

# Co-processing of sulfidic mining wastes and metal-rich post-consumer wastes by biohydrometallurgy

Anne-Gwenaelle Guezennec, Kathy Bru, Jérôme Jacob, Patrick d'Hugues

## ▶ To cite this version:

Anne-Gwenaelle Guezennec, Kathy Bru, Jérôme Jacob, Patrick d'Hugues. Co-processing of sulfidic mining wastes and metal-rich post-consumer wastes by biohydrometallurgy. Minerals Engineering, 2015, pp.10.1016/j.mineng.2014.12.033. 10.1016/j.mineng.2014.12.033 . hal-01102658

# HAL Id: hal-01102658 https://brgm.hal.science/hal-01102658

Submitted on 13 Jan 2015  $\,$ 

**HAL** is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés. Dear Author,

Please, note that changes made to the HTML content will be added to the article before publication, but are not reflected in this PDF.

Note also that this file should not be used for submitting corrections.

## **AUTHOR QUERY FORM**

	Journal: MINE	Please e-mail or fax your responses and any corrections to:
ELSEVIER	Article Number: 4565	E-mail: corrections.esch@elsevier.sps.co.in Fax: +31 2048 52799

Dear Author,

Please check your proof carefully and mark all corrections at the appropriate place in the proof (e.g., by using on-screen annotation in the PDF file) or compile them in a separate list. Note: if you opt to annotate the file with software other than Adobe Reader then please also highlight the appropriate place in the PDF file. To ensure fast publication of your paper please return your corrections within 48 hours.

For correction or revision of any artwork, please consult <u>http://www.elsevier.com/artworkinstructions.</u>

Any queries or remarks that have arisen during the processing of your manuscript are listed below and highlighted by flags in the proof. Click on the 'Q' link to go to the location in the proof.

Location in article	Query / Remark: <u>click on the Q link to go</u> Please insert your reply or correction at the corresponding line in the proof							
<u>Q1</u>	Your article is registered as a regular item and is being processed for inclusion in a regular issue of the journal. If this is NOT correct and your article belongs to a Special Issue/Collection please contact n.rajaj@elsevier.com immediately prior to returning your corrections.							
<u>Q2</u>	Please confirm that given name(s) and surname(s) have been identified correctly.							
<u>Q3</u>	Please check the edit made in affiliation.							
$\frac{Q^2}{Q^3}$ $\frac{Q^4}{Q^4}$	The citation 'Battaglia et al. (1994)' has been changed to match the author name/date in the reference list. Please check here and in subsequent occurrences, and correct if necessary.							
<u>Q5</u>	This section comprises reference that occur in the reference list but not in the body of the text. Please position each reference in the text or, alternatively, delete it. Any reference not dealt with will be retained in this section.							
<u>Q6</u>	The decimal comma has been changed to a decimal point in Table 1. Please check, and correct if necessary.							

#### **MINE 4565**

## **ARTICLE IN PRESS**

#### 2 January 2015

#### Highlights

• Bioleaching of sulfidic mining wastes enables to produce lixiviant solution that can be used further for e-wastes leaching. • Biological re-oxidation of iron greatly enhances metals dissolution kinetics and yields during PCBs leaching. • Copper extraction above 90% was achieved in 24 h of PCBs leaching. • Microbiologically assisted leaching of waste PCBs is a promising way for metals recycling. • Decoupling the lixiviant production from the leaching process enables to avoid toxicity issues.

**ARTICLE IN PRESS** 

MINERALS ENGINEERING

Minerals Engineering xxx (2015) xxx-xxx

Contents lists available at ScienceDirect



5 6

9 10 **Minerals Engineering** 

journal homepage: www.elsevier.com/locate/mineng

Please cite this article in press as: Guezennec, A.-G., et al. Co-processing of sulfidic mining wastes and metal-rich post-consumer wastes by biohydromet-

# <sup>3</sup> Q<sup>1</sup> Co-processing of sulfidic mining wastes and metal-rich post-consumer <sup>4</sup> wastes by biohydrometallurgy

7 92 Anne-Gwénaëlle Guezennec,\*, Kathy Bru, Jérôme Jacob, Patrick d'Hugues,

8 03 BRGM, Water, Environment & Ecotechnologies Division, 3, av. Claude Guillemin, BP 36009, 45060 Orléans Cedex 2, France

#### ARTICLE INFO

13 13 Article history: 14 Received 7 August 2014 15 Revised 19 December 2014 16 Accepted 22 December 2014 17 Available online xxxx 18 Keywords: 19 Bioleaching 20 Mining wastes

- 21 WEEE
- 22 PCBs
- 23 Co-processing 24

#### ABSTRACT

The consequence of a strong economic growth in emerging countries combined with the rise of the world population is an increase in the demand for raw materials, leading to growing concern regarding their availability and the global efficiency of the supply chain. These tensions reinforce the need to associate the development of the recycling industry to the identification of new resources which could be used for the recovery of valuable materials. The purpose of this study is to develop a novel biological co-processing approach for the recovery of strategic metals in both sulfidic mining wastes and post-consumer wastes (WEEE). The principle of this treatment is based on two steps: mine wastes are biologically oxidized, resulting in the production of a ferric iron-sulfuric acid lixiviant solution which is used to leach base and other soluble metals contained in e-scraps. Batch tests were carried out using flotation tailings wastes containing 60% of pyrite and grinded Printed Circuit Boards (PCB < 750 µm) with a solid load of 2.5%. Two series of tests were conducted in order to study the influence of the ferric iron concentration and of the bacterial activity on metals dissolution. Results showed that a higher ferric iron concentration led to an increase in the dissolution rate of copper which is the main metal contained in the PCBs. Moreover, a dissolution yield of 98.3% was reached for copper after 2 days when bacterial activity was observed, corresponding to an increase of about 20% compared to the tests without bacterial activity. Finally, this study highlights the importance of the availability of ferric iron and of the bacterial oxidation of ferrous iron for the feasibility of this bioleaching process dealing with the recycling of PCBs.

© 2014 Published by Elsevier Ltd.

#### 46

#### 47 1. Introduction

Among the different types of secondary post-consumption 48 wastes, e-wastes represent the fastest growing and most problem-49 atic waste stream in the world. In the EU, 9.7 million tons were 50 produced in 2009 and this is estimated to grow to a projected 51 12.3 million tons per year by 2020 (Huisman et al., 2008). Recover-52 ing metals from e-waste is potentially more energy efficient than 53 54 mining raw material. However, they are highly complex in terms of structure and composition as very little consideration is given 55 to end-of-life reprocessing during the design and construction of 56 electronic goods. As a result it is not always possible to feed such 57 waste into conventional recycling streams. Many of the metals 58 59 on the "criticality" list are found in significant quantities in e-60 wastes (EU, 2014). The majority of the value in e-waste is in the 61 printed circuit boards (PCBs). On average 90% of the intrinsic eco-62 nomic value of PCBs is in the precious metals that they contain (Cui and Zhang, 2008). They also contain some critical metals such as 63

allurgy. Miner. Eng. (2015), http://dx.doi.org/10.1016/j.mineng.2014.12.033

\* Corresponding author. Tel.: +33 (0) 2 38 64 31 36. *E-mail address:* a.guezennec@brgm.fr (A.-G. Guezennec).

http://dx.doi.org/10.1016/j.mineng.2014.12.033 0892-6875/© 2014 Published by Elsevier Ltd. gallium. 65% of the world's gallium production ends up in PCBs and no current process for its recovery exists. Copper is also very important: it is much more abundant in e-wastes and PCBs than the higher value metals, that is why its recovery and recycling are crucial given the increasing scarcity and complexity of copper ore.

