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Full scale amendment of a contaminated wood impregnation site with iron water treatment residues 1 2 Sanne Skov Nielsen^{a)}, Peter Kjeldsen(^(C)), Rasmus Jakobsen^{b)} 3 4 Department of Environmental Engineering, Technical University of Denmark, Miljovej 113, DK-2800 Kgs. 5 Lyngby, Denmark 6 e-mail: pekj@env.dtu.dk 7 8 (\boxtimes) Corresponding author: 9 Peter Kjeldsen 10 Tel. +45 45251561 11 Fax. +45 45932850 12 e-mail: pekj@env.dtu.dk 13 14 a) Present address: Orbicon, Ringstedvej 20, DK-4000 Roskilde, Denmark (e-mail: sann@orbicon.dk) 15 b) Present address: Geological Survey of Denmark and Greenland, Geological Survey of Denmark and 16 Greenland, Øster Voldgade 10, DK-1350 Copenhagen K, Denmark (e-mail: raj@geus.dk) 17 18 Abstract 19 Iron water treatment residues (Fe-WTR) are a free by-product of the treatment of drinking water with high 20 concentration of iron oxides and potential for arsenic sorption. This paper aims at applying Fe-WTR to a 21 contaminated site, measuring the reduction in contaminant leaching, and discussing the design of delivery 22 and mixing strategy for soil stabilization at field scale and present a cost-effective method of soil mixing by 23 common contractor machinery. Soil contaminated by As, Cr, and Cu at an abandoned wood impregnation site was amended with 0.22% (dw) Fe-WTR. To evaluate the full scale amendment a 100 m² test site and a 24 25 control site (without amendment) were monitored for 14 months. Also soil analysis of Fe to evaluate the 26 degree of soil and Fe-WTR mixing was done. Stabilization with Fe-WTR had a significant effect on 27 leachable contaminants, reducing pore water As by 93%, Cu by 91% and Cr by 95% in the upper samplers. 28 Dosage and mixing of Fe-WTR in the soil proved to be difficult in the deeper part of the field, and pore 29 water concentrations of arsenic was generally higher. Despite water logged conditions no increase in 30 dissolved iron or arsenic was observed in the amended soil. Our field scale amendment of contaminated 31 soil was overall successful in decreasing leaching of As, Cr and Cu. With minor improvements in the 32 mixing and delivery strategy, this stabilization method is suggested for use in cases, where leaching of Cu, 33 Cr and As constitutes a risk for groundwater and freshwater. 34 Keywords Field experiment, Iron oxide, Metal, Stabilization, Wood preservation sites 35 Running title: Amendment of a wood impregnation site with iron water treatment residues 36

37 1 Introduction

38 Throughout Europe brownfields are present as remnants of industrial production. As early industrial sites 39 from a time with no environmental protection laws, the sites may be heavily contaminated, and often the 40 contamination consists of heavy metals, that are, by nature, not degradable in the environment. The sheer 41 volume of soil is frequently too large for excavation and when the sites are situated in isolated locations, 42 development is not economically feasible. These sites may pose a threat to aquifers and surface waters by 43 leaching of contaminants. Cleaning and reutilization technologies of metal(loid) polluted soils including 44 soil washing [1, electrochemical remediation [2], phytoremediation [3] and combinations thereof are only 45 scarcely used [4].

Stabilizing or amending metal(loid) contaminated soil involves adding an amendment to the contaminated soil layers that reduces leaching of contaminants from the soil by sorption or precipitation, and is useful for brownfields and cases where there are no special requirements for the land use. In cases where such circumstances can be fulfilled, the method is considered a cost effective way to minimize the impact of mobile element contamination on the recipients [5]. Bioavailability of contaminants at the site may also be limited [5,6] which prevents spreading of contaminants in the food chain.

52 At former wood impregnation sites especially arsenic (As) is often very mobile in soil and 53 groundwater systems [7,8]. Iron oxides have a strong affinity for arsenic and other trace elements [9,10] 54 and several iron oxides has been evaluated as soil amendment for reducing arsenic and heavy metal 55 leachability: Ochre from mining activities [11], goethite [12], and water treatment residues [13,14]. In cases of multi element contamination a mix of amendments may prove the best option and can be assessed in a 56 57 strategic selection framework [15]. However, the long-term ability of the amendment to reduce leachability 58 should also be considered since slow transformation of the reactive minerals to less reactive minerals may 59 occur over longer time [16,17].

