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CONTAMINATION RELATED TO ANTHROPIC ACTIVITIES. CHARACTERIZATION AND REMEDIATION

#### Enrichment of trace elements in the clay size fraction 4 of mining soils 5

Patrícia Gomes · Teresa Valente · M. Amália Segueira Braga · J. A. Grande · M. L. de la Torre

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13Abstract Reactive waste dumps with sulfide minerals promote acid mine drainage (AMD), which results in water and 14soil contamination by metals and metalloids. In these systems, 1516contamination is regulated by many factors, such as mineralogical composition of soil and the presence of sorption sites 17on specific mineral phases. So, the present study dedicates 18 19itself to understanding the distribution of trace elements in different size fractions (<2-mm and <2-µm fractions) of min-20ing soils and to evaluate the relationship between chemical 2122and mineralogical composition. Cerdeirinha and Penedono, 23located in Portugal, were the waste dumps under study. The results revealed that the two waste dumps have high degree of 2425contamination by metals and arsenic and that these elements are concentrated in the clay size fraction. Hence, the higher 26degree of contamination by toxic elements, especially arsenic 27in Penedono as well as the role of clay minerals, jarosite, and 28goethite in retaining trace elements has management implica-2930 tions. Such information must be carefully thought in the reha-31bilitation projects to be planned for both waste dumps.

32	Keywords Mining soils · Contamination · Clay size fraction ·
33	Accumulation factor · Trace elements · Portugal

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P. Gomes · T. Valente (🖂) · M. A. S. Braga Centro de Investigação Geológica, Ordenamento e Valorização de Recursos (CIG-R), Universidade do Minho, Campus de Gualtar, 4710-057 Braga, Portugal e-mail: teresav@dct.uminho.pt

P. Gomes · T. Valente · J. A. Grande · M. L. de la Torre CIPIMS, Universidad de Huelva, Campus de la Rábida, Huelva, Spain

### Introduction

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Mining activity produces a large amount of not availed mate-35 rial that must be stored in appropriate conditions of stability, 36 security, and integration with the surrounding landscape 37 (Gomes 2011). These sterile materials, or mining wastes, are 38 generated at various stages of the production process. If they 39 are reactive, they will pose more environmental concern, due 40 to their participation in contaminant mineral-water interac-41 tions processes. So, abandoned mining lands represent one 42 of the most outstanding environmental risks, such as soil pol-43lution by potentially toxic metals and metalloids. These prob-44 lems affect many countries with historic mining activity 45(Thornton 1996; Fernández-Caliani et al. 2008), including 46Portugal. Across the country, about 200 abandoned mines 47 were identified (Santos Oliveira et al. 2002), which constitute 48 the national mining liability, whose abandonment occurred in 49an uncontrolled manner. In some of these mines, especially 50those which represented a great mobilization of sulfides, there 51is still a depreciation of environmental quality motivated by 52the reactive waste dumps (Valente 2004; Valente et al. 2011, 532012). In the weathering process of sulfides, oxidation gener-54ates acid mine drainage (AMD) mobilizing high amounts of 55acidity, sulfates, metals, and metalloids (Valente et al. 2009; 56 Q2 Gomes 2011; Sarmiento et al. 2012; Valente et al. 2012), from 57which results water and soil contamination. Also, the pollut-58ants may be transferred from these sites to nearby soils and 59water by wind-blown dispersion of fine particles. These 60 events are responsible for the impact of mining and ore treat-61 ment on the ecosystem, even at long distances. 62

The behavior of trace elements in AMD-affected systems is 63 regulated by many factors. First, element distribution into dif-64 ferent particle sizes depends of the presence of sorption sites 65 and of the mineralogical composition of soil (e.g., Acosta et al. 66

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2011). Fine particles, which include clav fraction, have high 67 specific area that retains high amounts of metals. In addition, 68 small particles are often soluble (Martínez-Martínez et al. 69 702010). Moreover, mobilization or fixation of elements in the soil can be regulated by the interactions between several bio-71geochemical processes, such as formation of secondary min-7273erals (Kabata-Pendias and Pendias 1984; Cox et al. 1995; Malpas et al. 2001). Besides, the Fe-rich materials that com-74pose the typical ochre-precipitates may adsorb or co-7576precipitate the toxic elements carried by AMD solutions 0377 (Valente et al. 2014). Therefore, to know the trace element distribution in fine soil particles is an important 78step for risk assessments of mining soils (Hardy and 79Cornu 2006; Acosta et al. 2009; Luo et al. 2011). 80