Pyrometallurgy is the traditional choice for metal refining from processed (usually upgraded) e-waste, resulting in the production of precious metal-bearing copper bullion (Tuncuk et al., 2012). It can be done within existing smelters treating mineral concentrates, where e-waste may be combined (10–15%) with a copper concentrate (Cui and Zhang, 2008). However, it is energy intensive and requires a relatively high grade feed material, and the ceramics from e-waste contribute to increase the final slag volume. Compared to pyrometallurgy, hydrometallurgical processes offer relatively low capital cost and are particularly suitable for small-scale installations (Tuncuk et al., 2012). An added advantage is their flexibility, offering a possibility for selective extraction of base and precious metals of interest in e-waste and PCBs. Since major metals exist in their elemental or alloy form in PCBs, their hydrometallurgical extraction has been tested

70

71

72

73

74

75

76

77

78

79

80

81

82

83

84

26

27

28

29

30

3 January 2015

157

158

159

160

161

162

163

164

165

166

167

168

169

170

171

172

173

180

181

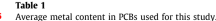
190

2

A.-G. Guezennec et al. / Minerals Engineering xxx (2015) xxx-xxx

85 using various oxidants (lixiviants; hydrogen peroxide, oxygen, 86 ferric iron, etc.) under acidic (HCl, H<sub>2</sub>SO<sub>4</sub>, HNO<sub>3</sub> etc.) or ammoni-87 acal and chloride leaching environments (Quinet et al., 2005; 88 Deveci et al., 2010; Huang et al., 2009; Kasper et al., 2011; 89 Tuncuk et al., 2012). These studies have shown a different degree of base and precious metals recovery efficiency. However, these 90 91 processes require the consumption of a high amount of chemi-92 cals. The use of microorganisms for the recovery of metals could 93 then be an economic and environmental alternative. Biohydro-94 metallurgy is an established technology for concentrates and 95 mineral processing wastes. It has been recognized as a potential 96 technology for the treatment of metallic wastes (Bryan et al., 97 2012). It has been used to recover copper and zinc from slags, gal-98 vanic sludge, fly ash and filter dust with recoveries close to 100% 99 (Krebs et al., 2006). Studies into the bioleaching of e-waste have 100 mainly involved the treatment of printed circuit boards (PCBs). 101 The use of organic acids produced by various fungi or biogenic 102 cyanide has been examined, particularly for the recovery of gold 103 and other noble metals (Brandl et al., 2001; Faramarzi et al., 2004; Brandl et al., 2008; Chi et al., 2011). However, such 104 105 approaches require the selective cultivation of specific microor-106 ganisms in circum-neutral media rich in organic substrates. This 107 often requires aseptic growth conditions and is unlikely to be 108 practical when treating large volumes of non-sterile e-wastes. 109 Further, the majority of the cyanide produced is consumed by 110 the copper which is in much higher concentrations relative to tar-111 get metals such as gold. Therefore, the use of ferric iron and/or 112 proton lixiviants produced by extreme acidophiles would be pref-113 erable and more practical. Indeed, there is no need for sterile con-114 ditions and culture media are relatively simple comprising key 115 nutrients such as sources of nitrogen, potassium and phosphorus. In mineral bioleaching the source of iron and sulfur which are 116 117 oxidized by the microbial community to produce the oxidizing acidic lixiviant solution is contained in pyrite or other sulfide 118 119 minerals. In the current e-waste bioleaching practices, this source 120 of iron and sulfur must be provided in addition to the nutritive 121 medium. This is usually realized in the form of ferrous sulfate 122 with acidity provided via pH control with sulfuric acid or through 123 addition of elemental sulfur. The studies published so far (Zhu 124 et al., 2011; Yang et al., 2009; Xiang et al., 2010; Liang et al., 125 2010; Karwowskaa et al., 2014; Hong and Valix, 2013; Ilyas et al., 2010; Cerruti et al., 1998) were carried out in shake flasks 126 in liquid media containing ferrous iron, sometime amended with 127 128 elemental sulfur and inoculated with pure or mixed cultures of iron- and sulfur-oxidizing microorganisms. Ground e-waste (usu-129 130 ally PCBs) was either added immediately, in a one-step process, 131 or following initial substrate oxidation (and thus lixiviant produc-132 tion) in an indirect two-step or multi-step process. The toxicity of 133 the e-waste on the microorganisms has been shown to be the 134 major problem preventing efficient leaching. Staggering the pro-135 duction of the lixiviant and the addition of the e-waste in a two-step process could then greatly increase leaching rates. Fur-136 thermore, PCBs are highly acid-consuming and require a high 137 138 degree of pH modification to maintain an acidic environment nec-139 essary for the microbial action and metal solubility. The necessary addition of chemical products will increase the operating cost of 140 141 such processes. In addition to deposits of secondary post-consumer wastes (the 142

143 classical target of the urban mining concept), old waste deposits



related to past mining and metallurgical activities can also be sig-144 nificant reserves of valuable base or strategic metals as well as 145 mineral substances. Before the 20th century, only a single or, at 146 best, a couple of metals were extracted from any given mine. The 147 other elements were either not detected by contemporary analyt-148 ical methodology or considered as mineralogical "exotica". As 149 demonstrated by European FP7 research project ProMine (http:// 150 ptrarc.gtk.fi/ProMine/default.aspx) these types of mining and met-151 allurgical residues contain not only rare and precious metals but 152 also appreciable amounts of "residual" base metals (Cu, Ni, Zn, 153 Co...) which must not be neglected in today's context of resource 154 scarcity and the environmental management of post mining 155 activities. 156

While current laboratory-scale studies provide evidence that biohydrometallurgical reprocessing of e-wastes and mining wastes are technically possible, the wider economics of such type of processes are unlikely to be favorable with the current state-of-theart and the competition with "classical" pyro and hydro options. Using an industrial ecology approach and including the potential costs associated with the "no action scenario" in terms of waste management, a co-processing concept was envisaged.

The purpose of this study is therefore to develop a novel biological co-processing approach for the recovery of strategic metals in both sulfidic mining wastes and post-consumer wastes (e-scraps). The principle of this treatment is based on two steps: mine wastes are biologically oxidized, resulting in the production of a ferric iron-sulfuric acid lixiviant solution which is used to leach base and other soluble metals contained in e-scraps. By decoupling lixiviant production from PCB leaching in a two-step process, it is assumed that issues of toxicity will be avoided.

