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Field sampling of soil pore water to evaluate the mobile fraction of trace elements in the Iglesiente area (SW Sardinia, Italy)

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- 1 Field sampling of soil pore water to evaluate the mobile fraction of trace elements in the
- 2 Iglesiente area (SW Sardinia, Italy)
- 3
- 4 Sara Concas¹, Carla Ardau¹*, Marcello Di Bonito², Pierfranco Lattanzi¹, Andrea Vacca¹
- 5 1. Dipartimento di Scienze chimiche e geologiche, Università di Cagliari, Italy
- School of Animal, Rural, and Environmental sciences, Nottingham Trent University, UK
 *corresponding author: carla.ardau@tiscali.it
- 8

9 Abstract

10 Field soil pore water monitoring was applied in a highly heavy-metal contaminated area in SW 11 Sardinia, Italy, as a direct, realistic measure of heavy metal mobility. The main chemistry of pore 12 waters well reflects the local characteristics of soils, ranging from Ca-SO₄ to (Ca)Mg-HCO₃ to 13 Ca(Na)-SO₄(Cl), with a wide range of conductivity. The mobility of Zn and Pb is apparently 14 controlled by equilibrium with minerals such as hydrozincite or smithsonite, and cerussite, 15 respectively. These results allow a correct estimate of the actual environmental risk associated with the presence of heavy metals in soils, and may serve as a supporting tool for phytoremediation 16 17 planning.

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19 Key words: soil pore water; heavy metal mobility; Iglesiente (Sardinia)

20

21 **1. Introduction**

The presence of potentially toxic chemical elements (PTEs) in soils, either naturally occurring or from anthropogenic activities (such as, amongst others, mining activity), represents a serious environmental threat, involving the entire system of relationships between hydrosphere, geosphere and biosphere (Nkongolo et al., 2013; Li et al., 2014). In the past century, mining was the driving sector of the economy in south-western Sardinia (Italy). Mining industry began to decline around the end of the century, leading to the closure of most mines over the 1970–1990 period. There are now in the region about 170 abandoned mines, and 200 million tons of mining-related wastes, covering a surface of 19 km² (RAS, 2003; Ardau et al., 2013). Due to the ineffective or totally absent management of mining-related wastes in the last decades, in several Sardinian areas the PTE contamination extended to the surrounding environment, affecting air, water and soils (e.g. Aru et al., 1995; Vacca & Vacca, 2001; Cidu et al., 2001; Concas et al., 2006; Cidu et al., 2011; Fanfani & Ardau, 2011; Vacca et al., 2012; Cidu et al., 2012; Ardau et al., 2013).

34 In abandoned mining areas, removal, treatment or confinement of mining-related wastes may 35 be not viable for several practical reasons, including the extremely high costs (Lomelin, 2002; 36 González et al., 2006; Ardau et al., 2013). A 'soft' alternative is represented by phytoremediation 37 (Barbafieri et al., 2011), that includes two different options: phytostabilization and phytoextraction 38 (Chaney et al., 1997). Phytoextraction may represent a valuable, cost-effective alternative to 39 traditional disruptive technologies (Cao et al., 2007), but it is often of limited application. A more 40 widely applicable technique is phytostabilization, because the quick establishment of a dense 41 vegetation cover is crucial to limit wind and water erosion. Moreover, plant roots may immobilize 42 metals by adsorption or accumulation, and provide a rhizosphere wherein metals precipitate and 43 stabilize (Mendez & Maier 2008; Cao et al., 2009; Nicoară et al., 2014).

44 One of the prerequisites for phytoremediation planning and management is to identify mobility and 45 phytoavailable fraction of (toxic) chemical components in soil/wastes. From many studies (Impelliteri et al., 2003; Tye et al., 2003; Cui et al., 2014), it is well established that total soil/waste 46 47 metal content alone is not an adequate measure of the potential mobility of a (toxic) element and, 48 consequently, of its possible uptake by plants (Vig et al., 2003; Di Bonito et al., 2008; Barbafieri et 49 al, 2011). Therefore, chemical methods to determine trace metal phytoavailability in soils are 50 usually based on the measurement of the extractable or labile fractions of those elements (De Siervi 51 et al., 2004). However, there is no agreement on how to evaluate at best the risk arising from the 52 fraction of pollutants that are most mobile in soils (Sneddon et al., 2006; Vázquez et al., 2008; Moreno-Jiménez et al., 2011). Among the possible methods of identification, the direct 53 54 measurement of the composition of soil pore waters is being increasingly applied, due to many

advantages in term of simplicity and costs (Meers et al., 2007; Di Bonito et al., 2008; Beesley et al.,
2010; Moreno-Jiménez et al., 2011; Shaheen et al., 2014).

57 In the present study, the composition (major element chemistry and selected heavy metal 58 contents) of soil pore water was determined in a highly heavy-metal contaminated area included in 59 the Iglesiente mining district (south-western Sardinia, Italy), one of the largest of the island, and of importance at the European scale. To the best of authors' knowledge, this is the first time that this 60 61 technique is applied in the district. The study area is located in a geographic region with a typically 62 Mediterranean climate, with very dry summers, and is characterized by natural geochemical 63 anomalies (particularly Pb, Zn, Ba), enhanced by centuries of mining activity (Bechstädt et al., 64 1994; Boni et al., 1999). The monitoring of temporal, local and spatial variations in pore water quantity and quality was carried on from a series of undisturbed soil/mine waste profiles. The study 65 is a significant part of a more extended project (Concas, 2014). In a companion paper (Concas et al., 66 67 submitted), the metal uptake by an endemic Mediterranean species (Pistacia lentiscus L.) was assessed. Although not part of this specific paper, some recall to that supplementary information 68 69 will be made for a more robust data discussion.

70

71 **2. Materials and methods**

72 2.1 Study site

The Iglesiente mining district (Sardinia, Italy) was for centuries up to the 1990s the seat of intense mining activity. The ore bodies are hosted in Cambrian dolomites and limestones, and produced some hundred million tons of Zn–Pb–Ag and Ba ores (Boni et al., 1999, and references therein). The cessation of mining activity left large quantities of mine wastes in dumps and flotation tailings basins, estimated at about 45 million m³ for the whole area (Cao et al., 2009).

Concas (2014) describes a total of 9 soil profiles in 5 sites in the basin of the San Giorgio river, having an extension of about 2,563 ha (Fig.1). Four main representative sites were selected for this study. The first site (*Campo Pisano*) is located near the town of Iglesias, the main urban centre of 81 the area. Campo Pisano is one of the hazard centres of the district because of the presence of waste 82 impoundments where, during mining activity in the 1900s, flotation wastes from several mines of 83 the district were collected, in particular from Monteponi mine (Cidu et al., 2001, RAS, 2003). The flotation wastes (about 8 million m³) contain Pb and Zn as the most abundant heavy metals. Due to 84 85 the high heavy metal concentrations, low organic carbon, and low cation exchange capacity, the flotation wastes area appears bare and unvegetated (Cao et al., 2009). Specifically, samples for this 86 87 study (sampling points P1 and P3 in Fig. 1) were collected from two different experimental plots 88 used for a previous revegetation test with endemic species (Pistacia lentiscus L.; Bacchetta et al., 89 2012). P1 samples are from a plot representing the untreated waste material, while P3 samples are 90 from a plot where some compost was added. Since planting in 2008, P. lentiscus survived only in 91 the P3 plot (Fig. 2).

92 The second and third sampling areas are two natural sites, not visibly affected by morphological 93 alterations due to mineral extraction, located in the natural substrate of Cambrian limestones and dolostones, that host the metalliferous bodies (sampling points P5 - Serra Merareddu, and P6 -94 Pozzo Santa Barbara - Fig. 1). These sites, characterised by natural geochemical anomalies 95 96 (particularly in Zn and Pb), though not intended to statistically represent the whole district, can give 97 an idea of pre-mining conditions. In both sites P. lentiscus grows spontaneously. The fourth site is a 98 lagoon deposit at Sa Masa, near the Funtanamare beach (sampling point P9 in Fig.1), final collector 99 of waters and pollutants from the entire watershed of the San Giorgio river (including mining 100 seepages, and sewage waters from the Iglesias town). Therefore, the high contamination at Sa Masa 101 (see below) is not just site specific, but induced by transportation of solid and liquid matrixes from 102 other localities within the San Giorgio basin. Even at Sa Masa, P. lentiscus grows spontaneously.