60 Iron water treatment residues (Fe-WTR) are a common waste product in Denmark, where the 61 water supply is based entirely on groundwater. With the main constituent (60-70%) being ferrihydrite, an 62 iron hydroxide, it is proposed here as a cheap and efficient soil amendment. No pH buffers are required as 63 for iron(II)sulphate, and the cost is limited to a control element analysis and transport to the site. Numerous 64 laboratory studies with soil amendments using iron oxides have been carried out, but few have tested the 65 methods at field scale [14,18] and applicability has not been discussed and evaluated. Laboratory 66 experiments are often carried out under optimal conditions, and their results may not directly be transferred 67 to full-scale scenarios in the field. Also impacts on the soil amendment such as geochemical processes in 68 the soil influenced by soil moisture and biota are lacking in laboratory experiments. For iron oxide 69 amendments prone to microbial iron reduction it is necessary to test their persistency under natural 70 conditions.

For full scale amendments large areas need to be treated so choosing a cost-effective and simple way to mix the treated soil and the amendment is crucial. In considering the options for soil mixing, the depth of the contaminated soil layers to be treated is an important factor. Soil mixing by augers is routinely 74 used in the delivery of zero valent iron (ZVI)-clay mixtures for degradation of chlorinated solvents [19] at 75 depths down to 8 m below ground. However, the equipment is highly specialized and therefore expensive 76 to rent. Trench ploughing, with a common use in forestry and a working depth of about 0.9 m, is an option, 77 but the trench plough has a considerable turning radius and will only reach the maximum working depth 78 after 50 m of ploughing. Another option is to use a screening bucket, a tool used in waste management and 79 soil treatment at construction sites. A screening bucket consists of a screen, capable to hold about 0.5 m^3 of 80 soil that is passed through the screen by rotating it. Soil fractions >50 mm (usually stones, but for fill also 81 bricks and concrete pieces) is then retained inside the screen and can be discarded. Adding Fe-WTR as 82 slurry, soil and slurry mixture would pass faster through the screening bucket.

This study evaluates the full scale amendment with Fe-WTR at a brownfield contaminated with high levels of As, chromium (Cr) and copper (Cu). To our knowledge this is a novel technology, not tested at full scale before. In designing the full scale approach, emphasis was put on making a low cost method easily applicable to a brownfield, thereby maximizing the options for wide-spread professional use. For evaluation of the retention capacity of the amendment, soil pore water samplers were installed at the site and trace element composition monitored for 12 months.

89

90 2 Materials and methods

91 2.1 Field site

92 The Collstrop wood impregnation site close to Hillerød, Denmark, has previously been used for soil 93 stabilization with WTR at a small scale [14]. Wood impregnation using As, Cr, and Cu was initiated in 94 1955. Heavily contaminated with As, Cr, and Cu, the site is left as a brownfield in the forest. A sketch of 95 the site with vegetation cover is presented in Fig 1. A profound correlation between vegetation cover and 96 contaminant concentration was found in an earlier study [20] with only very limited vegetation growing at 97 contamination hotspots. For this experiment a bare dripping pad in the eastern part of the site was chosen as 98 test site. At dripping pads the treated wood items were placed for drying. Fig. 1 shows the configuration of 99 the test site and location of the 10 pore water samplers, more than 40 soil samples and 4 shallow wells.

100

101 2.2 Amendment

Fe-WTRs were collected at Sjælsø Waterworks in Northern Zealand, Denmark, which supplies water to large parts of the greater Copenhagen area. Groundwater is extracted from wells 50 to 120 m below surface in a limestone aquifer, and the water is then aerated and passed through a sand filter for separating iron and manganese oxides from the drinking water. Freshly precipitated Fe-WTR slurry with a natural water content (91.2%, n=5) was used as amendment.