Bioleaching batch tests were performed using flotation tailings wastes containing 60% of pyrite in a first step, and, in a second step, grinded Printed Circuit Boards (PCB < 750  $\mu$ m) with the lixiviant produced by the bioleaching of the sulfidic wastes. The results were compared to those obtained in abiotic conditions using a synthetic leaching solution instead of a solution produced biologically. 179

#### 2. Materials and methods

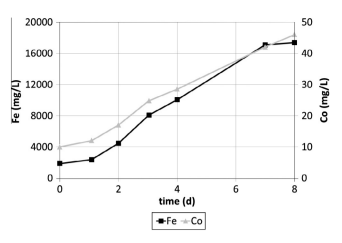
#### 2.1. Wastes

The mining wastes used in this study are flotation tailings 182 mainly composed of pyrite (60%). They contain also cobalt 183 (0.06%), copper (0.19%) and gold (1 g/t). These wastes have been 184 chosen for their high content of pyrite, which makes them partic-185 ularly suitable for bioleaching. The electronic wastes are computer 186 PCBs grinded first under 3 mm using a shear shredder and then 187 under 750 µm using a laboratory knife mill (Retsch, model 188 SM2000). 189

#### 2.2. Bacterial culture and nutrients

The tests were run using BRGM-KCC acidophilic and moderate 191 thermophilic (40 °C) microbial consortium which has already been 192 fully described (Battaglia et al., 1994; d'Hugues et al., 2003). The 193 predominant organisms in the culture are affiliated to the genera 194 *Leptospirillum, Acidithiobacillus* and *Sulfobacillus*. These bacteria are autotrophic and known as iron-oxidizer. They are also known 196 for their tolerance to high concentrations of various metals espe-197

Cu Fe Ni Pb Zn Sn Au Pd Ga Со Мо Ag g/kg ppm 215.1 24.3 2.4 17.5 14.6 34.6 393.6 143.9 42.4 7.9 29.2 11.2



**Fig. 1.** Iron and cobalt concentration vs. time in solution during the bioleaching of the sulfidic mining wastes.

cially copper (Guezennec et al., 2014). The culture used as an inoculum originated from BRGM stock culture, stored at -80 °C. The
culture was subcultured several times in batch mode and on pyrite
from 2 mL up to 2 L prior to the beginning of the batch test.

The culture was grown in a nutrient medium called "0Km" medium. It is a modified "9 K" medium (9 K without iron, "m" indicating modification of the basal salts) and was optimized for bacterial growth on cobaltiferous pyrite. Its standard composition is the following:  $(NH_4)_2SO_4$ , 3.70 g L<sup>-1</sup>; H<sub>3</sub>PO<sub>4</sub>, 0.80 g L<sup>-1</sup>; MgSO<sub>4</sub>·7H<sub>2</sub>O, 0.52 g L<sup>-1</sup>; KOH, 0.48 g L<sup>-1</sup>.

#### 208 2.3. Metal content in solid fractions

The metal content of initial PCBs and of leached residues was 209 210 determined using an analytical procedure derived from standard NF EN 13657. The size of the PCBs particles after comminution is 211 212 one of the main issues that must be encountered in PCBs analysis. 213 The size of the samples must be sufficient to avoid uncertainties 214 linked to the sampling representativeness. The analytical proce-215 dure was tested using several amounts of comminuted PCBs (1 g, 216 3 g and 5 g). From the results (data not shown) samples of 3 g were 217 finally selected for the following of the study. The quartered sam-218 ples of 3 g of materials (comminuted PCBs or leached residue) were dissolved in aqua regia using a closed system with reflow to avoid 219 220 reagents losses. The contact time between the samples and aqua regia was about 24 h at room temperature and then 2 h at 221 200 °C. After cooling, samples were carefully filtered and weighed. 222 223 The insoluble portion indicates ceramic and polymers. The leached portion represents the soluble metals and was analyzed by induc-224 tively coupled plasma optical emission spectrometry (ICP-OES) and 225 226 atomic adsorption spectrometry (AAS). The analyzed metals were: 227 copper, iron, nickel, lead, zinc, tin, silver, gold, palladium, gallium, 228 cobalt and molybdenum. The final solid residues were also charac-229 terized using optical microscopy as well as Scanning Electron 230 Microscopy to check the presence of undissolved metals, which 231 was not the case. The average metal content of PCBs samples used 232 in this study is given in Table 1.

#### 2.4. Laboratory apparatus

#### 2.4.1. Bioleaching of mining wastes

Bioleaching of the mining wastes was performed in order to produce the lixiviant solution used for the biotic leaching of PCBs. The tests were carried out in batch mode using 2 L laboratory scale glass reactors thermostated at 40 °C. The reactors were baffled; the agitation was performed using a dual impeller system consisting of a standard 6-blade flat Rushton turbine in combination with a 6blade 45° axial flow impeller. The impeller speed was set at 400 rpm. Air  $(120 L h^{-1})$  enriched with CO<sub>2</sub> (1%) was injected beneath the impeller at the bottom of the reactor via a stainlesssteel pipe. The tests were run in triplicate at 10% (w/w) pulp density and the inoculation was performed by adding 200 mL of the BRGM-KCC culture in each reactor. When the Fe<sup>3+</sup> concentration in the pregnant leach solution (PLS) reached the targeted value. reactors agitation and aeration were stopped. After natural decantation (around 12 h) the supernatant solution was filtered at 0.65 µm to remove particles while enabling bacteria to remain in the solution. This solution was then used as lixiviant solution for the biotic treatment of PCBs.

#### 2.4.2. PCBs leaching

Two series of PCBs leaching tests were performed in batch conditions using comminuted samples of PCBs with a solid load of 2.5%.

2.4.2.1. 1st series of PCBs leaching tests. The aim of the first series of experiments was to test the influence of initial Fe<sup>3+</sup> concentration as well as the type of lixiviant solutions (biological or chemical) on the PCBs leaching. For this purpose, the experiments were carried out in 500 mL Erlenmeyer flasks containing 10 g of comminuted PCBs and 400 mL of the biogenic lixiviant solution produced from the bioleaching of mining wastes (biotic tests) or 400 mL of a synthetic lixiviant solution (abiotic tests). For the biotic tests, two concentrations of  $Fe^{3+}$  were tested: 11 g L<sup>-1</sup> and 18 g L<sup>-1</sup>. The initial pH of the lixiviant solution was respectively 1.12 and 1.19, the initial Eh being respectively 862 and 826 mV (SHE). The synthetic solution was prepared by mixing deionized water with  $Fe_2(SO_4)3$ , 7 H<sub>2</sub>O salt in order to have a concentration of  $Fe^{3+}$  of about 18 g  $L^{-1}$ . The initial pH of the synthetic solution was 1.38 and the initial Eh was 914 mV (SHE). Biotic and abiotic tests were carried out in triplicate at ambient temperature and under magnetic stirring. Plastic paraffin film was used for sealing the Erlenmeyer flasks in order to prevent excessive evaporation.

275 2.4.2.2. 2nd series of PCBs leaching tests. The aim of the second series of experiments was to enhance the bacterial activity during PCBs 276 leaching by providing bacteria with more appropriate conditions 277 (i.e. temperature and oxygen transfer) and to compare the results 278 with those obtained in the first series of PCBs leaching experi-279 ments. The tests were carried out in duplicate using laboratory 280 scale glass reactors containing 1 L of the biogenic lixiviant solution 281 with a Fe<sup>3+</sup> concentration of 18 g L<sup>-1</sup> and 26 g of PCBs. The agita-282 tion device used in these tests had the same characteristics as 283 the one described for the production of the lixiviant solution. The 284 initial pH of the lixiviant solution was 0.90 and the initial Eh was 285

Table 2

Characteristics of the biological lixiviant solutions.