As previously noted, the climate of the area is typically Mediterranean, characterised by very dry summers. The maximum annual average temperature is 25°C, and the minimum 5°C. Average annual precipitation is ca. 800 mm (<u>http://my.meteonetwork.it/station/srd025/statistiche).</u>

106

107 2.2 Soil characterization

108 At each study site, a soil profile was opened and described, making a distinction in pedogenetic 109 horizons according to standard procedures of soil description (Schoeneberger et al., 2002); samples 110 of each horizon were collected for subsequent laboratory analyses. Even though Campo Pisano is 111 not classifiable as natural soil, for descriptive purposes the terms 'soil' and 'soil horizon' are also 112 used for that site, where the substrate is in fact mainly composed of mine tailings. This study 113 considers data from shallow, unsaturated horizons, according to the lysimeter or rhizometer depths 114 used for field soil pore-water extraction (see below). The complete set of soil data is available in 115 Concas (2014).

116 Bulk soil samples were air-dried and sieved (<2 mm, fine earth). Routine physical and chemical 117 analyses were carried out according to the procedures published by the Italian Ministry of Agriculture (MiPAF, 1998, 2000). Sand (2.00–0.05 mm), silt (0.050–0.002 mm) and clay (<0.002 118 119 mm) fractions were separated after the removal of organic matter by H₂O₂ treatment and dispersion 120 aided by Na-hexametaphosphate by the sieve and pipette methods. The soil pH was measured in a 121 1:2.5 soil/water suspension. The organic carbon (OC) was estimated by wet digestion with a 122 modified Walkley-Black procedure. Soils were classified according to IUSS Working Group WRB 123 (2014). Mineralogical analyses were performed on the <2 mm fraction by powder X-ray diffraction 124 (PXRD), on an automated Panalytical X'pert Pro diffractometer with Ni-filter Cu-K_{a1} radiation ($\lambda =$ 125 1.54060 Å), operating at 40 kV and 40 mA, using the X'Celerator detector (see 4.1 paragraph). The 126 main soil chemical components (major element oxides for ordinary rocks) were also determined on 127 the <2 mm fraction by a Philips PW1480 wavelength-dispersive X-ray fluorescence (WDXRF). For 128 the A horizon at the P9 location, it was not possible to obtain a suitable pellet for XRF analysis, 129 because of its loose incoherent texture. Information on its 'major oxide' content is not critical for 130 the purposes of this paper, and we made no further efforts to obtain these data. Raw intensities were corrected for matrix effects by the method of Franzini & Leoni (1972), making reference to 131 132 certified rock standards. The method is calibrated for ordinary rocks, with comparatively low heavy

133 metal contents. The correction routine does not adequately account for the unusually high contents of Pb and Zn typical of most of our samples. Therefore, XRF data should be taken as semi-134 135 quantitative. Loss of ignition (LOI) was determined by leaving an aliquot of the sample at 900°C 136 for 8 hrs, and expressed as percent weight difference before and after the treatment. Contributions 137 to LOI include carbonate decomposition, dehydration of clay minerals and/or Fe-Al oxyhydroxides, 138 and burning of organic matter. The total heavy metal contents of soil samples were determined on 139 the <2 mm fraction after digestion in microwave system (Milestone Ethos 1 oven) with a 140 HNO₃/HF/HClO₄ (5:2:1 v/v) mixture, as proposed by Brunori et al. (2005), analysing the obtained 141 solutions by an inductively coupled plasma optical emission spectrometer (ICP-OES Analyzer, 142 ARL Fisons 3520B). Soil samples were randomly digested and analysed in replicate to test the 143 precision of the method. In this case the average datum of the two aliquots was considered. The 144 accuracy was tested against the certified standard NIST 2710. Overall, heavy metal analyses in soils 145 are within +10% for both accuracy and precision.

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147 2.3 Soil pore water sampling and analyses

At each study site soil pore water samples were collected from unsaturated profiles, in the rooting zone. In order to reduce as much as possible soil disturbance during sample collection SPS 200 soil water samplers (SDEC, France) were used (Lekakis et al., 2015). The samplers, namely 'vacuum lysimeters', are characterised by a porous ceramic cup glued on the bottom of an empty PVC tube (31 mm in diameter), and are designed to replicate the suction function of plant roots.

The samplers were placed in all sampling points at 10-15 cm depth into the upper soil horizon, and, when possible, at 30-40 cm depth into the underlying horizon. At P1 and P3, only short lysimeters were used, because of the shallow depth of soils. Before starting the sampling, the lysimeters were left for 1 month in the field to equilibrate. One day before sampling, a vacuum of -75 kPa was created inside the tube with a vacuum pump (SPS400, SDEC, France), through a connecting tube hosted in a pierced rubber bung, in order to draw the pore water from the soil to the porous ceramic

159 cup and into the tube. To sample the pore water, the vacuum pump was used to collect the soil 160 solution contained in the tube, which was then poured into a sampling flask. Because of the climate 161 of the area, with very dry summers, four surveys were carried out between the rainiest period, 162 corresponding to the winter, and the starting of the dry season (from December 2012 to April 2013). 163 In fact, fall 2012 was unusually dry, and at the time of the first sampling (December 2012) it was 164 possible to obtain enough liquid for analysis only at Campo Pisano and Sa Masa. In the following 165 months (January - February 2013), there was enough rain infiltration to allow collection of 166 sufficient pore water from most lysimeters. In April 2013, with the beginning of the dry season, 167 only at Sa Masa, where the water table was always shallow, it was possible to collect soil pore 168 water. At this site in April 2013 three additional samples were collected using another type of 169 tension sampler, namely the 'rhizon sampler' or 'rhizometer' (Eijkelkamp Agrisearch Equipment, 170 The Netherlands) (Tye et al., 2003; Di Bonito et al., 2008). Using this device, pore water solution is 171 extracted by inserting it into the soil, and connecting a syringe for applying the vacuum and then a suction. Sampling points were two at Campo Pisano (sampling points 1 and 2, respectively at P1 172 173 and P3), three at the Serra Merareddu (P5) site (sampling points 6a, 6b and 6c), where letters a, b 174 ...indicate that lysimeters were positioned all around the same *P. lentiscus* individual; three at the Pozzo Santa Barbara (P6) site (sampling points 7a, 7b and 7c), and eight at the Sa Masa (P9) site, 175 176 respectively five sampled by lysimeters (sampling points 4a, 4b, 4c and 5a, 5b), two by rhizometer 177 (sampling points 19 and 22), and one sampled by both (sampling point 25).

Pore waters collected with lysimeters were filtered to 0.45 μ m; for solutions extracted by rhizon samplers this filtering is not necessary, because their porosity is less than 0.45 μ m. Labile parameters (temperature, pH, conductivity) were measured in the field. Because of the scarcity of collected liquid, Eh was not routinely determined; in the few samples where it was measured, the values indicate oxidising conditions (equilibrium with the atmosphere), as expected for unsaturated horizons. The solutions were stabilised to 1% v/v with HNO₃ for the subsequent analyses of major and trace elements, including heavy metals. Whenever enough solution was available, total

185 alkalinity was determined independently by HCl titration on another aliquot. The major elements were measured by Ion Chromatography, and the trace elements (Zn, Pb, Cd, Mn, Fe, Cu) by ICP-186 187 OES (Perkin Elmer Optima 2100 DV). Because of solution scarcity, anions were determined on the 188 same acidified aliquot used for cations; obviously, this made impossible the measure of nitrate in 189 solution. In Sardinia, high (>50 mg/L) nitrate content in groundwaters is mostly linked to 190 agriculture (RAS, 2010), which is not the land use of any of the studied sites. Moreover, ionic 191 charge balances (see below) do not suggest anion deficit in any of the analyzed pore waters. In 192 some cases (data marked with * in Tables 3a and 3b), when the solution from a single sampling was 193 not sufficient for the analyses, pore waters from two close sampling points were combined.