- 107
- 108 Chemical analysis and BET (Brunauer–Emmett–Teller) surface area of the solids can be found in Table 1.
- 109 For further information on the Fe-WTR composition, we refer to the earlier publication [14].
- 110

111 **2.3** Soil mixing procedure at field scale

112 Before considering the options for soil mixing, an estimate was made of the volume of soil that needed to 113 be treated at the site. The contaminants originate from dripping of impregnation liquids and as most of the 114 contaminant masses are present in the upper 1 m of the soil profile [14], an estimated 80% of the 115 contaminant mass would be captured by treating the upper 1 m of the soil. Additionally, previous studies at 116 the field site had revealed a hanging water table 1-1.5 m below ground [14]. Adding Fe-WTR below the 117 water table will have little effect, as the highly reactive ferrihydrite is expected to be easily dissolved by 118 reductive dissolution and presumably destroy the effect of the amendment, as the iron oxides are dissolved, 119 releasing all captured contaminants. Thus a treatment depth of 1 meter was chosen. A rotary screening 120 bucket mounted on an excavator was used to mix soil and Fe-WTR slurry. A test field of 10 by 10 m with 121 amendment of Fe-WTR was established. Adjacent to that a 10 by 10 m test site without amendment was 122 established, to act as a control plot for the evaluation of the Fe-WTR amendment efficiency. Pore water 123 was also sampled from an undisturbed soil profile within a few meters from the test fields. For configuration of the test fields see Fig. 1. Fig. 2 shows the rotary screening bucket in action. 124

125 A slurry-trailer carrying a 4.5 m³ basin, installed with mixers, kept the WTR slurry in suspension to make sure the solids were applied evenly at a rate of 0.12 m³·min⁻¹. Soil and Fe-WTR was mixed in 2 126 127 steps; excavating to 0.5 m and mixing with half the Fe-WTR slurry, then backfilling and mixing again down to 1 meter depth and adding the other half of the Fe-WTR slurry. A total of 4 m³ was added to the 128 129 amended field, which was the maximum amount possible to not increase the volume of the stabilized soil 130 significantly. The average added dry Fe-WTR percentage to the dry matter of soil in the 10 m by 10 m by 1 131 m is estimated to 0.22 %(dw). The control field was also mixed in 2 steps, but with no Fe-WTR addition 132 and therefore more difficult to mix because the slurry made it easier for the soil to pass through the 133 screening bucket.

During mixing greenish colored concrete pieces were separated from the soil, which are likely remains of the impregnation basins and their greenish color presumably copper-arsenate precipitates. For brownfield management, removing waste fractions like this will improve the esthetic value of the site.

137

138 2.4 Sampling and monitoring

139 After soil mixing in late April 2011, the site was left to settle. Soil samples were taken at random spots in 140 the amended and the control plots in May 2011. In total nine composite samples representing the depth 0-141 0.5 meters depth and 0.5-1 meter depth were sampled for both plots (in total 36 samples). The average 142 composition of the soil in the mixed plot and in the control plot was compared using t-test (P=0.05, n=18). 143 Groundwater wells and soil pore water samplers were installed in early May 2011. As according to the 144 manufacturer of samplers the first 2 L of samples has to be discarded; only samples after June 2011 were 145 analyzed. The experiment was concluded in July 2012 after collecting and analyzing more than 200 water 146 samples.

147 To monitor the composition of soil pore water, 10 PrenArt® pore water samplers were installed at 148 an angle of 45° (to limit rapid flow along the tubing) at depth 0.5 and 1 m below surface. Each sampler was 149 connected to a 1 L Bluecap bottle with 1 kPa vacuum. The bottle was changed every 21-31 d, when the 150 collected amount was sufficient for analysis. Total sample volume and pH were measured and 20 mL 151 samples were filtered through a 0.45 µm hydrophilic PTFE filter and stored at 5°C until analysis. The soil 152 pore water concentrations in the amended and the unamended field plot and in an undisturbed reference site 153 a few meters away were sampled over a 14 month period to monitor potential temporal changes. Both test 154 fields were divided in a 3x3 sampling grid (Fig. 1). Soil samples were collected using a hand auger. A 100 g composite sample was taken from each of the 0.5 m homogenized cores at depths 0-0.5 m and 0.5-1.0 m. 155 The samples were dried at 105°C for 24 h and homogenized in an agate mortar. From the homogenized 156 157 sample 2 g was taken for acid digestion in 7 N HNO₃.

158 A hanging secondary water table was present at the site due to clay layers in the glaciofluvial 159 formation [14]. At the time of soil mixing in late spring it was observed about 1 meter below the soil surface. To measure the water table 2 shallow wells (PE tubing Ø2.2 cm) were installed by hand augering 160 161 at 1 m depth with 20 cm screen. Water table measurements were done with the same frequency as water 162 sampling.