So, the main purpose of the present study comes in line 81 82 with this environmental concern. It deals with the subject of trace elements distribution in two distinctive sulfide 83 waste dumps, abandoned after decades of W and Au ex-84 85 ploitation, respectively, Cerdeirinha and Penedono (Portugal). These two sites (Fig. 1) were not yet the subject of 86 technical intervention for their environmental rehabilita-87 tion. The specific objectives of the present study are to 88 89 evaluate the degree of contamination by trace elements and to analyze the role of the clay size fraction in the 90 retention of contaminants. 91

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### Material and methods

Sites description

The two selected sites are located in regions with strong min-<br/>ing tradition for metals and so with a long historic in water and<br/>soil contamination by AMD (Valente and Leal Gomes 2009;<br/>Valente et al. 2012).9497

Cerdeirinha (Fig. 1) was exploited for W, in a skarn deposit 98 with sulfides, mainly iron sulfides. The waste dumps have 99 been formed by successive accumulation of sterile material 100from the treatment of tungsten ores. Penedono (Fig. 1), in a 101 granitic context, was exploited, mainly for Au, in quartz veins 102with sulfides. It was an important mining center for gold and 103other elements, such as arsenic, and the waste dumps are com-104 posed by sterile materials resulted from milling, hydrogravitic, 105and flotation operations. 106

Cerdeirinha has a more diverse mineral association, due to 107 the skarn nature of the ore deposit. Here, phyrrhotite and py-108 rite are the most abundant sulfides, while at Penedono, there is 109arsenopyrite with minor pyrite. A main difference between the 110two ore deposits relies on the absence of carbonates and cal-111 cium silicates in Penedono, which act as neutralizing minerals 112in Cerdeirinha. A more complete paragenetic description of 113the two ore deposits can be found in Valente et al. (2012). 114



Fig. 1 Localization of two study sites

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115 Sampling and analytical methods

The methodology includes sampling and preparation procedures followed by a general characterization of the mining soils, regarding texture, general properties, mineralogy, and chemistry in two grain size fractions: <2 mm (named bulk

120 sample) and the clay size fraction (<2  $\mu$ m).

121 In the field, the samples represent the first 20 cm of sub-122 strate on which plants grow on the waste dumps. At each 123 mine, the sampling sites represent the spatial variability, con-124 trolled by different aspects, including by the differences in the 125 vegetation cover.

Analytical procedures followed classical methodological
approaches for texture and general soil properties (Gomes
2011).

Mineralogy was analyzed by X-ray diffraction (XRD) ap-129130plied to both fractions. XRD was performed with a Philips X'pert Pro-MPD diffractometer, using Cu-Ka radiation, op-131erated with a 2 theta step size of 0.02° and a counting time of 1321.25 s. For the identification of clay minerals, the <2-µm 133fraction was separated by sedimentation and analyzed in ori-134ented aggregates of air-dried, ethylene gycol-treated, and 135136thermal-treated samples. The relative abundance of the mineral phases in both fractions was estimated by measuring the 137intensity of diagnostic diffraction peaks (Valente et al. 2012). 138139For the study of compositional and morphological aspects, scanning electron microscopy (on gold coated samples) was 140used, with a LEICA S360 microscope combined with an en-141 142ergy dispersive system (SEM-EDS). Also, transmission elec-143 tron microscopy was applied to provide compositional spectra, mineral identification by microdiffraction and morpholo-144145gy properties.

The chemical composition (Fe, Al, Mn, Cu, Ni, Pb, Zn, As, 146Ba, Bi, Cr, Sr, and W) of both size fractions was analyzed by 147inductively coupled plasma/mass spectrometry (ICP/MS) at 148Activation Laboratory, Ltd. (Actlabs, Canada). For that, sam-149 150ples were pulverized in an Agata mortar and submitted to 151aqua regia digestion. Duplicate samples and check precision blanks as well as certified standards (GXR series) for accuracy 152153were used.