194 The dissolved organic carbon (DOC) was determined by TOC/TN analyser (Analytical Sciences 195 Thermalox program), a specific instrument for total organic carbon and nitrogen contents in water 196 solutions. The instrument actually determines the total carbon (TC); whereas the dissolved 197 inorganic (carbonate) carbon (DIC) is removed by acid addition, and the remaining DOC is 198 measured by a second pass in the TOC analyser. The inorganic carbon is calculated as the 199 difference between TOC and DOC. When it was not possible to collect enough pore water for an 200 independent determination of alkalinity, the value of DIC was used to calculate alkalinity as HCO_3^{-1} . 201 Quality control of pore water chemical analyses is intrinsically difficult. To the best of authors' 202 knowledge, there are no specific certified international standards, and replicate analyses were 203 seldom possible. Preliminary release tests with deionised water on new lysimeters and rhizometers 204 indicated negligible (below detection limits for ICP-OES) amounts of released elements, except for 205 Ba. For this element, high values were recorded in lysimeter blanks, therefore data for Ba were not 206 included in this study. Field blanks were obtained by introducing deionised water in the same 207 devices used for pore water samples, and treating the collected volumes as independent samples. 208 The accuracy of ICP-OES analyses was checked with calibration standard solutions. A further 209 check of quality of analysis for major elements are ionic balances. From data in Table 3b, it is

210 possible to calculate ionic balances: most are within $\pm 10\%$, and only two are within $\pm 15\%$. Overall, 211 it is estimated that most of analyses are within $\pm 20\%$ error for both accuracy and precision.

The collected data were elaborated with two different speciation programs, PHREEQC 3.0.2 (Parkhurst et al., 1980; Parkhurst, 1995) and WHAM 7.0.2 (Tipping, 1994); specifically, the second accounts for complexation of metals by organic matter.

- 215
- 216 **3. Results**
- 217
- 218 *3.1 Soil characterization*

Field observations, summarized in Table 1, showed that (i) the soil derived from untreated waste 219 220 material (P1) is shallow, and characterized by a very thin cemented and hardened surface horizon 221 overlying two mineral horizons slightly enriched in organic carbon (OC); (ii) the soil derived from the waste material treated with compost (P3) is shallow, and characterized by a ploughed epipedon 222 223 enriched in OC; (iii) soils derived from dolomitic limestones (P5 and P6) are shallow or moderately 224 deep, with A horizon at surface and a well expressed Bw horizon, and lithic contact toward the underlying bedrock; (iv) the soil derived from Holocene lagoon deposits at Sa Masa (P9) is deep, 225 226 and characterized by a sequence of different sedimentary cycles (marked by differences in texture).

The differences in texture among the soils (Table 1) reflect their origin from different parent materials, and confirm the field assessment. Reaction is generally neutral to slightly alkaline, with the exception of the strongly acid 2^C horizon of P1 and P3, and the moderately alkaline 5C4 and 6C5 horizons of P9. OC contents are highest in the surface horizons and decrease regularly with depth.

In the World Reference Base (WRB), the studied soils classify as follows: Spolic Technosol (P1
and P3), Endoleptic Luvisol (Abruptic, Humic, Clayic, Rhodic) (P5), Epileptic Luvisol (Abruptic,
Humic, Chromic) (P6), and Haplic Fluvisol (Calcaric, Ruptic) (P9).

9

The main physicochemical characteristics of the studied soils are reported in Table 1, while mineralogical composition is described hereafter. At Campo Pisano (P1 and P3), tailings mineralogy is dominated by the presence of quartz, gypsum and carbonates (mainly dolomite, secondarily calcite and traces of ankerite), with minor muscovite and microcline, and residual pyrite. Hydrozincite, cerussite, as well as traces of sphalerite and galena, were also detected in some horizons.

Samples from Serra Merareddu (P5) and Pozzo Santa Barbara (P6) contain a clay component (kaolinite and montmorillonite), as expected in *sensu stricto* soils. The main mineral at both sites is however quartz (possibly of aeolic deposition: Concas, 2014). Dolomite is the predominant carbonate; barite occurs in all profiles. At Sa Masa (P9), carbonates (calcite, dolomite, and subordinate ankerite) and quartz represent the main minerals. Minor minerals include hemimorphite, barite, and cerussite. As observed above, at Sa Masa some mineral component could result from transport from other localities.

The main chemical composition (Table 2a) is substantially in agreement with the mineralogy of 248 soils. High LOI and CaO values in samples from Campo Pisano and Sa Masa are indicative of the 249 250 presence of a major carbonatic fraction, while in P5 and P6 profiles the prevalence of the quartz and phyllosilicate fraction is well pinpointed by high values of SiO₂ and Al₂O₃. Iron (as Fe₂O₃) is 251 significant, as expected, in the tailing samples from Campo Pisano (P1 and P3), where, other than in 252 253 residual pyrite or ankerite, it could be also present as poorly crystalline Fe-oxide-hydroxides from sulphides oxidation, such as ferrihydrite (Blowes et al., 2003; Marescotti et al., 2012; Ardau et al., 254 255 2014), not easily identifiable by PXRD analyses in complex matrixes; the site with the lowest Fe is 256 Sa Masa.

Among heavy metals (Table 2b), Zn concentration is around thousand mg kg⁻¹ at Pozzo S. Barbara (P6), from thousand to ten thousand mg kg⁻¹ at Campo Pisano (P1 and P3) and Serra Merareddu (P5), from thousand to hundred thousand mg kg⁻¹ at Sa Masa (P9). Lead concentrations vary from thousand mg kg⁻¹ at Campo Pisano and Pozzo S. Barbara, and from thousand to ten thousand at Serra Merareddu and Sa Masa. Cadmium and Cu are present in amounts around tens mg kg⁻¹ in all
sites, except at Sa Masa where concentrations can reach hundreds mg kg⁻¹.

As it can be observed from chemical results, the natural soils at P5 and P6 sites show heavy metal 263 264 contents of the same order of magnitude as the mining flotation waste at Campo Pisano. Some 265 values are in fact higher than median values of stream sediments in the district (Boni et al., 1999), which can be taken as an indication of the local post-mining geochemical baseline. Concas (2014) 266 267 reports that at these sites the rock substrates (R layer) show appreciably lower metal contents, 268 suggesting that either the sites are in some way contaminated by anthropic activities (e.g., by wind 269 transport from nearby mine sites), and/or pedogenetic processes led to a (passive) enrichment in 270 some heavy metals. This issue is beyond the purpose of this paper, and it will not be further 271 discussed.

272

273 *3.2 Pore water chemistry*

The overall major element chemistry of pore water samples, distinguished for location, is depicted in the Piper diagram of Fig. 3. Tables 3a and 3b and 4a and 4b report the main physical and chemical parameters (including major and selected trace components) of the studied pore waters, divided for location and sampling periods.

Fig. 3 points out a clear difference among pore water major chemistry of Campo Pisano samples (P1 and P3) from the natural substrate of Cambrian limestones and dolostones (P5 and P6), and from Sa Masa (P9). In terms of descriptive hydrochemical facies (e.g., Back, 1966), the Campo Pisano waters belong to a Ca-SO₄ type, those of P5 and P6 to a Mg-HCO₃ type (with a subordinate Cl component for some samples), and those at Sa Masa again to a Ca-SO₄ type, but with a significant Na-Cl component, presumably reflecting a marine input, either from seawater ingression or from sea spray deposition (see further discussion).

285 Conductivity of pore waters is highly variable, ranging from ~570-900 μ S cm⁻¹ in samples from the 286 natural substrate, ~1200-2000 μ S cm⁻¹ at Campo Pisano, up to ~2400-3700 μ S cm⁻¹ at Sa Masa, indicating that the overall amount of soluble minerals, influencing pore water components, is rather different in the three sites. Moreover, at Sa Masa a direct influence of the sea water component (see below) in the high conductivity is likely. The pH of pore waters shows a moderate range of variation (from circumneutral to alkaline) for different sites and/or different sampling times.

291 With respect to trace elements, zinc presents the highest concentration at Sa Masa, in the order of thousand $\mu g L^{-1}$, in particular at the deepest levels (up to 5700 $\mu g L^{-1}$, with the only exception of the 292 December 2012 sampling, when Zn does not exceed hundreds $\mu g L^{-1}$). At Campo Pisano, Zn 293 concentrations are around hundreds $\mu g L^{-1}$ (no substantial differences are observed between plots 294 P1 and P3). As observed at Sa Masa, concentrations are one order of magnitude lower during 295 December 2012 sampling. The lowest Zn concentration (averagely of tens μ L⁻¹) is recorded at P5 296 297 and P6, where December 2012 sampling was not possible for the absence of sufficient water (see § 298 2.3).

Lead concentrations are in the order of tens μ g L⁻¹ in all sampling sites, in agreement with the known low mobility of this metal at pH values >7. The highest concentrations are registered in January 2013 at Serra Merareddu, which is also the site with the highest total Pb values in the soil. Manganese concentrations may show a certain variability in samples of the same site (e.g. 4a, 4b, 4c); in general, this metal is more abundant at Sa Masa (up to hundreds μ g L⁻¹). At Campo Pisano Mn does not exceed tens of μ g L⁻¹, while it is close or below the limit of detection in the natural soils (P5 and P6).