Type of PCB leaching test	pН	Eh (SHE) mV	${\rm Fe}_{\rm tot}~{\rm mg}~{\rm L}^{-1}$	$\mathrm{Fe}^{2+}~\mathrm{mg}~\mathrm{L}^{-1}$	$\mathrm{Cu}^{2+}\mathrm{mg}\mathrm{L}^{-1}$	$\mathrm{Ni}^{2+}~\mathrm{mg}~\mathrm{L}^{-1}$	$Pb^{2+}$ mg $L^{-1}$	$\mathrm{Zn}^{2+}~\mathrm{mg}~\mathrm{L}^{-1}$	$\mathrm{Co}^{2+}~\mathrm{mg}~\mathrm{L}^{-1}$
Erlenmeyer flasks (500 mL)	1.1 1.2	862 826	10966 18145	8 42	109 135	8.5 16	1.5 6	6.4 11	28 45
Reactor (1 L)	0.9	800	18430	151	112	14.7	5.9	15	46

Please cite this article in press as: Guezennec, A.-G., et al. Co-processing of sulfidic mining wastes and metal-rich post-consumer wastes by biohydrometallurgy. Miner. Eng. (2015), http://dx.doi.org/10.1016/j.mineng.2014.12.033

3

233

234

235

236

237

238

239

240

241

242

243

244

245

246

247

248

249

250

251

252

253

254

255

256

257

258

259

260

261

262

263

264

265

266

267

268

269

270

271

272

273

274

4

A.-G. Guezennec et al. / Minerals Engineering xxx (2015) xxx-xxx

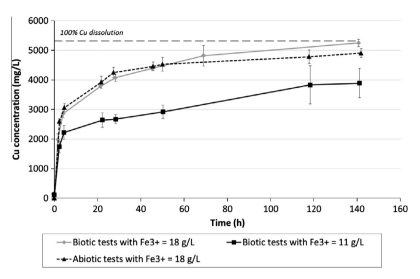


Fig. 2. Cu<sup>2+</sup> concentration vs. time during biotic and abiotic leaching of PCBs in Erlenmeyer flasks.

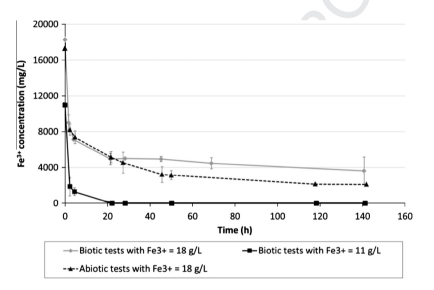


Fig. 3. Fe<sup>3+</sup> concentration vs. time during biotic and abiotic leaching of PCBs in Erlenmeyer flasks.

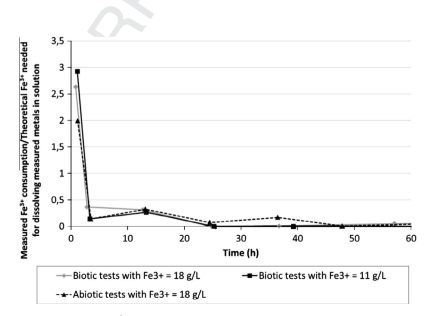


Fig. 4. Ratio "Measured Fe<sup>3+</sup> consumption/Theoretical Fe<sup>3+</sup> needed for dissolving measured metals in solution" vs. time during biotic and abiotic leaching of PCBs.

A.-G. Guezennec et al./Minerals Engineering xxx (2015) xxx-xxx

5

800 mV (SHE). The temperature of the reactors was maintained at
40 °C and the air flow rate was set at 0.5 L/h.

#### 288 2.5. Reactor monitoring

289 Each type of reactors was monitored regularly for temperature, 290 pH (adjusted below 1.8 by adding H<sub>2</sub>SO<sub>4</sub> (96%)) and redox poten-291 tial. Several samples of solution were taken and filtered at 292 0.45 µm in order to monitor the concentrations of Fe, Cu, Ni, Zn, Pb and Co (measured by AAS) and of Fe<sup>2+</sup> (determined by titration 293 using cerium(IV) sulfate). Bacterial cells were counted regularly in 294 biotic slurry samples. The leach residue was collected at the end of 295 296 each batch test and analyzed using the procedure described in Section 2.3. The final liquor of the biotic tests was also analyzed by 297 ICP-OES to determine Sn, Au, Ag, Ga, Mo and Pd final concentra-298 299 tions. These data were used further for the calculation of mass balance and metal dissolution yields. 300

#### 301 3. Results

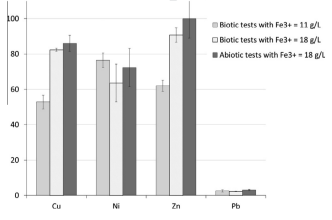
# 302 3.1. Production of the lixiviant solution by bioleaching of sulfidic 303 mining wastes

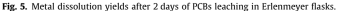
Fe and Co concentration follow-up during the bioleaching of the 304 305 sulfidic mining waste is shown in Fig. 1. The operating conditions 306 were not optimized since the objective of these tests was only to produce lixiviant solution, but the curve is similar to the one 307 308 obtained in previous study using KCC-BRGM microbial consortium 309 on this pyritic waste with first a lag phase of one day, and then a 310 regular increase of metal content. Experiments were stopped when 311 the Fe concentration in the solution reached the targeted value: 11 or 18 g L<sup>-1</sup>. These values were chosen by calculating the stoichiom-312 etric amount of oxidant (Fe<sup>3+</sup>) required to leach the metals con-313 tained in the PCBs. Indeed several authors showed that in 314 presence of Fe<sup>3+</sup>, the dissolution of PCBs metals is governed by 315 an oxidative mechanism based on the following equation (Yang 316 et al., 2014; Zhu et al., 2011; Bas et al., 2013): 317 318

320 
$$M^0 + 2Fe^{3+} = M^{2+} + 2Fe^{2+}$$
 (1)

Ferric iron in aqueous solution exhibits a standard reduction potential (Eh) as high as +0.77 V and could readily solubilize base metals contained in electronic scraps. Knowing the content of Cu, Ni, Zn, Pb, Fe and Sn in the PCBs it can be easily calculated from Eq. (1) that Fe<sup>3+</sup> concentration of 11 g L<sup>-1</sup> corresponds to a stoichiometric ratio of 1.2 and 18 g L<sup>-1</sup> to a stoichiometric ratio of 2.

The lixiviant solution obtained from the bioleaching of the sulfidic wastes is characterized by low pH (<1.3) and high redox





potential (800–862 mV), which indicates a good biological oxidiz-<br/>ing activity (Table 2). This is confirmed by the low concentration of<br/>Fe<sup>2+</sup> compared to total iron. The solution also contains small<br/>amounts of Cu, Ni, Pb, Zn and Co (Table 2).329<br/>330331

#### 3.2. PCBs leaching tests

# 3.2.1. Effects of $Fe^{3+}$ concentration and of lixiviant solution type (chemical or biological) on leaching

The first series of leaching tests aimed first at studying the influence of lixiviant solution production method (chemical or biological) on the metals dissolution by performing biotic PCBs leaching tests using the biogenic lixiviant solution obtained from the bioleaching of mining wastes and abiotic tests using a synthetic solution containing ferric iron and sulfuric acid. The concentration of Fe<sup>3+</sup> was set to 18 g L<sup>-1</sup> in both solutions. The 2nd aim was to study the influence of Fe<sup>3+</sup> concentration on the metals dissolution. Therefore biotic tests were also carried out using the biogenic lixiviant solution with a ferric iron concentration of 11 g L<sup>-1</sup>. Biotic and abiotic experiments were performed in Erlenmeyer flasks at ambient temperature and without air injection.