#### 154 **Results and discussion**

The two studied sites have acidic soils (pH<4), with low con-155tents of nutrients. They differ in the contents of organic matter, 156presenting Cerdeirinha higher concentration (5 %) compared 157with Penedono, whose samples do not surpass the 1 %. Both 158soils can be considered as coarse soils, being classified as 159sandy loam, in accordance with the USDA texture classifica-160161tion system. So, the general properties are relatively similar in the two waste dumps. A detailed characterization of the min-162163 ing soils can be found in Valente et al. (2012).

#### Mineralogy

The mineralogy of the mining soils is presented in Table 1. 165According to XRD study, the most common minerals in bulk 166soil (<2-mm fraction) are quartz, k-feldspar, plagioclase, and 167 mica. In a general way, marcasite, pyrrhothite, pyrite, and 168 jarosite are prominent minerals at Cerdeirinha. In contrast, 169Penedono shows clearly lower variability in this fraction, with 170quartz and feldspar as dominant phases. This result reflects the 171differences in the two ore paragenesis. In fact, at Cerdeirinha, 172the skarn nature of the ore deposit in association with massive 173sulfides contributes to higher mineralogical diversity (Gomes 1742011), with minerals such as wollastonite and apatite. Table 2 175reveals the differences in  $<2-\mu m$  fraction in the two waste 176dumps. The clay size fraction is also more diverse at 177Cerdeirinha. Here, the iron-rich minerals jarosite and goethite 178are the most abundant phases, followed by a set of clay min-179erals, including kaolinite and 2:1 clay minerals. On the other 180 hand, at Penedono, kaolinite dominates, followed by jarosite 181 and clay minerals, while goethite is vestigial. 182

Geochemistry

#### Metal concentrations in different size fraction

Chemical composition of the soil is shown in Fig. 2, where 185concentrations are compared in the two size fractions: <2 mm 186 and  $<2 \mu m$ . It is possible to assess that elements such as Fe, 187 Al, and As are present in high quantities in the two waste 188 dumps (i.e., in a range between 1537.5 mg/kg for Al in the 189<2-mm fraction at Penedono and 1,765,000 mg/kg for Fe in 190the <2-µm fraction at Cerdeirinha). Despite this fact, Fe was 191the most abundant element in soil of the two waste dumps and 192 As has higher values at Penedono (with 25,875 mg/kg in the 193<2-µm fraction). Results suggest that the ore deposits have 194 influence in the concentrations of trace elements in the soil. 195Cerdeirinha is dominated by iron sulfides, thus increasing the 196concentration of Fe. The element As is present at Penedono in 197higher amount, due to dissolution of arsenopyrite, which is the 198main sulfide phase present in the ore deposit. 199

Furthermore, Ni, Ba, Cr, and Sr appear in low concentra-200tions at both sites. As opposed to Cerdeirinha, Penedono has 201lower concentrations of Mn, Cu, Zn, and Bi, showing a less 202 mineralogical and metal diversity (Table 1), also because soils 203enriched in quartz tend to have lower levels of trace elements 204(Silva et al. 2013). On the other hand, Penedono has higher 205concentration of Pb. Moreover, in this site, all elements are 206concentrated in <2-µm fraction, without exception. In con-207trast, some elements such Mn and W are in larger amount in 208the bulk soil fraction of Cerdeirinha. These exceptions can be 209 explained by the presence of wolframite crystals as a conse-210quence of low recuperation rates in the ore treatment. 211

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t1.1	Table 1         Mineralogical assemblage of the two size fractions: <2-µm fraction (clay minerals and associated minerals) and <2-mm fraction (bulk soil										
t1.2	Size fraction	Cerdeirinha ( <i>n</i> =6)	Penedono (n=8)								
t1.3	<2 μm	Jt >>II>Go>S>Gb>M-V>Cl>K	K>Jt>Il>S>Go								
t1.4	<2 mm	Q >>P>F>Mi>Mar>Pry=Mgs> Py=Go=Jt=Ap>Cm=Am=Wo>To	Q >>>F>Mi>P>Cm=Jt								

Jt jarosite, Il illite, Go goethite, Gb gibbsite, ML mixed-layered mica-vermiculite, S smectite, Cl chlorite, K kaolinite, Q quartz, F k-feldspar, P plagioclase, Mi mica, Cm clay minerals, Py pyrite, Mar marcasite, Pyr pyrrhotite, Mgs magnesite, Ap apatite, Am amphibole, Wo wollastonite, To Tourmaline

Although not detected in XRD study, such a presence was confirmed by SEM study.