163

164 2.5

Element analysis of soil and water

165 Element content in soil and water samples was determined using a Varian 5000 ICP-OES. A standard reference soil was included for every 30 samples and all measurements of those found to be within 10%. 166 Detection limits for water samples were 0.7 μ g·L⁻¹ and for soil samples 35 mg·kg⁻¹ at wavelengths 259.940 167 (Fe) 293.931 (Mn) 327.395 (Cu) 205.560 (Cr) and 188.980 (As) nm. 168

169 For the last 3 rounds of pore water sampling the speciation of Cr(VI)/Cr(III) and As(V)/As(III) 170 was measured. Visual light spectrophotometric screening for Cr(VI) was done using s-diphenylcarbazide as an indicator (EN 196-10:2006, 2006) with at detection limit of 10 μ g·L⁻¹. As(V)/As(III) speciation was 171 done by filtering through an aluminosilicate-containing cartridge, retaining As(V) [21]. The As(V) 172 173 concentration was then calculated as the difference between As(III) and the total As content.

174

175 2.6 Physical properties of soil and WTR

176 BET surface area was measured by the N_2 adsorption method [22]. For soil and WTR properties such as 177 geological characterization, grain size and distribution see [14].

178

179 3 **Results and discussion**

180 3.1 Distribution of the amendment

181 The analysis of the homogenized core samples showed only a slight difference in iron content with an average of 4815 mgFe·kg⁻¹ (range 1193-6124 mgFe·kg⁻¹) in the control field and 5360 mgFe·kg⁻¹ (range 182 3805-13304 mgFe·kg⁻¹) in the amended (Table 1). A difference of 0.13 wt % Fe was found in the upper 0.5 183

m of soil between the treated and the untreated field and there was no significant (*t*-test, P=0.05, n=18) difference between iron concentrations in the lower parts of the two fields. The iron amendment was not sufficiently distributed, as the upper 0.5 m in the treated field has a mean value of 6140 mgFe·kg⁻¹ and the lower only slightly more than the natural background value of 4815 mgFe·kg⁻¹. Iron distribution in soil layers is shown in Fig. 3, which also reveals a high variability in the background level of iron content with high iron content at the bottom of the control field, may be a soil horizon formed by the natural pseudogleyic conditions in the soil.

191

192 **3.2** Leaching of contaminants in the amended soil

193 Soil pore water concentrations are a tool to evaluate the most mobile fraction of trace elements in soils [23]. 194 Pore water concentrations of As (Fig. 4b+c), Cr (Fig. 5a+b) and Cu (Fig. 5c+d) are decreased as the 195 addition of Fe-WTR significantly reduces the mobile fraction of contaminants in the soil. Comparing all 196 data points for the full year of sampling, the soil amendment reduces pore water concentrations in the mixed fields with 93% for As, 91% for Cu and 95% for Cr in the shallow (50 cm below ground) samplers 197 198 and 75% for As, 89% for Cu and 97% for Cr in the deeper samplers when comparing with the control site. 199 This difference shows that the upper soil has obtained a larger retention for As and that the lower part of 200 the treated field did not receive a sufficient amount of the Fe-WTR. In terms of the Danish groundwater quality criteria, the dose of Fe-WTR was not sufficient. The average As concentration for amended soil 201 (upper samplers) is 328 μ g·L⁻¹ and does not comply with the groundwater criteria of 8 μ g·L⁻¹[24], but the 202 leaching of contaminants has been significantly reduced. Studies using iron oxide amendment for 203 204 stabilization has typically used a higher percentage of iron oxides added to the soil to obtain better 205 contaminant retention: 5-15 wt% [25] or 1-8% [18] oxygen scarfing granulate (69% magnetite) or 2.5-5.0 206 wt% ochre (likely ferrihydrite, calcite and aragonite) [11]. Average Fe-WTR addition obtained in this full 207 scale study was about 0.22 wt % of the amended soil. The amendment is to be increased to a range of 1-2 208 wt% in future full scale projects.

