomicrobia and pollution levels
as observed in our study.

Further analysis at lower taxonomical level led to the identification of 57 OTUs 435 responsible of α - and β - diversity changes observed between different sample types. 436 437 Discriminant OTUs were found in similar abundances in reference- and contaminatedsamples. Interestingly, OTUs barely found in reference- and contaminated-samples were 438 439 significantly more abundant in OMW, suggesting that these OTUs may be able to grow in a 440 PTEs contaminated environment rich in organic matter. OTUs related to the B12-WMSP1 group described by Costello and Schmidt (Costello and Schmidt, 2006) in fumarolic soils 441 (Costello et al., 2009; Schmidt et al., 2018; Tebo et al., 2015) were found in higher amount in 442 dumps than in the reference-samples. 443

444

445 **5. Conclusions**

The importance of soil health and quality in ecosystem services underlines the need to 446 447 understand the processes governing the behavior and effects of PTEs on soil microorganisms and supported functions. This assumption is especially relevant in highly polluted areas such 448 449 as mining areas. Here, we evaluated the degree of contamination in a long-term mining site 450 and assessed the ecotoxicological effects of PTEs on bacterial communities. We observed that low PTEs concentrations can have a strong impact on the diversity and composition of 451 452 microbial communities while physicochemical parameters such as organic content can either help counteract or enhance their negative effects. Understanding the relationship between 453 physicochemical parameters and diversity and composition of microorganisms can contribute 454

to the environmental risk assessment of long-term polluted sites and help create strategies torestore contaminated sites and mitigate the pollution dissemination.

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465 ABBREVIATIONS

- 466 ARS, Arylsulfatase enzyme activity
- 467 DHA, Dehydrogenase enzyme activity
- 468 EC, Electric conductivity
- 469 OMW, Olive-oil mill waste
- 470 OM, organic matter
- 471 PhA, Phosphomonoesterase enzyme activity
- 472 PTEs, Potentially toxic elements
- 473 β-Gal, β-galactosidase enzyme activity

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Figures



Figure 1. Geographical area and sampling sites (reference, green; contaminated, red; olive mill waste, yellow; tailings, blue and dumps, grey) in the study area.



Figure 2. Enzymatic activities (mean \pm S.D): dehydrogenase (DHA) (µg TPF g⁻¹ DW day⁻¹), arylsulfatase (ARS) (µg PNF g⁻¹ DW day⁻¹), acid phosphomonoesterase (PhA 6.5) (µg PNF g⁻¹ DW day⁻¹), alkaline phosphomonosterase (PhA 11) (µg PNF g⁻¹ DW day⁻¹) and β-Galactosidase (β-Gal) (µg PNF g⁻¹ DW day⁻¹) for reference-, contaminated-samples, olive mill waste (OMW), tailings and dumps. ANOVA followed by Tukey's test and Kruskal Wallis tests (*) were performed. For DHA, ANOVA was performed on log-transformed data. For each parameter, different letters within the same line indicate significant differences (p <0.05). n=4



Figure 3. Bacterial α -diversity indices (mean values \pm standard deviation) derived from amplicon sequencing of the 16S rDNA gene copies from reference-, contaminated-samples, olive mill waste (OMW), tailings and dumps. ANOVA test followed by Tukey's test was performed. For each diversity index value, different letters indicate that samples are significantly different for each treatment and time. For dominance, ANOVA was performed on log-transformed data. n=4

a

b

Figure 4. Bacterial β -diversity analysis for reference-, contaminated-samples, olive mill waste (OMW), tailings and dumps using Bray Curtis distance matrix of 16S rRNA sequences and (a) PTEs concentrations or (b) enzymatic analysis as explanatory factors. The variance explained by each axis is given as percentage. For each sample type, the four replicates are represented with the same color.

Figure 5. Relative abundance of the major bacterial phyla (expressed as % of the total number of OTUs) for reference-, contaminated-samples, olive mill waste (OMW), tailings and dumps. Phyla whose relative abundance was below 5% were grouped as "others". ANOVA followed by Tukey's test and Kruskal Wallis tests (Proteobacteria, Chloroflexi, Acidobacteria and Cyanobacteria) were performed. For Firmicutes and TM7, ANOVA was performed on log-transformed data. n=4

Figure 6. Heatmap plot representing relative abundances of OTUs correlated to physicochemical and enzymatic analysis (correlation cut-off r>0.4) in reference-, contaminated-samples, olive mill waste, tailings and dumps.