Therefore, the results obtained for soil chemistry reflect the original differences in the ore paragenesis as already noted in the mineralogy. Cerdeirinha has the highest concentrations of Fe, reaching 15 %, which is in accordance with the abundance of iron sulfides. On the other hand, the highest contamination by As occurs at Penedono, with contents up to 3 %.

These results also show that both sites have a similar trend in what concerns the relation between the two size fractions. In fact, the highest concentrations are observed in the clay size fraction for most of the elements, with the exception of manganese and tungsten in the Cerdeirinha samples.

#### 225 Correlation between trace elements

226 Correlation analysis provides an effective way to reveal the
227 relationships between trace elements in soil. Tables 3 and 4
228 show the correlation matrixes for Cerdeirinha and Penedono,
229 respectively, considering the concentrations obtained in the
230 <2-μm fraction.</li>

Strong correlations occur between concentrations of some 231232trace elements in this fine fraction. However, the elements behave somewhat differently at both sites. At Cerdeirinha 233(Table 3), Fe is significant and positively correlated with Bi 234235(0.890). Al is correlated with Ba (0.819); Mn had close rela-236tionships with Zn (0.815) and As (0.823). Cu and Ni are re-237lated with Cr. These high correlations indicate that these elements associate in the parent materials and suggest similar 238geochemical behavior during weathering process, as observed 239

t2.1 **Table 2** Mineralogical composition of the mining waste dump soils estimated by XRD in the <2-μm fraction (clay minerals and associated minerals)

t2.2	Waste dumps	<2-µm fraction (%)								
t2.3			С	Il	M-V	Sm	K	Go	Jt	Gi
t2.4	Cerdeirinha	Average	5	10	6	8	4	9	51	7
t2.5	Penedono		-	19	-	13	40	1	27	-

*C* chlorite, *Il* illite, *M-V* mixed-layer mica-vermiculite, *Sm* smectite, *K* kaolin mineral, *Go* goethite, *Jt* jarosite, *Gi* gibbsite

for Bi. Bismuth occurs at the ore deposit of Cerdeirinha as 240native bismuth and bismuthinite (Valente et al 2012). Howev-241 er, in this Fe-rich paragenesis, dissolution of bismuth followed 242by precipitation of iron-rich minerals may take place. Convo-243lute bands of iron oxyhydroxides (probably goethite) with Bi 244inside iron-rich crusts were observed by SEM. In fact, previ-245ous studies revealed the tendency of Bi to co-precipitate with 246hydrous Fe oxides (Augustithis 1995), hence justifying the 247observed correlation between Fe and Bi. 248

Close correlation between Cu, Cr, and Ni indicates the 249sulfides, such as pyrite, chalcopyrite, sphalerite, gersdorfite, 250and other complex sulfides from the ore paragenesis (Valente 251et al. 2012) as the main common source of these elements. The 252same happens for Mn with As and Zn, reflecting the associa-253tion between ore minerals and sulfides. Identically, in geo-254chemical processes, Ba is commonly associated with K, in 255alkaline feldspars and biotite. So, the high correlation 256(0.819) of the pair Ba-Al is reflecting this lithophilic affinity. 257

At Penedono, the correlation matrix (Table 4) indicates that 258 Al is effectively correlated with Mn (0.939) and Sr (0.782) 259 while Mn appears correlated with Ba (0.765) and Pb is related 260



Fig. 2 Concentration and distribution of metals in different particle size fractions