The dissolved  $Cu^{2+}$  and  $Fe^{3+}$  concentrations were monitored with time (Figs. 2 and 3). Copper dissolution and ferric iron consumption are very similar in biotic and abiotic tests with an initial  $Fe^{3+}$  concentration of 18 g L<sup>-1</sup>. The  $Cu^{2+}$  concentration increased rapidly during the first 5 h. A slower dissolution was then observed before the curve leveled off after 120 h with a final copper dissolution rate of about 96%. Ferric iron follows an inverse trend with a sharp decrease in the first hours of the tests followed by a slower decrease afterwards. In biotic tests no re-oxidation of  $Fe^{2+}$  ions in  $Fe^{3+}$  ions was observed. Moreover, the number of bacterial cells displayed no significant changes during the biotic tests remaining to a level closed to  $10^8$  bact/mL.

These results underline that there was no significant bacterial activity in tests performed with the biogenic lixiviant solution. This could be explained by the fact that the ambient temperature and the oxygen content in solution were too low for any activity of moderate thermophilic bacteria. Therefore, it can be concluded that, if carried out at ambient temperature, the ferric ion production method (chemical or biological) had no influence on the metals dissolution.

Regarding the influence of the  $Fe^{3+}$  concentration, Fig. 2 shows that higher  $Fe^{3+}$  concentration leads to increase the kinetics of the copper dissolution and the final recovery. Similar results were found by several authors (Yang et al., 2009; Zhu et al., 2011; Bas et al., 2013).

By comparing Figs. 2 and 3 it can be seen that in the biotic test with a Fe<sup>3+</sup> concentration of 11 g L<sup>-1</sup>, Cu dissolution rate was lower than the consumption rate of Fe<sup>3+</sup>. Indeed, even if the initial Fe<sup>3+</sup> concentration of 11 g L<sup>-1</sup> corresponds to a stoichiometric ratio of 1.2, all Fe<sup>3+</sup> was consumed after one day whereas Cu dissolution rate was only about 50% at that time. In biotic and abiotic tests with a Fe<sup>3+</sup> concentration of 18 g L<sup>-1</sup>, the consumption of Fe<sup>3+</sup> is not complete: Fe<sup>3+</sup> concentration decreases more slowly after 5 h of leaching and reaches a steady state even if all metals are not completely dissolved. This seems to indicate that ferric iron is no more involved in the leaching. One hypothesis could be that ferric iron forms complexes and is therefore not available for metals dissolution.

The ferric iron consumption was analyzed by calculating the ratio "Measured Fe<sup>3+</sup> consumption/Theoretical Fe<sup>3+</sup> needed for dissolving measured metals in solution" vs. time (see Fig. 4). This ratio take into account only the dissolution of Cu, Pb, Ni and Zn since they are the main metals contained in PCBs. In the first hours of leaching, this ratio is equal to 2 in the abiotic tests, which confirms that the metal dissolution corresponds to the oxidative process

333 334

335

336

337

338

339

340

341

342

343

344

345

346

347

348

349

350

351

352

353

354

355

356

357

358

359

360

361

362

363

364

365

366

367

368

369

370

371

372

373

374

375

376

377

378

379

380

381

382

383

384

385

386

387

388

389

390

391

392

6

### **ARTICLE IN PRESS**

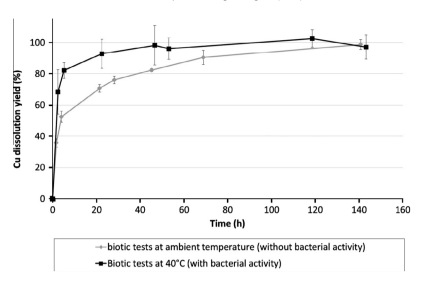


Fig. 6. Evolution of Cu dissolution yields with time during biotic tests at 40 °C and at ambient temperature (initial Fe<sup>3+</sup> = 18 g L<sup>-1</sup>).

**Table 3** Metal dissolution yields (in%) and absolute standard deviation  $\sigma$  (in%) after 2 days and at the end of tests of PCBs leaching at 40 °C and at ambient temperature (initial Fe<sup>3+</sup> = 18 g L<sup>-1</sup>).

	After 2 days of leaching		At the end of the leaching tests (about 6 days)			
	Biotic test at 40 °C (with bacterial activity)	Biotic test at ambient temperature (without bacterial activity)	Biotic test at 40 °C (with bacterial activity)	Biotic test at ambient temperature (without bacterial activity)		
Cu	98.3 ( <i>σ</i> = 12.8)	82.3 ( $\sigma$ = 0.9)	97.7 ( $\sigma$ = 5.3)	98.6 ( $\sigma$ = 3.2)		
Ni	$100 (\sigma = 2.8)$	$63.5 (\sigma = 10.7)$	$100 (\sigma = 5.9)$	84.3 ( $\sigma$ = 11.8)		
Zn	95.3 ( $\sigma$ = 14.0)	90.7 ( $\sigma$ = 4.1)	95.2 ( $\sigma$ = 6.7)	98.5 ( $\sigma$ = 5.8)		
Pb	2.7 ( $\sigma = 0.1$ )	2.3 ( $\sigma = 0.1$ )	$3.3(\sigma = 0.1)$	2.6 ( $\sigma = 0.1$ )		
Ga	NA	NA	$68.6 (\sigma = 5.3)$	42.9 ( $\sigma$ = 7.4)		
Sn	NA	NA	59.4 ( $\sigma$ = 1.6)	$6.8(\sigma = 1.8)$		

NA: Not analyzed.

406

415

presented in Eq. (1). In biotic tests, whatever the Fe<sup>3+</sup> concentra-393 tion, the ratio is higher than 2 in the first hours, underlining the 394 presence of other reactions consuming Fe<sup>3+</sup>. One hypothese which 395 396 is reported by several authors (Yang et al., 2014; Zhu et al., 2011; Xiang et al., 2010; Vestola et al., 2010; Wang et al., 2009) could 397 398 be the formation of jarosite precipitates (see Eqs. (2) and (3)), a 399 reaction which is favored by the presence of ions such as K<sup>+</sup> or NH<sup>4+</sup> brought by the nutrient medium: 400 401

403 
$$2SO_4^{2-} + 3Fe^{3+} + 6H_2O + K^+ = KFe_3(SO_4)_2(OH)_6 + 6H^+$$
 (2  
404

$$4/3SO_4^{2-} + 2Fe^{3+} + 4H_2O + 2/3NH_4^+ \\$$

$$= 2/3(NH_4)Fe_3(SO_4)_2(OH)_6 + 4H^+$$
(3)

After about 5 h, this ratio falls under 0.5 meaning that metals dissolution not only occurs through the oxidative action of Fe<sup>3+</sup> initially
present in the flasks but also through other phenomena. For example, part of zero-valence metals could be directly leached out based
on the following overall reaction, with H<sup>+</sup> coming from sulfuric acid:

$$2M^0 + 4H^+ + O_2 \rightarrow 2M^{2+} + 2H_2O \tag{4}$$

As mentioned by several authors (Hong and Valix, 2013; Zhu et al., 2011; Yang et al., 2014; Bas et al., 2013) Eq. (4) has a low kinetics which could explain the decrease in copper dissolution rate after the rapid consumption of  $Fe^{3+}$  at the beginning of the leaching experiments.