Tables

Sample type	Sample	X (UTM)	Y (UTM)	Altitude	Other Characteristics
Reference	SQD-9	388986	4298003	666	Recently fertilized
Reference	SQD-13	390156	4298068	664	
Reference	SQD-18	390256	4297642	671	Dry land, plowed
Reference	SQD-19	390207	4297137	669	Fallow, unplowed
Contaminated	SQD-8	388935	4297593	687	Agricultural
Contaminated	SQD-11	389261	4297675	675	2 meters from a plant
Contaminated	SQD-12	389392	4297805	678	Developed soil
Contaminated	SQD-14	389798	4297560	663	
Olive Mill Waste	SQD-2	389170	4297396	655	
Olive Mill Waste	SQD-4	389103	4297409	659	
Olive Mill Waste	SQD-6	389059	4297448	657	
Olive Mill Waste	SQD-20	389238	4297521	671	With visible dump debris
Tailings	SQD-1	389208	4297287	646	Aeolic erosion
Tailing	SQD-15	389704	4297503	667	
Tailing	SQD-16	389521	4297623	667	
Tailing	SQD-17	389553	4297525	673	
Dumps	SQD-3	389129	4297398	658	Heterogeneous granulometry
Dumps	SQD-5	389081	4297558	658	Heterogeneous granulometry
Dumps	SQD-7	389033	4297470	662	Flooded area
Dumps	SQD-10	389032	4297607	674	Heterogeneous granulometry

Table 1. Distribution (geographic coordinates) and main characteristics of samples

Table 2. Physicochemical characteristics of reference-, contaminated-samples, olive mill waste, tailings and dumps. Values were mean \pm standard deviation. ANOVA followed by Tukey's test were performed. For EC and OM, ANOVA was performed on log-transformed data. For each parameter, different letters within the same line indicate significant differences (p <0.05). For EC and OM, ANOVA performed on log-transformed data. n=4

	Reference	Contaminated	Olive Mill Waste	Tailings	Dumps	
pН	$6.1 \pm 0.3a$	$6.8 \pm 1.0a$	5.2 ± 0.6 ab	6.7 ± 1.7a	$3.9 \pm 0.5b$	
EC (mV)	$62.4 \pm 27.8a$	$90.6 \pm 49.9a$	651.3 ± 489.7ab	415.6 ± 378.8ab	2282.3 ± 1786.4b	
OM (%)	$2.9 \pm 1.9a$	8.1 ± 8.9a	$60.6 \pm 19.7 \mathrm{b}$	$2.0 \pm 0.4a$	$1.6 \pm 0.5a$	
Sample type	sandy loam	sandy loam	loamy sand	silt loam	sandy loam	
EC 1. this can be divide OM and a sector						

EC: electric conductivity, OM: organic matter

Table 3. PTEs concentrations (mg/kg) (mean values ± standard deviation) and Pollution Load
Index (PLI) in reference-, contaminated-samples, olive mill waste, tailings and dumps.
ANOVA test followed by Tukey's test and Kruskal Wallis followed by pairwise comparisons
with Bonferroni correction (*) were performed. Values indicated by different letters are
significantly different within the same line. For Ni and Cu ANOVA was performed on root
square transformed data. For Sb, ANOVA was performed on inverse transformed data,
respectively. For Zn, Hg and Th, ANOVA performed on log-transformed data, respectively.
n=4

	Reference	Contaminated	Olive Mill Waste	Tailings	Dumps
Pb	315.8 ± 332.6a	$3696.5 \pm 4406.2a$	$38890 \pm 20687.0b$	$4030.2 \pm 7170.8a$	48597.5 ± 16835.1b
Zn	$204.8 \pm 153.8a$	1522.4 ± 2165.2ab	86050 ± 99094.2c	2024.5 ± 2492.1ab	$4640.0 \pm 2259.5b$
Hg	$0 \pm 0a$	$55.4 \pm 92.1a$	$3605.8 \pm 6752.0a$	557.1 ± 430.8a	$130.8 \pm 142.5a$
Sb	$16.8 \pm 1.8a$	37.4 ± 31.0 abc	$330.9 \pm 435.2 bc$	65.5 ± 87.7 ab	$433.6 \pm 198.9c$
Со	$221.7 \pm 76.4a$	268.3 ± 80.0 ab	$451.7 \pm 78.4 \text{bc}$	192.5 ± 27.9a	$503.9 \pm 180.6c$
Cu	$17.3 \pm 10.0a$	$165.6 \pm 297.6a$	963.7 ± 466.2b	$113.4 \pm 196.7a$	$1000.6 \pm 805.0b$
Mn*	688.2 ± 238.9 ab	1572.5 ± 908.6a	1637.4 ± 1866.4ab	423.5 ± 136.5ab	$210.4 \pm 90.8b$
Ni	$31.4 \pm 18.7a$	$38.9 \pm 31.4a$	$132.1 \pm 95.6a$	$37.0 \pm 14.0a$	$27.6 \pm 8.1a$
Th	9.7 ± 3.9ab	4.6 ± 4.1 ab	$0 \pm 0b$	3.8 ± 6.9 ab	$35.1 \pm 23.7a$
PLI	1.0 ± 0.1	3.2 ± 2.8	21.1 ± 23.9	10.5 ± 4.8	7.4 ± 2.4

PLI: Pollution Load Index

SAN QUINTÍN MINE

Analytical chemistry

Microbial enzyme activities

Composition and diversity of bacterial community

Ecotoxicological risk assessment of San Quintín mine area