306 Iron does not show high mobility, in spite, for example, of significant bulk soil concentrations at 307 Campo Pisano. In fact, as expected at pH values measured in the pore waters, Fe results strongly 308 bound to the solid fraction of the soil (e.g. Ardau et al., 2009), and therefore, often below the 309 detection limit in the pore water. When detectable, the saturation index (SI^1) modelling results (not 310 shown) indicate high oversaturation with respect to the most abundant Fe-oxide-hydroxide in

¹ Saturation index (SI) = log (IAP/K_{sp}), where IAP (ion activity product) is the product of the chemical activities of the dissolved ions of the mineral, and K_{sp} is the solubility product. SI shows whether a particular mineral is in equilibrium with solution (SI null), or will tend to dissolve (SI negative), or precipitate (SI positive)

311 mineralised areas, such as goethite or ferrihydrite. Iron is also the metal showing the highest local 312 variability as a function of sampling period (see for example P1), indicating a strict seasonality 313 relationship.

314 As expected from their comparatively low bulk soil concentrations, Cd and Cu are definitely315 subordinate in pore waters of all sites.

The OC content varies from undetectable ($<0.04 \text{ mg L}^{-1}$) up to 12.6 mg L⁻¹ at Campo Pisano. OC contents in pore waters are not clearly related to bulk OC in soils. Presumably, this finding reflects different degrees of solubility of organic matter in the different sites.

319 Speciation equilibria, as calculated by PHREEQC and WHAM, are summarised in Table 5. 320 Calcium and Mg are mostly present as free ions (average 70-80% in all samples). In the natural 321 substrate (P5 and P6), Ca and Mg also show a subordinate association with carbonate species (CO₃) 322 and HCO₃), while in sulphate-rich pore waters (Campo Pisano and Sa Masa) a subordinate 323 association with SO₄ is shown. As a general rule, all samples reach equilibrium or oversaturation 324 conditions with respect to calcite and dolomite (Fig 4a,b). Gypsum equilibrium is close to be 325 attained for samples from Sa Masa and Campo Pisano, while samples from natural substrate result 326 undersaturated with respect to it (Fig 4c).

Trace metals show a variety of species, the abundance of which is related either to the main water composition, or to the specific behaviour of the element. $PbCO_3$, for example, is generally the most abundant Pb species both in carbonate and in sulphate waters; a subordinate species for this metal is PbOH⁺. All samples are close or in equilibrium with cerussite (Fig. 4d), and undersaturated with respect to anglesite (Fig. 4e).

332 Zinc mainly occurs as Zn^{2+} and subordinately as $ZnSO_4$. Several samples are close to equilibrium 333 with hydrozincite, but a few are strongly undersaturated (Fig. 4f). Hydrozincite was in fact 334 identified by PXRD in some samples. Smithsonite (Fig. 4g) is consistently undersaturated, but a 335 number of samples at Sa Masa are close to equilibrium with it. This mineral was never positively 336 recognized in the studied samples, but its widespread occurrence in the district is well documented. 337 Specifically, smithsonite has been recognized both in the 'red muds' tailings at Monteponi and at 338 Campo Pisano, often in association with hydrozincite and hemimorphite (the so-called 'calamine' 339 ores of the Iglesiente mining district were typically composed of Zn (hydroxy-) carbonates and 340 silicates; Boni et al., 2003). Runoff from these wastes may well be transported to Sa Masa by the 341 San Giorgio river.

Manganese occurs as free ion, or as ion pair with SO_4 and CO_3 . Several pore waters are close to or above saturation with rhodocrosite (Fig. 4h). This phase was not recognized in the studied samples; alternatively, the phase(s) controlling Mn in pore waters could be solid solution(s) with calcite and/or dolomite.

Cadmium mainly occurs as $CdCl^+$ and Cd^{2+} , and copper as $CuCO_3$ and $CuOH^+$; moreover, this metal is the only one for which the fraction bound to colloidal fulvic acids (as calculated by the WHAM program) is important (up to 99%). It is difficult to identify the possible phases controlling the mobility of Cd and Cu, since these elements are presumably present as minor substituting ions both in primary and secondary phases, therefore they do not appear in mineral theoretical formulas as identified by the speciation program.

352

353 **4. Discussion**

354 This study supports field monitoring of soil pore water chemistry as a useful method to evaluate the environmental risk associated to the mobility of (toxic) elements. With respect to indirect methods 355 356 (selective and/or sequential extractions), it is comparatively rapid, because it does not require the extraction step(s); moreover, it directly records field conditions, that may imply different metal 357 358 mobilities in the same soil (as shown in this study by differences in sampling events at the same sites). An additional important aspect is that pore water chemistry well reflects the 'history' of the 359 associated soil/waste, being not merely the expression of the local mineral and chemical 360 composition of soil/waste, but also of its 'use', geographic and climatic context. In fact, from pore 361 water data treatment emerged some distinctive marks to distinguish among the natural mineralised 362

363 soils, the mine waste 'soils', and the soil from a 'collecting' area, selected for this study. If mineralogical composition is generally the main factor influencing pore water chemistry, an 364 365 interesting exception, linked to the geographic context, is represented by Sa Masa swamp soil. At 366 Sa Masa, the influence of the near sea in the main chemistry of pore water overlaps that of soil 367 mineralogy, while trace elements show the mark of the accumulated material from different mined 368 areas present in the S. Giorgio basin, in particular for the high presence of Zn, which turned out to 369 be the most mobile metal. Sa Masa site represents, therefore, a clear example on how the mere use 370 of the total metal content in order to evaluate potential environmental risk of a contaminated soil 371 may offer indication far from those obtained by the use of pore water.

At Campo Pisano (P1 and P3), we are not in presence of a *sensu stricto* soil, but the substrate is composed of mining wastes. This nature of the 'soil' is well pointed out by a main chemistry dominated by Ca-SO₄, reflecting the abundant presence of gypsum, secondary product of sulphide oxidation (as a source of sulphate), and carbonates (as a source of calcium). The heavy metal contents reflect their mobility at the pore water pH values (for example with a prevalence of Zn with respect to Pb) and of their solid-speciation (see below).

378 In the natural substrate of Cambrian limestones and dolostones (P5 and P6), pore water chemistry 379 well marks the difference with respect to mined areas, mainly for a limited conductivity, indication 380 of a reduced reactivity of mineral phases. However, the field pore water is perfectly able to identify 381 the presence of a geochemically anomalous area. The main chemistry is dominated by the carbonatic nature of the substrate, while sulphates, indicative of possible sulphide presence and of 382 383 their subsequent oxidation, are limited. The lower contents in Zn in pore waters from this site with 384 respect to Campo Pisano well reflect a limited reactivity of minerals in an undisturbed soil with 385 respect to a mining waste dump. Lead is an exception since, regardless of its total concentration in 386 soils, its pore water concentration is similar and comparatively low for all sites, probably because of saturation with respect to cerussite. 387

388 In sites where lysimeters were positioned in more than one point (see paragraph 2.3), pore waters 389 variations in different sampling points are not significant, with the only exception of Mn. Variations related to the sampling period were more significant. Salinity at Sa Masa was higher in December 390 391 2012, and lower in February and April 2013. As shown in Fig. 5, this trend of salinity variation is 392 matched by seasonal variations of both Na and Cl (and of SO_4 – not shown), suggesting that sea 393 water contributions prevail at this site. For lagoon deposits like Sa Masa, contributions of seawater 394 may be linked either to marine ingression, and/or to dissolution and infiltration of sea spray 395 deposited on the soil surface. Both phenomena are usually more pronounced during dry periods. The input of sea salts from marine aerosol in groundwater (and surface water) close to the coast is a 396 397 well known phenomenon due to the dissolution by the rain of the particulate deposited on the 398 ground, and subsequent infiltration across the soil (e.g., Whipkey et al., 2000; Lorrai & Fanfani, 2007, and references therein). The current data may thus suggest that either marine ingression. 399 400 and/or sea spray infiltration increase the pore water marine component during drier seasons, whilst 401 pore water is subjected to a sort of 'dilution effect', affecting sea water components (and salinity) 402 during rainy periods, due to a washout of the first soil centimetres, and transportation of the soluble 403 sea component toward deeper horizons.