209

210 3.3 Speciation of As and Cr

211 As toxicity and mobility in soil and groundwater is hugely dependent on speciation, a chemical speciation 212 As and Cr was done in the last 3 rounds of pore water sampling as mentioned in the material and methods 213 section. As no Cr(VI) was found in the pore water, all chromium at the site is assumed to be Cr(III) which 214 is consistent with the low solubility of chromium hydroxides in soil [26] and the observed low pore water 215 concentrations. Only two samples contained more than 5% As(III), which means that the overall speciation 216 of arsenic at the site is consistent with aerobic conditions. These two samples with 8.5% and 20% As(III) 217 were from the deep sampler in the untreated field. This correlates well with the fact that the deep samplers 218 at 90 cm below ground are likely to be below the water table and that this field had the high iron 219 concentrations in the pore water, suggesting somewhat reducing conditions.

221 **3.4 Effect of water logged conditions**

Reductive dissolution of iron oxides can happen as the soil gets saturated with water and release of sorbed arsenic takes place accordingly [27-29]. The rate of reductive dissolution of iron (hydr-)oxides depends on mineral type, crystallinity and aggregate size [30,31]. For Fe-WTR amendment the reaction rate is expected to be very high as the main constituent is ferrihydrite, the most reactive iron hydroxide for the above mentioned reasons.

227 Water table measurements are shown in the top right corner of Fig. 4a, which also shows the total 228 iron concentration of the sampled pore water in Fig 3 d+e. Only for the unamended field did iron concentrations in the pore water increase (up to around 400 $\mu g \cdot L^{-1}$) in the period from late August to 229 December. Small amounts of iron (up to 100 μ g·L⁻¹) were found after December, despite the water table 230 231 only slowly decreased during the months of January to March. Temperature dependence of iron reduction 232 rates in soil with the release of Fe and As have been reported to strongly slow down when the temperature 233 decreases from 23°C and 14°C to 5°C [29]. If this is part of the explanation it implies that a coincidence of 234 the cold season with the infiltration season, as seen here, is beneficial where waterlogging may occur.

235 As no increase in dissolved iron is seen in the iron amended soil, not even during water logging, it 236 suggests that the iron amendment is resistant to iron reducing conditions. Microbial activity in the soil may 237 be low due to the high contents of contaminants [32]. Especially copper has been shown to have a toxic 238 effect on iron reducing bacteria, thereby halting iron reduction [33]. However as the copper concentration is 239 almost identical in the fields, this cannot be the sole explanation. If the reduction of iron oxide in the soil is 240 not complete due to an excess of iron oxide in the soil (which is likely to be the case for the Fe-WTR amended soil), it is possible that iron and arsenic is retained in the soil. Fe²⁺ competes for arsenate on the 241 ferrihydrite surface [34], so the Fe^{2+} released by iron reduction may not be released to the water phase in 242 243 the amended soil. Likewise the release of As to the aqueous phase does not happen until the number of 244 surface sites on the ferrihydrite surface is too small to adsorb all arsenic [28].

Another approach is to study the manganese concentrations in the soil pore water (Fig. 3f+g). Fe-WTR contains a substantial amount of manganese oxides, precipitated with the iron oxides, and the background concentration of manganese (in average 198 mg·kg⁻¹) is relatively smaller compared to the amount of WTR added to the soil. Manganese reduction is thermodynamically favored over iron reduction, so a high porewater concentration of manganese indicates WTR dissolution, as can be seen in Fig. 3, where a peak in the Mn concentration is correlated with a high water table (Fig. 4a).

251

252 **3.5** Mechanical treatment of soil

Based on the As and Fe (Fig. 4bcde) as well as the Cr and Cu (Fig. 5abcd) pore water concentrations, the mechanical treatment of mixing the soil without adding Fe-WTR has a significant effect on the levels of all measured elements in the soil pore water. This indicates that the mixing of contaminated soil increases the leaching of elements, possibly due to release of fine colloids mobilized with the mechanical treatment of the soil. Addition of iron oxides to a soil do change the mechanical properties of soil and ferrihydrite, the 258 main constituent of Fe-WTR, has even been proposed as a clearing agent to reduce turbidity in fresh water 259 [35]. A cementing effect of Fe-WTR may limit the colloidal transport of elements and this effect can 260 account for some of the difference in metal(loid) leaching in the two mechanically treated fields. This 261 observation suggest that the mechanical treatment of soil, for instance in the case of excavation and 262 landfilling, may increase the leaching of contaminants, but this effect can be avoided with the addition of 263 Fe-WTR.