The metal dissolution yields obtained after 48 h of leaching are summarized in Fig. 5. After 2 days, the Cu extraction was 82.5% in biotic tests with an initial  $Fe^{3+}$  concentration of 18 g L<sup>-1</sup>, 86% in 423 abiotic tests and 52.8% in biotic tests with an initial Fe<sup>3+</sup> concentra-424 tion of 11 g L<sup>-1</sup>. Zn showed the same behavior as Cu in all types of 425 experiments. Fe<sup>3+</sup> concentration seems to have no influence on the 426 dissolution of Ni. Pb dissolution yields are significantly lower than 427 what can be expected given the pH and the redox of the solution, 428 probably because of Pb precipitation phenomenon (Ilyas et al., 429 2010; Ilyas et al., 2013). Ilyas et al. (2013) reported in particular 430 the formation of PbSO<sub>4</sub> precipitates during PCBs bioleaching in 431 shake flasks. 432

433

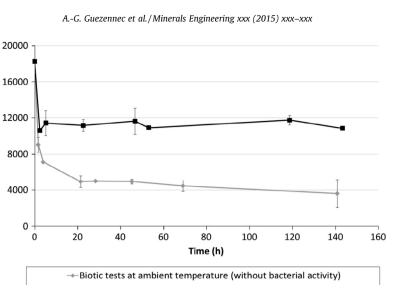
#### 3.2.2. Effects of bacterial activity on leaching

Since no bacterial activity was detected in the first series of bio-434 tic tests, a second series of leaching tests was run in order to 435 enhance bacterial activity and to favor biological oxidation of fer-436 rous iron produced by the oxidative leaching of PCBs. The experi-437 ments were performed using the biogenic lixiviant solution 438  $([Fe^{3+}]_{t0} = 18 \text{ g } \text{L}^{-1})$  at 40 °C which is known to be the optimal tem-439 perature for BRGM-KCC consortium. A small amount of air was 440 injected  $(0.5 L h^{-1})$  in order to provide oxygen to bacteria. The 441 results obtained were compared to those obtained with the biotic 442 tests of the first series of PCBs leaching experiment run with the 443 same biogenic lixiviant solution in shake flasks at ambient temper-444 ature. Cu concentration over time is shown in Fig. 6. A higher dis-445 solution yield of copper is observed. After 4 h of leaching, Cu 446 dissolution yield reached about 76% in these tests against 53% in 447 the tests in flasks at ambient temperature. A complete extraction 448 of copper, nickel and zinc is obtained after 2 days of leaching, 449 whereas the extraction in flask tests is only 82.3% for Cu, 63.5% 450

Fe<sup>3+</sup> concentration (mg/L)



7



---Biotic tests at 40°C (with bacterial activity)

Fig. 7. Fe<sup>3+</sup> concentration vs. time during biotic tests at 40 °C and at ambient temperature (initial Fe<sup>3+</sup> = 18 g L<sup>-1</sup>).

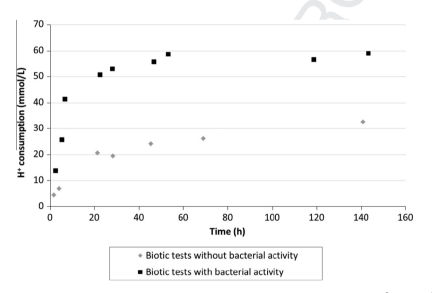


Fig. 8. H<sup>+</sup> consumption vs. time during biotic tests with and without bacterial activity (initial Fe<sup>3+</sup> = 18 g L<sup>-1</sup>).

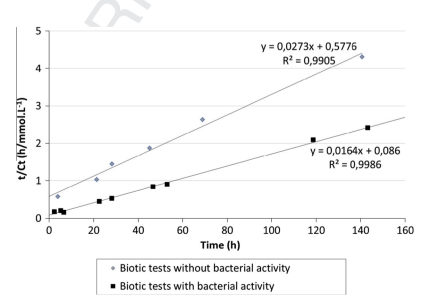


Fig. 9. Second-order model of H<sup>+</sup> consumption during biotic tests with and without bacterial activity (initial Fe<sup>3+</sup> = 18 g L<sup>-1</sup>).

3 January 2015

505

506

507

508

528

8

471

A.-G. Guezennec et al./Minerals Engineering xxx (2015) xxx-xxx

Table 4

Comparison between Cu extraction obtained by several authors and our results.

	One-step leaching					Two-step leaching			
	Xiang et al. (2010)	Ilyas et al. (2007)	llyas et al. (2013)	Wang et al. (2009)	Zhu et al. (2011)	Yang et al. (2009)		This study	
PCB content (g $L^{-1}$ )	20	10	100	7.8	8	25		25	
Time (day)	16	18	12	5	3	1	1.5	1	2
% Diss. Cu	95%	74%	90%	70%	92.9%	90%	100%	93%	99%

451 for Ni and 90.7% for Zn (Table 3). As previously observed. Pb disso-452 lution vield is very low, probably due to phenomenon of precipita-453 tion. Results of metals extraction at the end of the tests (about 454 6 days) is also given in Table 3. After about 6 days, both cases have similar dissolution yields for Cu, Zn and Pb. Regarding extraction of 455 Ni, Ga, Sn (both are only analyzed at the end of the experiments) 456 457 after 6 days of leaching, higher values are reached in tests at 458 40 °C; in particular, the Sn extraction rate is more than 8 times 459 higher. As can be expected with an acid ferric lixiviant solution, 460 no dissolution was obtained for Au, Ag, Pd and Mo (results not shown). Fig. 7 shows that this is correlated with a higher level of 461 Fe<sup>3+</sup> in solution. Indeed, even if both cases exhibit a drastic 462 decrease of Fe<sup>3+</sup> concentration in the first 2 h, Fe<sup>3+</sup> stabilizes then 463 at about 11 g L<sup>-1</sup> all along the tests at 40 °C whereas the concentra-464 tion of Fe<sup>3+</sup> decreases in the tests at ambient temperature. This dif-465 ference is probably linked to the re-oxidation of ferrous iron by the 466 467 oxidizing bacteria contained initially in the biological lixiviant solution according to the following equation: 468 469

$$4Fe^{2+} + O_2 + 4H^+ \xrightarrow{bacteria} 4Fe^{3+} + 2H_2O$$
(5)

According to Bas et al. (2013),  $Fe^{2+}$  is the only energy-yielding sub-472 strate for bacteria to support the bacterial growth and activity in 473 bioleaching process. A cycle between Fe<sup>3+</sup> and Fe<sup>2+</sup> is then built 474 475 and metals are continually leached out. This could explain the 476 increase in dissolution kinetics and in dissolution rates observed 477 in tests with bacterial activity compared to the ones carried out at 478 ambient temperature where no bacterial activity was detected. 479 The ratio between FeIII and FeII in the solution ([FeIII]/[FeII]) is also an important parameter which must be encountered to interpret 480 these results. In the test at 40 °C this ratio decreases to a value of 481 482 1.1 after 30 min and then increases to reach a value of 1.6 at the 483 end of the experiments (7 days of leaching). In the tests at ambient 484 temperatures this ratio decreases constantly to reach a value of 0.2 485 at the end of the experiments, which is much less favorable to metal 486 leaching. These results highlight the central role of ferric iron in the 487 metal dissolution process as well as the influence of the biological 488 recycling of ferric iron on the leaching kinetics of PCBs.