404 Compositional variations are also registered in the other sampling sites, as for example, Campo 405 Pisano. In this case the element variations are more linked to precipitation/dissolution of soluble 406 salts, effects of dilution, or possibly interaction time influence. In fact, under comparatively dry 407 conditions, circulation of soil fluids is not pervasive, and the system does not approach an 408 equilibrium state; another important aspect to be considered is that Zn, an essential micronutrient, is 409 constantly taken up by plants (including *P. lentiscus* – see below). Plant physiology is therefore at 410 the same time influencing and influenced by element concentration in soil pore water.

411 An added value of field pore water chemistry monitoring is also the possibility to perform element 412 speciation in solution, offering a further 'tile' for the understanding of interface processes between 413 hydrosphere and biosphere. In fact, as pinpointed by other authors (e.g. Moreno-Jiménez et al., 414 2011 and references therein), the soluble fraction will interact differently with plant roots as a 415 function of element speciation; for instance Shahid et al. (2012, 2014) observed how organic ligands are capable to modify Pb speciation by forming organo-metallic complexes of different 416 417 bioavailability. In this study we observe how the same concentration of some element in pore water 418 (e.g. Zn) could be characterised by differences in speciation at different sites/times. Though beyond 419 the purpose of this study, the understanding of how aqueous speciation may influence metal uptake by P. lentiscus could be an interesting subject of further investigation. The associated use of 420 421 PHREEQC and WHAM programs also permitted to identify which heavy metals are preferentially 422 present as free ions or bound to colloidal fulvic acids, and possible speciation differences from site 423 to site.

424 In general, from equilibrium calculations, the soil/pore water contact time appears sufficient for reaching equilibrium conditions with several mineral phases, mostly carbonates, present in the soil 425 426 (e.g. Ca-Mg carbonates, hydrozincite, cerussite, gypsum). This allows to evaluate which minerals 427 are mostly responsible for controlling element mobility. This control is not expected to vary in a 428 short time. However, if mineral abundance is expected to vary, e.g. because of alteration, it is 429 possible to hypothesise a change of pore water quality in medium/long-time term. This is 430 particularly true for 'soil' constituted by residual mining material as in Campo Pisano, more 431 subjected to rapid alteration.

A potential limitation of this field-based approach could be that, in Mediterranean climates, during dry periods, pore water is not enough to be sampled, dictating and altering the sampling period distribution depending on climatic conditions. However, information on variations due to seasonal, climatic and morphological conditions is critical for the understanding of the physiological response of plants in the geographic context in which they occur. Laboratory-based methods fail to supply such information, and therefore long-term field studies, such as soil pore water monitoring, are required. 439 In terms of environmental risk, the concentrations of metals in pore waters can be usefully compared with limits established by Italian laws (D.lgs 152/2006) for subsurface waters (Tables 4a 440 and 4b). These limits are exceeded for Pb in most samples, and in many samples at P9 also for Zn, 441 442 Mn, and in three samples also for Cd; Mn limits are exceeded in some samples at Campo Pisano 443 (P1 and P3). Therefore, based on present data, there is a potential impact for heavy metals in the 444 area, especially at Sa Masa. It is interesting to notice that Campo Pisano, considered one of the 445 'hazard centres' of the district, does not show the highest levels of heavy metals neither in soils, nor 446 in pore waters. On the other hand, with respect to the other studied sites, the absence at Campo 447 Pisano of a vegetation cover limiting wind and water erosion makes this site a potential centre of 448 mechanical dispersion of contaminants.

449 Considering the potential mobilization to the biosphere, Concas (2014) showed that Pb is much less mobile than Zn with respect to uptake by P. lentiscus. This limited Pb uptake was ascribed to a 450 451 combination of the lower Pb mobility, and of a physiological mechanism of exclusion (Zn is an essential micronutrient, whereas Pb has no known physiological function, and may have phytotoxic 452 453 effects - Shahid et al., 2012). Concas (2014) also showed that Pb and Zn uptake of P. lentiscus 454 varies seasonally. Since field pore water and plant sampling did not occur at the same time (plants 455 were sampled in May and September 2012), it is not possible to compare directly plant uptake and 456 metal concentration of the pore water. Moreover, plant uptake is clearly influenced also by the total 457 amount of available pore water, and not only by metal concentrations. This aspect could well be worthy a future step of investigation. The authors notice here that this kind of information could be 458 459 useful for the management of a mining waste dump subjected to phytostabilization. For example, a 460 water-saving watering system could be planned in specific periods of the year in order to modify pore water composition, to obtain a decrease of some element when necessary. As a consequence of 461 462 Zn mobilization to the biosphere, there is a potential risk that this metal could enter the trophic chain. On the other hand, phytostabilization actions using P. lentiscus as a pioneer species should 463

464 not be impaired, inasmuch as this species is well tolerant to Zn (Bacchetta et al., 2012; Concas,
465 2014).

466

467 **5.** Conclusions

468

469 Field sampling and chemical monitoring of soil pore water in the major abandoned mining district
470 of Iglesiente (SW Sardinia) confirmed that pore water is a rapid and easy method to collect reliable
471 information on the potential risk of a contaminated soil.

Soil pore waters chemistry in the studied sites reflects the mineralogy of the local soil, and suggests
an approach to equilibrium with coexisting solid phases. This aspect permits to easily evaluate
which minerals are likely responsible for controlling (toxic) element mobility.

475 With respect to laboratory tests on metal mobility, the analysis of pore water permits to evaluate 476 trace elements mobility, and the consequent environmental risk, also following pore water seasonal 477 changes both in qualitative and quantitative terms. Pore water represents the most available fraction 478 of the overall soil matrix for plant uptake, therefore allowing a direct comparison with possible 479 seasonal changes in metal uptake by plants. In fact, in a companion study of the local species P. 480 *lentiscus*, local and seasonal variations of metal (Zn, Pb and Hg) concentrations in different parts of 481 the plant were detected. Such variations may be partially a response to local/seasonal changes of 482 pore water quantitative/qualitative characteristics. This kind of information could be useful for the 483 management of a mining waste dump interested by phytostabilization.

484

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661	Table captions
662	
663	Table 1. Selected soil properties at the sampling sites
664	
665	Table 2a and 2b. Major and selected trace element analysis of soils at the sampling sites,
666	distinguished for horizons
667	
668	Table 3a and 3b. Pore water parameters (T°, pH, conductivity) and major chemistry distinguished
669	for sampling site, sampling time, and sampling method.
670	
671	Table 4a and 4b. Selected trace elements in pore waters distinguished for sampling site, sampling
672	time, and sampling method. Bold font indicates values above limits established by Italian laws
673	(D.lgs 152/2006) for subsurface waters.
674	

Table 5. Main element speciation in pore waters as determined by PHREEQC and WHAM codes.

- 676 Figure captions
- 677

678 Figure 1. Schematic map of the study area, with location of sampling points.

679

680 Fig. 2. a) general view of the Campo Pisano experimental plots established in 2008 by Bacchetta et

- al. (2012), pointing those selected for this study; b) and c) vertical sections of 'soils' in Plot 1 and
- 682 Plot 3; in the first (untreated material) *P. lentiscus* planted in 2008 did not survive, whereas several
- 683 individuals survived in Plot 3, where compost was added as amendment.
- 684
- Fig. 3. Piper diagram illustrating the variability in major ion composition of the studied pore waters.
- Fig. 4. Saturation indexes for selected minerals, distinct for sampling site (symbol), and ordered for

688 sampling time (from right to left). The solid line (SI = 0) indicates equilibrium between mineral and

689 solution. See text for the assessment of the role of selected minerals from (a) to (h) in controlling

- 690 element mobility in pore waters.
- 691

Fig. 5. Na vs Cl plot of pore water samples from Sa Masa site (P9). The linear trend suggests a co-variation of the two elements due to seasonal variations.