264

265 4 Conclusions

266 Stabilization with Fe-WTR in full scale proved to be fairly simple to apply to the contaminated site. 267 Despite insufficient application of amendment, the field experiment provides valuable insight in the 268 practical application of an iron oxide based soil amendment. Leaching of As, Cr, Cu, as measured by pore 269 water concentrations, was significantly reduced in the Fe-WTR amended soil compared to both a 270 mechanically mixed, but not amended, soil and an undisturbed soil. Even with an iron addition of only 0.22 271 wt%, the amendment caused a significant stabilization, but more Fe-WTR than applied in this study is 272 needed to comply with ground water quality standards. Despite water logged conditions in the lower parts 273 of the soil, no clear signs of iron reduction and subsequent contaminant release were observed. Manganese 274 was however released from the treated soil, suggesting that the Fe-WTR is subject to partial reductive 275 dissolution.

276 The distribution of soil amendment achieved with a screening bucket was satisfying for the upper part of 277 the soil, but it proved more difficult, at least in this very first attempt, to amend the soil from 0.5 to 1 m 278 below surface. In any case, mechanical treatment or mixing of metal(loid) contaminated soil should be 279 avoided as it mobilizes dissolved (or colloidally transported) contaminants into the soil pore water. 280 However Fe-WTR treatment prevents this, possibly because of inter-particulate cementation processes. As 281 the dose of WTR and difficulties with proper mixing are rather easy obstacles to overcome, we propose this 282 method for use in situ at other contaminated sites.

283

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Tables

Table 1 Average solid concentrations and BET surface area of soil and Fe-WTR and average speciation of

380 pore water in the amended field, control field and undisturbed soil. Number of samples, *n*, are given in

381 parenthesis). For pore water numbers in parenthesis are lowest and highest measured value of As(III) in

 $\mu g \cdot L^{-1}$. Basic soil properties are given in [Error! Bookmark not defined.].

		As	Cr	Cu	Mn	Fe	BET	As(III)/As(V)	Cr(III)/Cr(VI)
		[mg·kg ⁻¹]				[mg·kg ⁻¹]	$[m^2 \cdot g^{-1}]$	[-]	
Amended field	0-0.5 m (<i>n</i> =9)	563	205	598	325	6140	1.654	2.3 (0.1-3.7) (<i>n</i> =7)	>99.9 (<i>n</i> =7)
	0.5-1 m (<i>n</i> =9)	430	133	423	227	4580	1.632		
Control field	0-0.5 m (<i>n</i> =9)	539	156	378	198	4830	1.859	11.1 (4.2-20) (<i>n=3</i>)	>99.9 (n=3)
	0.5-1 m (<i>n</i> =9)	572	149	386	189	4800	1.911		
Undisturbed field		-	-	-	-	-	-	1.2 (0.4-3.5) (<i>n</i> =4)	>99.9 (<i>n</i> =4)
Fe-WTR	(n=5)	46.1	13.0*	36.6	8489.8	28.6%	160.3	-	-

*below analytical quantification limit - not measured

Figure captions

Fig. 1 Sketch of vegetation Collstrop Brownfield, a former wood impregnation plant. Vegetation is drawn based on observations from [Error! Bookmark not defined.], but is generally in agreement with author observation within recent years (left). Configuration of the full scale soil stabilization experiment with location of pore water samplers, shallow wells and soil samples. Two soil samples were taken at each sampling point (top right). Fig. 2 Pictures from the amendment action showing the used rotary screening bucket. Fig. 3 Total iron concentration in the two test fields after application of water treatment residues (WTR) amendment to the northern field. The size of the bubble denotes the iron concentration Fig. 4 Soil pore water concentrations of redox sensitive elements As(b+c), Fe(c+d) and Mn(e+f) and average water table (a). Left column shows pore water samplers in 50 cm depth and the right 100 cm depth. Data points are average of two samplers for elements and four for the water table. Error bars denotes the high and low value. If no error bars are shown, the data point consists of a single measurement from one sampler Fig. 5 Soil pore water concentrations of Cr (a+b) and Cu (c+d). Data points are average of two samplers. Error bars denotes the high and low value. If no error bars are shown, the data point consists of a single

measurement from one sampler









Fig. 2b