<b>Q4</b> t3.1	t3.1	Table	3 Correla	tion matrix	of trace elem	ents concent	tration in Ce	rdeirinha (<2	2-µm fractio	n)				
	t3.2		Fe	Al	Mn	Cu	Ni	Pb	Zn	As	Ba	Bi	Cr	Sr
	t3.3	Fe	1.000											
	t3.4	Al	-0.722	1.000										
	t3.5	Mn	0.306	0.112	1.000									
	t3.6	Cu	-0.785	0.253	-0.359	1.000								
	t3.7	Ni	-0.176	-0.377	-0.283	0.512	1.000							
	t3.8	Pb	0.131	-0.672	0.014	0.439	0.721	1.000						
	t3.9	Zn	0.146	0.120	0.815	-0.378	-0.037	-0.029	1.000					
	t3.10	As	0.452	-0.206	0.823	-0.300	-0.391	0.240	0.507	1.000				
	t3.11	Ba	-0.375	0.819	0.341	-0.039	-0.195	-0.621	0.355	-0.180	1.000			
	t3.12	Bi	0.890	-0.826	0.208	-0.464	-0.100	0.422	-0.089	0.560	-0.646	1.000		
	t3.13	Cr	-0.548	-0.098	-0.495	0.921	0.745	0.653	-0.472	-0.386	-0.268	-0.232	1.000	
	t3.14	Sr	0.767	-0.820	-0.310	-0.472	0.254	0.249	-0.290	-0.164	-0.503	0.687	-0.103	1.000

261 with Bi (0.783). In turn, the clear association between Ni and 262 Zn (0.809) and Ni and Cr (0.976) stands out, which reveals a 263 common origin for all, possibly represented by the sulfides. At 264 this site, Bi correlates more strongly with Pb, which may also be reflecting its chalcophile affinity, which results in its 265266partitioning into sulfides, especially galena. In fact, Bi is com-267mon in late Pb-Zn veins or perigranitic hydrothermal vein deposits as that of Penedono. The strong correlations among 268Mn, Al, Sr, and Ba may reveal the lithophilic affinity of these 269 270elements and their link with plagioclase and micas, highly 271abundant in the clay size fraction as demonstrated by the 272XRD study.

#### 273 Accumulation factor

The enrichment in the clay size fraction with respect to the bulk soil can be expressed through the determination of an accumulation factor, which reflects the ratio between contents in both fractions. Figure 3 represents the accumulation factor  $(AF_x)$  for a set 278 of elements that pose greater environmental concern, calculat-279 ed by using Eq. (1) (Acosta et al. 2009). 280

$$AF_x = X_{\text{fraction}} / X_{\text{bulk}} \tag{1}$$

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As can be seen in Fig. 3, the accumulation factor (AF) for 284 Cerdeirinha showed consistent values  $\sim 1.0$  for Cu, Zn, and 285 As. Here, Ni and Bi and specially Pb showed higher enrichment trends (AF=14). At Penedono, the AF is generally 287 higher than at Cerdeirinha. Ni stands out with an AF around 288 28. 289

In a general way, the concentrations (Fig. 2) and AFs 290 (Fig. 3) of trace elements in the two fractions demonstrated 291 an enrichment of the finer fraction. Preliminary works reported that the preferential partitioning of trace metals is in the 293 clay fraction (Moore et al. 1989; Fernández-Caliani et al. 294

t4.1 Table 4 Correlation matrix (Person coefficients) of trace elements concentration in Penedono (<2-µm fraction)

t4.2		Fe	Al	Mn	Cu	Ni	Pb	Zn	As	Ba	Bi	Cr	Sr
t4.3	Fe	1.000											
t4.4	Al	-0.457	1.000										
t4.5	Mn	-0.240	0.939	1.000									
t4.6	Cu	-0.375	0.308	0.325	1.000								
t4.7	Ni	0.552	-0.170	-0.024	-0.187	1.000							
t4.8	Pb	-0.417	-0.238	-0.433	0.453	-0.392	1.000						
t4.9	Zn	0.418	0.264	0.360	-0.333	0.809	-0.709	1.000					
t4.10	As	0.102	0.583	0.556	-0.490	-0.084	-0.591	0.397	1.000				
t4.11	Ва	0.349	0.584	0.765	0.129	0.508	-0.547	0.682	0.435	1.000			
t4.12	Bi	-0.376	-0.423	-0.533	0.123	-0.334	0.783	-0.673	-0.624	-0.578	1.000		
t4.13	Cr	0.598	-0.307	-0.122	-0.116	0.976	-0.360	0.713	-0.245	0.431	-0.271	1.000	
t4.14	Sr	-0.579	0.782	0.649	0.335	-0.460	-0.139	-0.006	0.442	0.098	-0.461	-0.540	1.000

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Fig. 3 Average values for accumulation factor in two waste dumps