Table	1
1	-

		Depth		Texture		Textural	pН	OC
Soil	Horizon	-	Sand	Silt	Clay	class ^a	(H_2O)	
		cm		%		_		g kg ⁻¹
P1	^ACm	0-0.3/0.5	77.7	20.9	1.4	ls	7.6	11
	^AC1	0.3/0.5-8/14	82.5	16.7	0.8	1 s	7.5	9
	^AC2	8/14-16.5/20.5	69.6	29.3	1.1	sl	7.3	14
	2^C	16.5/20.5->30	68.4	28.3	3.3	sl	4.5	2
P3	^Ap1	0-1.5/1.8	73.8	24.8	1.4	ls	7.2	61
	^Ap2	1.5/1.8-15/17	77.9	21.3	0.8	1 s	7.1	66
	2^C	15/17->22	66.8	31.1	2.1	sl	4.8	5
P5	А	0–15	44.1	41.6	14.3	1	7.3	43
	AB	15-28	30.7	39.9	29.4	cl	7.3	27
	Bw	28-60/65	13.6	26.6	59.8	с	7.3	11
	R	>60/65						
P6	А	0–10	44.1	43.8	12.1	1	7.1	37
	Bw	10-37/50	31.5	41.2	27.3	cl	7.5	18
	R	>37/50						
P9	А	0–8	47.2	49.8	3.0	1	7.5	25
	2C1	8-12	18.4	76.2	5.4	sil	7.8	9
	3C2	12–25	73.6	24.6	1.8	ls	7.8	6
	4C3	25-40	53.0	45.9	1.1	sl	7.8	5
	5C4	40–59	34.1	63.6	2.3	sil	8.0	6
	6C5	59->120	86.9	12.1	1.0	ls	8.1	5

^a ls = loamy sand; sl = sandy loam; l = loam; cl = clay loam; c = clay; sil = silt loam.

Soil	Horizon	Depth (cm)	LOI	Na ₂ O	MgO	Al2O ₃	SiO ₂	P_2O_5	K ₂ O	CaO	TiO ₂	MnO	Fe ₂ O ₃	Other*	Tot
		(cm)	%	%	%	%	%	%	%	%	%	%	%	%	%
P1	^ACm	0-0.3/0.5	28.84	0.40	13.9	3.81	14.3	0.04	0.59	23.2	0.13	0.24	13.5	1.52	100
	^AC1	0.3/0.5-8/14	26.69	0.33	13.5	4.13	14.5	0.04	0.60	25.0	0.12	0.25	10.7	1.47	97.0
	^AC2	8/14-16.5/20.5	24.25	0.61	15.2	2.84	11.9	0.04	0.53	26.0	0.13	0.23	17.0	1.69	100
	2^C	16.5/20.5->30	13.70	0.25	0.50	2.57	9.21	0.03	0.49	23.9	0.12	0.02	48.5	0.79	100
P3	^Ap1	0-1.5/1.8	25.32	0.41	5.43	5.71	21.9	0.89	0.93	20.6	0.16	0.20	17.6	1.12	100
	^Ap2	1.5/1.8-15/17	24.71	0.52	9.46	4.28	18.9	0.25	0.82	23.2	0.16	0.17	16.8	1.00	100
	^2C	15/17->22	12.75	0.11	0.18	2.25	7.67	0.05	0.40	25.8	0.09	0.02	50.2	0.65	100
P5	А	0-15	12.56	0.23	1.19	20.8	52.1	0.15	1.92	0.62	0.44	0.49	8.15	1.55	100
	Ab	15-28	10.65	0.25	1.41	21.3	50.9	0.13	2.27	0.47	0.53	0.51	9.79	1.94	100
	Bw	28-60/65	10.50	0.22	1.32	26.9	44.7	0.08	2.59	0.34	0.45	0.40	11.7	0.90	100
P6	А	0-10	11.43	0.60	1.96	16.9	54.0	0.15	2.84	0.95	0.81	0.96	8.91	0.61	100
	Bw	10-37/50	8.51	0.49	1.80	21.2	52.8	0.12	2.79	0.60	0.76	0.87	9.38	0.68	100
P9	2C1	8-12	28.89	0.65	5.33	3.28	23.1	0.04	0.39	32.4	0.17	0.18	3.68	2.69	100
	3C2	12-25	31.68	0.48	6.61	2.48	18.2	0.03	0.25	35.7	0.17	0.22	2.71	2.23	100
	4C3	25-40	33.48	0.41	6.61	1.17	14.4	0.02	0.19	40.1	0.09	0.22	2.23	1.73	100
	5C4	40-59	35.37	0.46	7.99	1.30	10.5	0.02	0.19	38.7	0.09	0.24	3.89	1.96	100
	6C5	59->120	31.77	0.62	8.22	1.31	10.3	0.02	0.21	35.2	0.09	0.20	9.24	4.13	100

* Except the A horizon at the P9 location – see text

*Sum of elements in Table 2b

Table 2a

Profile	Horizon	Depth (cm)	Cd	Cu	Ni	Pb	Zn
					mg/kg		
	^ACm	0-0.3/0.5	70	52	< 16	2180	12932
D1	^AC1	0.3/0.5-8/14	72	44	< 16	2180	12402
PI	^AC2	8/14-16.5/20.5	78	92	< 16	2126	14646
	2^C	16.5/20.5->30	15	56	< 16	2698	5119
	^Ap1	0-1.5/1.8	54	54	< 16	1760	9360
P3	^Ap2	1.5/1.8-15/17	54	51	< 16	1566	8327
	^2C	15/17->22	16	57	< 16	2776	3666
	А	0-15	6	56	39	13642	1779
DC	Ab	15-28	<4	54	39	17697	1655
P5	Bw	28-60/65	<4	56	53	7134	1822
	R	60/65->60/65	10	10	< 16	1754	426
DC	А	0-10	30	37	41	1360	4680
Po	Bw	10-37/50	30	31	44	1453	5244
	R	37/50->37/50	<4	<8	< 16	<d.l.< td=""><td>1097</td></d.l.<>	1097
	А	0-8	265	330	< 16	37616	220946
	2C1	8-12	153	72	< 16	4300	22400
DO	3C2	12-25	89	49	< 16	3401	18805
F 7	4C3	25-40	70	45	< 16	3398	13792
	5C4	40-59	92	45	< 16	2335	17122
	6C5	59->120	146	83	< 16	5927	35173

Table 2b

			t (°C)				pН			λ	(µS cn	n ⁻¹)		alkali	nity (n	ng L ⁻¹)		Orga	nic Carbon	$(mg L^{-1})$	
Site	sample	Dec	Jan	Febr	Apr	Dec	Jan	Febr	Apr	Dec	Jan	Febr	Apr	Dec	Jan	Febr	Apr	Dec	Jan	Febr	Apr
P1	1	14.5	11.2	14.5	-	9.21	8.26	7.15	-	-	1818	2060	-	18	-	57	-	11.4	< 0.04	3.3	-
P3	2	17.2	10.9	15.5	-	8.73	8.51	7.38	-	-	1240	2020	-	44	89	147	-	12.6	7.1	10.7	-
	4a	13.9	11.2	13	-	8.12	7.39	7.31	-	-	2630	2770	-	69	180	307	-	6.48	1.9	1.8	-
P9 long lysimeter	4b	13.7	11.3	12.2	-	7.91	7.41	7.11	-	-	3750	2840	-	47	158	314	-	6.93	2.3	2.1	-
10119 19 01110001	4c	13.7	10.8	11.3	-	7.72	7.03	7.26	-	-	3240	2740	-	111	167	307	-	3.25	1.6	1.6	-
Р9	5a	13.7	10.7	12.6	-	8.31	7.88	7.55	-	-	2900	2450	-	55	114	237	-	8.1	< 0.04	2.9	-
short lysimeter	5b	-	10.6	12.3	-	-	7.64	7.39	-	-	2600	2420	-	-	136	245	-	-	1.2	4.8	-
P9 rizhometer P9	19	-	-	-	24	-	-	-	7.93	-	-	-	2440	-	-	-	151	-	-	-	5.1
rizhometer P9	22	-	-	-	16.8	-	-	-	7.78	-	-	-	2180	-	-	-	152	-	-	-	5.62
rizhometer P9	25	-	-	-	20	-	-	-	7.62	-	-	-	2460	-	-	-	154	-	-	-	3.97
short lysimeter	25	-	-	-	18.7	-	-	-	7.5	-	-	-	2620	-	-	-	192	-	-	-	4.79
	6a	-	-	15.2	-	-	-	8.87	-	-	-	566	-	-	-	243	-	-	-	3.3	-
P5	6b	-	11.8*	14.3	-	-	9.14*	8.87	-	-	577*	706	-	-	178*	275	-	-	4.3*	4.1	-
	6c	-	11.8*	15.9	-	-	9.14*	8.8	-	-	577*	696	-	-	178*	252	-	-	4.3*	3.8	-
	7a	-	-	14.2	-	-	-	8.9	-	-	-	797	-	-	-	278	-	-	-	4.0	-
P6	7b	-	11.4	14.8	-	-	9.16	9.02	-	-	910	961	-	-	210	339	-	-	< 0.04	4.7	-
	7c	-	-	12.5	-	-	-	9.05	-	-	-	679	-	-	-	401	-	-	-	2.4	-