#### 3.2.3. H<sup>+</sup> consumption during PCBs leaching 489

Since the recycling of ferrous iron into ferric iron consumes H<sup>+</sup> 490 (see Eq. (5)), the consumption of protons during PCBs leaching and 491 492 its influence on metal dissolution efficiency were examined. As indicated by Yang et al. (2014), the PCB leaching process can lead 493 to several pathways of H<sup>+</sup> consumption: a direct H<sup>+</sup> consumption 494 495 through reactions with substances of PCB such as alkaline sub-496 stances, and an indirect H<sup>+</sup> consumption with bacterial oxidation 497 of Fe<sup>3+</sup>. H<sup>+</sup> consumption over time is plotted in Fig. 8. Fig. 8 con-498 firms that H<sup>+</sup> consumption is faster and higher in tests with bacte-499 rial oxidation of ferrous iron compared to the ones without this 500 oxidation. Yang et al. (2014) suggested that H<sup>+</sup> consumption kinetics can be described by the following second-order kinetic model: 501 502

$$\frac{t}{C_t} = \frac{t}{C_e}$$

 $+\frac{1}{k_2C_e^2}$ 

where t represents the time;  $C_t$  represents the amount of H<sup>+</sup> consumption;  $k_2$  represents the reaction rate constant of second-order kinetics; C<sub>e</sub> represents the total H<sup>+</sup> consumption in the whole process of bioleaching.

Fitting results of this second-order kinetic model are shown in 509 Fig. 9. In both cases, the correlation coefficient is over 0.99 suggest-510 ing that the second-order kinetic model is able to adequately 511 describe the progress of H<sup>+</sup> consumption during leaching. Calcula-512 tion of the reaction rate constant  $k_2$  gives 0.0031 [mmol/L]<sup>-1</sup> [h]<sup>-1</sup> 513 for tests with bacterial activity and 0.0013 [mmol/L]<sup>-1</sup> [h]<sup>-1</sup> for 514 tests without bacterial activity. Eq. (5) also allows evaluating the 515 total H<sup>+</sup> consumption in the whole process of leaching and then 516 the potential of metal dissolution rate. Potential value of the total 517 H<sup>+</sup> consumption is strongly increased when bacteria are allowed to 518 oxidize ferrous iron with a value of 60.9 mmol L<sup>-1</sup> compared to 519 36.6 mmol L<sup>-1</sup> for tests without bacterial activity, meaning that 520 favorable conditions for bacterial oxidation of Fe<sup>2+</sup> greatly 521 enhances the overall extraction of metals. It must be noted also 522 that given the importance of proton consumption during PCBs 523 leaching, lixiviant solutions produced from the bioleaching of sul-524 fidic mining wastes are not only interesting for their content in fer-525 ric iron but also because they provide high amounts of protons 526 under the form of sulfuric acid. 527

#### 3.3. Comparison of this study with results from the literature

Cu extraction obtained in the 2nd series of PCBs leaching exper-529 iments were compared to results from the literature. This compar-530 ison is not easy since experimental conditions are rather different. 531 For example, the initial PCBs content in the reactor can vary from 532 less than 1% to more than 10%. The particle size distribution is also 533 very different: PCBs are usually grinded under 250 µm; in some 534 cases only the finest fraction is used in the tests. These factors 535 can of course have a great influence on the results of leaching. It 536 must be noted also that in most of the papers dedicated to PCBs 537 bioleaching, experimental tests are performed in one step mode, 538 which means that the lixiviant solution is not produced in a first 539 reactor prior to PCBs leaching. Only a few papers propose PCBs 540 leaching in two steps mode. As can be seen in Table 4, in the case 541 of one-step leaching the final copper dissolution yield varies 542 between 70% and 95% for a leaching time ranging from 3 to 18. 543 The two-steps mode enables to reach better kinetics and higher 544 metal dissolution yield since the copper dissolution is almost com-545 plete in 2 days of leaching. Only Zhu et al. (2011) reached extrac-546 tion yield above 90% after 3 days of one-step leaching, but initial 547 PCBs content was 3 times smaller (8 g  $L^{-1}$  against 25 g  $L^{-1}$ ). In most 548 of papers dealing with one-step leaching problems of metal toxic-549 ity impacting bacterial activity were mentioned to explain low 550 leaching efficiency. The results obtained by Yang et al. (2009) in 551 2-steps mode using a culture of A. ferrooxidans are very similar to 552 the results obtained in this study, which highlights the interest 553 of this method of leaching. It confirms that decoupling lixiviant 554 production and leaching process could limit problems of metal tol-555 erance and thus greatly enhance leaching efficiency in terms of 556 kinetics but also in terms of metals dissolution yields. 557

Please cite this article in press as: Guezennec, A.-G., et al. Co-processing of sulfidic mining wastes and metal-rich post-consumer wastes by biohydrometallurgy. Miner. Eng. (2015), http://dx.doi.org/10.1016/j.mineng.2014.12.033

(6)

9

609

610

611

612

613

614

615

616

617

618

619

620

621

622

623

624

625 626

627

628

629

630

631

632

633

634

635 636

637

638 639

640

641 642

643 644

645

646

647

648

649

650

651

652

653

654

655

656

657

658

659

660

661

662

663

664

665

666

667

668

669

670

#### A.-G. Guezennec et al./Minerals Engineering xxx (2015) xxx-xxx

#### 558 **4. Conclusions**

559 The lixiviant solutions obtained from the bioleaching of sulfidic mining wastes using KCC-BRGM consortium showed a good per-560 formance for base metal recovery from waste PCBs (Cu, Ni, Zn, 561 Pb, Sn, Ga). The tests were carried out using a two steps approach 562 (bio-generation of the lixiviant oxidative solution/microbiologi-563 564 cally assisted leaching of the PCBs). The results indicated that 565 microbial assisted leaching was significantly more effective than a pure chemical leaching of PCBs. The extraction of metals was 566 mainly accomplished indirectly through oxidation by ferric iron. 567 The bacterial activity helped regenerating and maintaining a 568 569 higher concentration of ferric iron in solution. The comparison with data from the literature is significantly encouraging since 570 the rate and kinetics obtained in this study were much higher 571 and quicker than those obtained in previous studies. It must be 572 noted that in most of them, bioleaching was carried out using a 573 one-step approach, which confirms the interest of decoupling the 574 lixiviant production from the leaching process to avoid toxicity 575 576 issues.

#### 577 5. Uncited reference

578 **Q5** Fujita et al. (2013).