2008; Luo et al. 2011). These might be attributed to greater 295surface area per unit of mass of the fine particles, which in-296crease the adsorption capacity of this fraction (Wong et al 2972006). Furthermore, finer soil particles have greater content 298 299 of secondary minerals (clay minerals, Fe, Mn, and Al oxides and hydroxides and hydroxysulfates). In addition, metal en-300 301 richment in clay fraction was especially noticeable for Ni, Bi, and Pb at both waste dumps. Sposito (1989) suggested that 302 duo to isomorphous substitution of Ni for Fe or Mg in the 303 octahedral sheet of illite. Ni tends to accumulate in the finest 304 305 fractions. This may be an explanation for having a higher amount of these elements in the <2-µm fraction, especially 306 307 in Penedono, where there is a significant quantity of illite. 308 Also, Kohut and Warren (2002) indicated that Ni may occur in Mg-bearing minerals such as smectite. Moreover, chlorite is 309 normally composed by Mg, Al, and Fe and can also contain 310 311elements, such as Cr, Ni, and Cu, according to the same au-312 thor. This can justify the higher contents of these elements because they can be related to chlorite which was found in 313 314soils of the study area in Cerdeirinha (Table 2). Although, the accumulation of these elements are not limited to clay min-315erals but also appear in Fe oxides (Kataeva et al. 2004). As 316 317 well known (e.g., Bigham et al. 2002), oxide-hydroxides have 318 high sorption capacity of trace elements, such as Ni, Pb, Cu, 319 and Zn. In accordance with Ni et al. (2009), it is thought that 320 the main adsorbent for Pb are oxides phases, under conditions 321 of weathering, which can occur in both waste dumps. How-322 ever, it is known that quartz and feldspars are very weak ad-323 sorbent for trace metals, and in turn, the bulk soil fraction is largely composed by these primary minerals. So, trace ele-324 ments are not concentrated in this fraction, as seen in Figs. 2 325326 and 3.

In addition, clay fraction showed the highest levels of Fe 327 328 for both soils (Fig. 2), which mean that clay-sized minerals such as jarosite and goethite (Table 2) are significant sources 329 330 of Fe. Also, according with Arocena et al. (1995), these two phases can retain metals such as Pb and so play a significant 331role in metal enrichment in fine fraction. 332

333 The obtained results suggest a higher capacity of retention by the clay size fraction present at Penedono. Although 334 oxyhydroxides, such as goethite, are known by their high 335

sorption capacity of trace elements, in Penedono, goethite is 336 vestigial. Nevertheless, the mineralogical assemblage com-337 posed by kaolinite, illite, smectite, and jarosite seems more 338 effective than the one present at Cerdeirinha. In addition to 339 smectite, illite may also play a significant role in the retention 340 of trace elements, such as Ni, by means of isomorphous sub-341stitution of Fe in the octahedral sheet of illite. This may be an 342 explanation for having a higher amount of this element in the 343 <2-µm fraction, especially in Penedono, where there is more 344 quantity of illite. 345

Conclusion

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Metal and metalloid concentrations in soil reveal that ele-347 ments, such as Fe, Al, and As were present in high quantities 348 in the two waste dumps, being Fe the most abundant element. 349 The XRD study of Penedono shows lower mineralogical var-350 iability in the two analyzed fractions, in contrast with 351Cerdeirinha. These results suggest that the ore deposits have 352 strong influence on the mineralogy of the soil. 353

The accumulation factor in Penedono presents values 354higher than 1 for all studied elements. In Cerdeirinha, this 355 accumulation occurs for almost all analyzed elements, with 356 exception of Mn and W. This study indicated that the two 357 waste dumps are heavily polluted with metals and arsenic 358 and that these elements are concentrated in the finer fraction. 359Thus, it is believed that clay minerals and associated minerals, 360 especially jarosite, may play an important role in sorption of 361 toxic elements at both soils. On the other hand, this enrich-362 ment may enhance the environmental risk for pollutant dis-363 persion by wind, water, and animal activity. Hence, the higher 364 degree of contamination by toxic elements, especially arsenic 365 in Penedono, as well as the role of clay minerals, jarosite, and 366 goethite in retaining trace elements has management implica-367 tions. Such information must be carefully thought in the reha-368 bilitation projects to be planned for both waste dumps. 369

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