* = 6b + 6c average; - = not analysed

Table 3a

Site	sample		Cl (m	g L ⁻¹))		SO ⁴ (m	g L ⁻¹)		(Ca ²⁺ (r	ng L ⁻¹	^I)	Ν	1g ²⁺ (1	ng L ⁻¹)	l	Na ⁺ (n	ng L ⁻¹)		K ⁺ (r	ng L ⁻¹))
		Dec	Jan	Feb	Apr	Dec	Jan	Feb	Apr	Dec	Jan	Feb	Apr	Dec	Jan	Feb	Apr	Dec	Jan	Feb	Apr	Dec	Jan	Feb	Apr
P1	1	30	16	24	-	855	1480	1290	-	316	460	469	-	34	98	54	-	18	11	14	-	1	1	0.0	-
P3	2	14	43	10	-	783	1044	1082	-	314	393	468	-	39	62	45	-	7	22	9	-	2	6	0.9	-
DO	4a	443	368	272	-	1385	1186	962	-	433	322	279	-	156	132	105	-	326	268	205	-	26	21	17	-
long lysimeter	4b	448	437	279	-	1429	1404	1070	-	438	376	294	-	161	157	106	-	332	310	205	-	27	24	18	-
0,1	4c	439	357	270	-	1382	1184	978	-	422	332	275	-	148	122	100	-	319	262	198	-	24	21	18	-
P9	5a	398	337	255	-	1252	1187	920	-	338	295	259	-	135	133	103	-	285	255	194	-	21	21	16	-
short lysimeter	5b	-	346	258	-	-	1186	924	-	-	326	271	-	-	119	98	-	-	253	197	-	-	21	17	-
P9																									
rizhometer P9	19	-	-	-	274	-	-	-	757	-	-	-	219	-	-	-	75	-	-	-	168	-	-	-	16
rizhometer P9	22	-	-	-	260	-	-	-	893	-	-	-	262	-	-	-	83	-	-	-	184	-	-	-	16
rizhometer P9	25	-	-	-	302	-	-	-	753	-	-	-	286	-	-	-	78	-	-	-	178	-	-	-	17
short lysimeter	25	-	-	-	256	-	-	-	919	-	-	-	259	-	-	-	91	-	-	-	187	-	-	-	17
	6а	-	-	23		-	-	7		-	-	36		-	-	36	-	-	-	26	-	-	-	10	-
P5	6b	-	31*	30		-	6*	4		-	23*	23		-	41*	67	-	-	23*	34	-	-	9*	11	-
	6c	-	31*	18		-	6*	15		-	23*	42		-	41*	55	-	-	23*	26	-	-	9*	9.9	-
	7a	-	-	64		-	-	14		-	-	36		-	-	66	-	-	-	33	-	-	-	6.2	-
P6	7b	-	168	123		-	14	16		-	27	31		-	120	69	-	-	68	62	-	-	28	25	-
	7c	-	-	40		-	-	13		-	-	0		-	-	54	-	-	-	43	-	-	-	6.7	-

*= 6 b + 6c average; - = not analysed

Table 3b

Site	sample		Znμ	ug L ⁻¹			P	bμg L ⁻¹		Cd μ g L ⁻¹				
Site		Dec	Jan	Febr	Apr	Dec	Jan	Febr	Apr	Dec	Jan	Febr	Apr	
P1	1	20	126	172	-	22	17	18	-	-	< 0.4	2.3	-	
Р3	2	38	145	206	-	36	18	32	-	-	< 0.4	3.6	-	
DO	4a	127	6226	4324	-	21	30	<5	-	-	< 0.4	1.4	-	
long lysimeter	4b	117	6261	4613	-	12	28	<5	-	-	< 0.4	2.3	-	
0,7	4c	481	7861	5666	-	20	23	14	-	-	< 0.4	3.7	-	
Р9	5a	60	1497	1516	-	16	23	18	-	-	< 0.4	2.5	-	
short lysimeter	5b	-	2530	3195	-	-	19	23	-	-	< 0.4	7.1	-	
P9														
rizhometer P9	19	-	-	-	1665	-	-	-	64	-	-	-	5	
rizhometer P9	22	-	-	-	2056	-	-	-	10	-	-	-	7	
rizhometer P9	25	-	-	-	2274	-	-	-	18	-	-	-	10	
short lysimeter	25	-	-	-	1597	-	-	-	<5	-	-	-	3	
	6a	-	-	35	-	-	-	<5	-	-	-	<1.3	-	
P5	6b	-	31*	41	-	-	64*	18.6	-	-	<0.4*	<1.3	-	
	6с	-	31*	127	-	-	64*	<5	-	-	<0.4*	<1.3	-	
	7a	-	-	23	-	-	-	<5	-	-	-	<1.3	-	
Рб	7b	-	24	30	-	-	46	<5	-	-	< 0.4	<1.3	-	
	7c	-	-	18	-	-	-	13	-	-	-	<1.3	-	

*= b + c average; - = not analysed

Table 4a

Site	sample		Mn	ι μg L ⁻¹			Fe	μg L ⁻¹			Cu µ	ιg L⁻¹	
bite		Dec	Jan	Febr	Apr	Dec	Jan	Febr	Apr	Dec	Jan	Febr	Apr
P1	1	9	10	< 0.01	117	0.4	13	119	-	8.0	< 0.5	<1.2	-
P3	2	16	13	< 0.01	200	98	19	<5.3	-	26	4.8	2	-
D O	4a	140	226	141	74	47	11	7	-	4.9	5.9	11	-
long lysimeter	4b	68	230	62	73	< 0.1	13	<5.3	-	4.6	1.9	3	-
27	4c	63	92	5	70	< 0.1	6.0	<5.3	-	2.2	1.7	2	-
P9	5a	15	59	3	77	15	2.2	<5.3	-	6.0	4.6	8.9	-
short lysimeter	5b	-	22	< 0.01	-	-	15	<5.3	-	-	0.7	4	-
P9 rizhometer P9	19	-	-	-	<0.01	-	-	-	-	-	-	-	<1.2
rizhometer P9	22	-	-	-	< 0.01	-	-	-	-	-	-	-	<1.2
rizhometer P9	25	-	-	-	< 0.01	-	-	-	-	-	-	-	4
short lysimeter	25	-	-	-	< 0.01	-	-	-	-	-	-	-	2
	6a	-	-	-	-	-	-	-	-	-	-	-	-
P5	6b	-	9*	-	-	-	23*	<5.3	-	-	7*	4.3	-
	6c	-	9*	-	-	-	23*	<5.3	-	-	7*	<1.2	-
	7a	-	-	-	-	-	-	<5.3	-	-	-	<1.2	-
P6	7b	-	9	-	-	-	24	<5.3	-	-	8	<1.2	-
	7c	-	-	-	-	-	-	<5.3	-	-	-	<1.2	-