#### 579 **References**

- Bas, A.D., Deveci, H., Yazici, E.Y., 2013. Bioleaching of copper from low grade scrap TV circuit boards using mesophilic bacteria. Hydrometallurgy 138, 65–70.
- Battaglia, F., Morin, D., Ollivier, P., 1994. Dissolution of cobaltiferrous pyrite by Thiobacillus ferrooxidans and Thiobacillus thiooxidans – factors influencing bacterial leaching efficiency. J. Biotechnol. 32, 11–16.
- Brandl, H., Bosshard, R., Wegmann, M., 2001. Computer-munching microbes: metal leaching from electronic scrap by bacteria and fungi. Hydrometallurgy 59 (2-3), 319-326.
- Brandl, H., Lehmann, S., Faramazi, M.A., Martinelli, D., 2008. Biomobilization of silver, gold, and platinum from solid waste materials by HCN-forming microorganisms. Hydrometallurgy 94 (1-4), 14–17.
- Bryan, C.G., Watkin, E., McCredden, T., Wong, Z., Harrison, S., Kaksonen, A., 2012.
   From Wii to WEEE: the application of biohydrometallurgy to electronic wastes.
   Biohydromet '12, Falmouth, UK, June 18–20.
- Cerruti, C., Curutchet, G., Donati, E., 1998. Bio-dissolution of spent nickel-cadmium batteries using *Thiobacillus ferrooxidans*. J. Biotechnol. 62, 209–219.
- Chi, T.D., Lee, J., Pandey, B.D., Yoo, K., Jeong, J., 2011. Bioleaching of gold and copper from waste mobile phone PCBs by using a cyanogenic bacterium. Miner. Eng. 24 (11), 1219–1222.
- Cui, J., Zhang, L., 2008. Metallurgical recovery of metals from electronic waste: a review. J. Hazard. Mater. 158 (2–3), 228–256.
- Deveci, H., Yazıcı, E.Y., Aydın, U., Yazıcı, R., Akcil, A., 2010. Extraction of copper from scrap TV boards by sulphuric acid leaching under oxidizing conditions. In: Proceedings of Going Green-CARE.
- d'Hugues, P., Battaglia-Brunet, F., Clarens, M., Morin, D. 2003. Microbial diversity of
   various metal-sulfides bioleaching cultures grown under different operating
   conditions using 16S-rDNA analysis. In: Tsezos, M., Remoudaki, E.,
   Hatzikioseyian, A. (Eds.), International Biohydrometallurgy Symposium IBS
   2003. Hellas, Athens, pp. 1313–1324.

- EU 2014. REPORT ON CRITICAL RAW MATERIALS FOR THE EU. Report of the Ad hoc Working Group on defining critical raw materials, May 2014.
- Faramarzi, M., Stagars, M., Pensini, E., Krebs, W., Brandl, H., 2004. Metal solubilization from metal-containing solid materials by cyanogenic Chromobacterium violaceum. J. Biotechnol. 113 (1–3), 321–326.
- Fujita, T., Ono, H., Dodbiba, G., 2013. Recycling and evaluation of printed circuit board by physical separation and heat treatment. Proceedings Sardinia 2013, Fourteenth International Waste Management and Landfill Symposium, S. Margherita di Pula, Cagliari, Italy; 30 September – 4 October 2013.
- Guezennec, A.G., Hanke, M., Chmielarz, A., Joulian, C., Ménard, Y., d'Hugues, P., 2014. Bio-hydrometallurgy: an alternative to pyrometallurgy for copper recovery in a polymineral concentrate. Proceedings of Hydrometallurgy 2014 Congress – June 22–25, Victoria, Canada.
- Hong, Y., Valix, M., 2013. Bioleaching of electronic waste using acidophilic sulfur oxidizing bacteria. J. Cleaner Product. 65, 465–472.
- Huang, K., Guo, J., Xu, Z., 2009. Recycling of waste printed circuit boards: a review of current technologies and treatment status in China. J. Hazard. Mater. 164 (2–3), 399–408.
- Huisman, J. et al., 2008. Review of Directive 2002/96 on Waste Electrical and Electronic Equipment (WEEE) Final Report, 2008. United Nations University, Bonn, Germany.
- Ilyas, S., Anwar, M.A., Niazi, S.B., Ghauri, M.A., 2007. Bioleaching of metals from electronic scrap by moderately thermophilic acidophilic bacteria. Hydrometallurgy 88, 180–188.
- Ilyas, S., Lee, J., Chi, R., 2013. Bioleaching of metals from electronic scrap and its potential for commercial exploitation. Hydrometallurgy 131–132, 138–143.
- Ilyas, S., Ruan, C., Bhatti, H.N., Ghauri, M.A., Anwar, M.A., 2010. Column bioleaching of metals from electronic scrap. Hydrometallurgy 101, 135–140.
- Karwowskaa, E., Andrzejewska-Morzucha, D., Łebkowskaa, M., Tabernackaa, A., Wojtkowskab, M., Telepkob, A., Konarzewskab, A., 2014. Bioleaching of metals from printed circuit boards supported with surfactant-producing bacteria. J. Hazard. Mater. 264, 203–210.
- Kasper, A.C., Berselli, G.B.T., Freitas, B.D., Tenório, J.A.S., Bernardes, A.M., Veit, H.M., 2011. Printed wiring boards for mobile phones: characterization and recycling of copper. Waste Manage. 31 (12), 2536–2545.
- Krebs, W., Brombacher, C., Bosshard, P., Bachofen, R., Brandl, H., 2006. Microbial recovery of metals from solids. FEMS Microbiol. Rev. 20 (3–4), 605–617.
- Liang, G., Mo, Y., Zhou, Q., 2010. Novel strategies of bioleaching metals from printed circuit boards (PCBs) in mixed cultivation of two acidophiles. Enzyme Microbial Technol. 47 (7), 322–326.
- Quinet, P., Proost, J., Van Lierde, A., 2005. Recovery of precious metals from electronic scrap by hydrometallurgical processing routes. Miner. Metall. Process. 22 (1), 17–22.
- Tuncuk, A., Stazib, V., Akcila, A., Yazicic, E.Y., Devecic, H., 2012. Aqueous metal recovery techniques from e-scrap: hydrometallurgy in recycling. Miner. Eng. 25, 28–37.
- Vestola, E.A., Kuusenaho, M.K., Närhi, H.M., Tuovinen, O.H., Puhakka, J.A., Plumb, J.J., Kaksonen, A.H., 2010. Acid bioleaching of solid waste materials from copper, steel and recycling industries. Hydrometallurgy 103, 74–79.
- Wang, J.W., Bai, J.F., Xu, J.Q., Liang, B., 2009. Bioleaching of metals from printed wire boards by Acidithiobacillus ferrooxidans and Acidithiobacillus thiooxidans and their mixture. J. Hazard. Mater. 172, 1100–1105.
- Xiang, Y., Wua, P., Zhua, N., Zhang, T., Liu, W., Wu, J., Li, P., 2010. Bioleaching of copper from waste printed circuit boards by bacterial consortium enriched from acid mine drainage. J. Hazard. Mater. 184 (1–3), 812–818.
- Yang, Y., Chen, S., Li, S., Chen, M., Chen, H., Liu, B., 2014. Bioleaching waste printed circuit boards by *Acidithiobacillus ferrooxidans* and its kinetics aspect. J. Biotechnol. 173, 24–30.
- Yang, T., Xu, Z., Wen, J., Yang, L., 2009. Factors influencing bioleaching copper from waste printed circuit boards by *Acidithiobacillus ferrooxidans*. Hydrometallurgy 97 (1–2), 29–32.
- Zhu, N., Xiang, Y., Zhang, T., Wu, P., Dang, Z., Li, P., Wu, J., 2011. Bioleaching of metal concentrates of waste printed circuit boards by mixed culture of acidophilic bacteria. J. Hazard. Mater. 192 (2), 614–619.

671 672 673