*= b + c average; - = not analysed

Table 4b

Site	sampler	Species* (PHREEQC)	fraction bound to colloidal FA (WHAM)
	1 - Dec	Ca ²⁺ , CaSO ₄ , CuOH ⁺ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , MnSO ₄ , PbOH ⁺ , PbCO ₃ , Pb ²⁺ , Zn ²⁺ , ZnSO ₄	Cu (80%), Zn (3.5%), Pb (0.2%)
P1	1 - Jan	Ca ²⁺ , CaSO ₄ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , MnSO ₄ , PbOH ⁺ , PbCO ₃ , Pb ²⁺ , Zn ²⁺ , ZnSO ₄	no O.C.
	1 - Febr	Ca ²⁺ , CaSO ₄ , Mg ²⁺ , MgSO ₄ , Pb ²⁺ , PbCO ₃ , Zn ²⁺ , ZnSO ₄	Zn (2.4%), Cd (1.15%), Pb (0.015%)
	2 - Dec	Ca ²⁺ , CaSO ₄ , CuCO ₃ , Mn ²⁺ , MnSO ₄ , PbCO ₃ , Zn ²⁺ , ZnSO ₄	Cu (60%), Zn (5%), Pb (0.1%)
P3	2 - Jan	$Ca^{2+}, CaSO_4, CuOH^+, CuCO_3, Mg^{2+}, MgSO_4, Mn^{2+}, MnSO_4, MnCO_3, PbCO_3, PbOH^+, Zn^{2+}, ZnSO_4, CuOH^+, CuCO_3, Mg^{2+}, MgSO_4, Mn^{2+}, MnSO_4, MnCO_3, PbOH^+, Zn^{2+}, ZnSO_4, Mn^{2+}, MnSO_4, Mn^{2+}, MnSO_4, Mn^{2+}, MnSO_4, Mn^{2+}, MnSO_4, Mn^{2+}, MnSO_4, Mn^{2+}, MnSO_4, Mn^{2+}, Mn^{2$	Cu (55%), Zn (3.2%), Pb (0.06%)
	2 - Febr	Ca^{2+} , $CaSO_4$, $CuCO_3$, Mg^{2+} , $MgSO_4$, $PbCO_3$, Zn^{2+} , $ZnSO_4$	Cu (99%), Zn (8.8%), Cd (3.6%), Pb (70%)
	6b+c - Jan	Ca ²⁺ , CaSO ₄ , CuCO ₃ , Cu(CO ₃) ₂ ²⁻ , Mg ²⁺ , MgSO ₄ , MnCO ₃ , Mn ²⁺ , PbCO ₃ , Pb(CO ₃) ₂ ²⁻ PbOH ⁺ ,Zn(OH) ₂ , Zn ²⁺ , ZnCO ₃	Cu (39%), Zn (3.5%)
P5	6a - Febr	Ca ²⁺ , CaCO ₃ , Mg ²⁺ , MgCO ₃ , Zn(OH) ₂ , Zn2+, ZnCO ₃	Zn (4.8%)
	6b - Febr	Ca ²⁺ , CaCO ₃ , CuCO ₃ , Cu(CO ₃) ₂ ²⁻ , Mg ²⁺ , MgCO ₃ , PbCO ₃ , Pb(CO ₃) ₂ ²⁻ , Zn ²⁺ , ZnCO ₃	Cu (60%), Zn (3%), Pb (0.9%)
	6c - Febr	Ca ²⁺ , CaCO ₃ , Mg ²⁺ , MgCO ₃ , Zn(OH) ₂ , Zn ²⁺ , ZnCO ₃	Zn (3%)
	7b - Jan	Ca ²⁺ , CuCO ₃ , Cu(CO ₃) ₂ ²⁻ , Mg ²⁺ , MgCO ₃ , MnCO ₃ , Mn ²⁺ , PbCO ₃ , Pb(CO ₃) ₂ ²⁻ , PbOH ⁺ , Zn(OH) ₂ , Zn ²⁺ , ZnCO ₃	no O.C.
P6	7a Febr	Ca ²⁺ , CaCO ₃ , Mg ²⁺ , MgCO ₃ , Zn(OH) ₂ , Zn ²⁺ , ZnCO ₃	Zn (5%)
	7b - Febr	Ca ²⁺ , CaCO ₃ , Mg ²⁺ , MgCO ₃ , Zn(OH) ₂ , Zn ²⁺ , ZnCO ₃	Zn (5.5%)
	7c - Febr	Ca ²⁺ , CaCO ₃ , Mg ²⁺ , MgCO ₃ , PbCO ₃ , Pb(CO ₃) ₂ ²⁻ , Zn(OH) ₂ , Zn ²⁺ , ZnCO ₃	Zn (5%), Pb (1%)
Р9	4a - Dec	Ca ²⁺ , CaSO ₄ , CuCO ₃ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , MnSO ₄ , PbCO ₃ , Zn ²⁺ , ZnSO ₄	Cu (20%), Zn (1%)
long lysimeter	4a - Jan	Ca ²⁺ , CaSO ₄ , CuCO ₃ , Cu ²⁺ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , MnSO ₄ , PbCO ₃ , Zn ²⁺ , ZnSO ₄	Cu (4.4%), Zn (0.05%)
5	4a - Febr	Ca ²⁺ , CaSO ₄ , CuCO ₃ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , Zn ²⁺ , ZnSO ₄	Cu (8.1%), Zn (0.4%), Cd (0.3%)

	4b - Dec	Ca ²⁺ , CaSO ₄ , CuCO ₃ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , MnSO ₄ , PbCO ₃ , Pb ²⁺ , Zn ²⁺ , ZnSO ₄	Cu (12%), Zn (0.4%), Cd (0.3%)
	4b - Jan	Ca ²⁺ , CaSO ₄ , CuCO ₃ , Cu ²⁺ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , MnSO ₄ , PbCO ₃ , Pb ²⁺ , Zn ²⁺ , ZnSO ₄	Cu (5.8%), Zn (0.06%)
	4b - Febr	Ca^{2+} , $CaSO_4$, $CuCO_3$, Mg^{2+} , $MgSO_4$, Mn^{2+} , $MnSO_4$, Zn^{2+} , $ZnSO_4$	Cu (12%), Zn (0.4%), Cd (0.3%)
	4c - Dec	Ca^{2+} , $CaSO_4$, $CuCO_3$, Mg^{2+} , $MgSO_4$, Mn^{2+} , $MnSO_4$, $PbCO_3$, Pb^{2+} , Zn^{2+} , $ZnSO_4$	Cu (13%), Zn (0.6%)
	4c - Jan	Ca^{2+} , $CaSO_4$, $CuCO_3$, Mg^{2+} , $MgSO_4$, Mn^{2+} , $MnSO_4$, $PbCO_3$, Pb^{2+} , Zn^{2+} , $ZnSO_4$	Cu (2.7%), Zn (0.2%)
	4c - Febr	Ca ²⁺ , CaSO ₄ , CdCl ⁺ , Cd ²⁺ , CuCO ₃ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , MnSO ₄ , PbCO ₃ , Zn ²⁺ , ZnSO ₄	Cu (9.7%), Zn (0.4%), Cd (0.2%), Pb (1.2%)
	5a+5b - Dec	Ca ²⁺ , CaSO ₄ , CuOH ⁺ , CuCO ₃ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , MnSO ₄ , PbCO ₃ , Zn ²⁺ , ZnSO ₄	Cu (66%), Zn (4%), Pb (0.05%)
	5a - Jan	Ca ²⁺ , CaSO ₄ , CuCO ₃ , CuOH ⁺ , Cu ²⁺ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , MnSO ₄ , PbCO ₃ , Pb ²⁺ , Zn ²⁺ , ZnSO ₄	no O.C.
	5a - Febr	Ca ²⁺ , CaSO ₄ , CdCl ⁺ , Cd ²⁺ , CuCO ₃ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , MnSO ₄ , PbCO ₃ , Zn ²⁺ , ZnSO ₄	Cu (22%) Zn (1%), Cd (0.7%), Pb (2%)
	5b - Jan	Ca ²⁺ , CaSO ₄ , CuCO ₃ , CuOH ⁺ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , MnSO ₄ , PbCO ₃ , Pb ²⁺ , Zn ²⁺ , ZnSO ₄	Cu (0.06%), Zn (0.3%)
P9	5b - Febr	Ca^{2+} , $CaSO_4$, $CdCl^+$, Cd^{2+} , $CuCO_3$, Mg^{2+} , $MgSO_4$, $PbCO_3$, Zn^{2+} , $ZnSO_4$	Cu (22%), Zn (1%), Cd (0.7%), Pb (2%)
	ryzhom 19 Apr	Ca^{2+} , $CaSO_4$, $CdCl^+$, Cd^{2+} , Mg^{2+} , $MgSO_4$, $PbCO_3$, $PbOH^+$, Pb^{2+} , Zn^{2+} , $ZnSO_4$	Zn (2.8%), Cd (1.3%), Pb (5.6%)
	ryzhom 22 Apr	Ca ²⁺ , CaSO ₄ , CdCl ⁺ , Cd ²⁺ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , MnSO ₄ , PbCO ₃ , PbOH ⁺ , Pb ²⁺ , Zn ²⁺ , ZnSO ₄	Zn (2.3%), Cd (1.3%), Pb (4%)
	ryzhom 25 Apr	Ca ²⁺ , CaSO ₄ , CdCl ⁺ , Cd ²⁺ , CuCO ₃ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , MnSO ₄ , PbCO ₃ , PbOH ⁺ , Pb ²⁺ , Zn ²⁺ , ZnSO ₄	Zn (2.3%), Cd (0.6%), Pb (37%)
	lysim 25 Apr	Ca ²⁺ , CaSO ₄ , CdCl ⁺ , Cd ²⁺ , CuCO ₃ , Mg ²⁺ , MgSO ₄ , Mn ²⁺ , MnSO ₄ , Zn ²⁺ , ZnSO ₄	Cu (94%), Zn (3%), Cd (0.9%)

*listed for each element in order of relative abundance; only species present in amounts >5% are reported

Table 5.







Fig 2



Fig. 3







 \triangle P9 Dec \bullet P9 Jan \Box P9 Feb \diamond P9 Apr

Fig. 5