IMPROVEMENT OF SELECTIVE COPPER EXTRACTION FROM A HEAT-TREATED CHALCOPYRITE CONCENTRATE WITH ATMOSPHERIC SULPHURIC-ACID LEACHING

IZBOLJŠANJE SELEKTIVNE EKSTRAKCIJE BAKRA IZ TOPLITNO OBDELANEH KONCENTRATI HALKOPIRITA Z LUŽENJEM Z ŽVEPLENO KISLINO NA ZRAKU

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The present work focuses on the extraction of Cu, suitable for production, directly from a leach solution of a chalcopyrite concentrate as an alternative to the conventional smelting and refining method. Firstly, the main aspects of the kinetics of chalcopyrite leaching in a sulphuric-acid solution and transformation reactions of chalcopyrite at high temperatures were briefly reviewed. Secondly, atmospheric-acid leaching experiments were performed as a function of the acid content and temperature under agitation. Direct leaching experiments resulted in very low and scattered Cu recoveries, hence revealing a sulphide passivation layer on the chalcopyrite. In order to get rid of this layer, the sulphides in the chalcopyrite were transformed into sulphates with heat treatment at 500 °C prior to the leaching process, in which a high dissolution of species was obtained. Finally, the leaching of the pre-heated concentrate resulted in a 99.82 % Cu extraction. Consequently, optimized process parameters were proposed by comparing the Cu extraction and the increased purity of the pregnant solution without Fe impurities.

Keywords: chalcopyrite concentrate, leaching, heat treatment, passivation layer, copper extraction, selectivity

1 INTRODUCTION

Copper has been one of the most important metals for over five thousand years.¹ Production of copper was easy for high-grade copper ores, for which traditional batch-smelting techniques were used before their grades were degraded. Afterwards, beneficiation of lower-grade ores was realised with the flotation technique prior to the smelting process. However, the traditional smelting technique became costly as the ore grades further decreased. On the other hand, there is currently an imbalance between the copper supply and the world demand.² In dealing with this problem, nowadays researchers are working hard to decrease process costs.

Chalcopyrite (CuFeS₂) is one of the most abundant and widely spread copper-bearing minerals,³ accounting for approximately 70 % of the Earth’s copper.⁴ In traditional smelting processes, the chalcopyrite concentrate of a desired grade is obtained with a multi-stage flotation of a sulphide ore, having fine grains of chalcopyrite dispersed in a matrix of various sulphide minerals and quartz.¹⁴ The chalcopyrite concentrate is then smelted in reverberatory flash furnaces.⁵,⁶ A major problem with smelting is the pollution of the environment, especially with sulphur dioxide.¹ At the same time, there is a decline in the copper-ore grades, often remarked upon as a future challenge in the production of copper for industry.⁷

Hydrometallurgical processes, an alternative to smelting, offer a high potential for treating chalcopyrite concentrate, apart from heap leaching, since they result in increased metal recoveries and reduced air pollution.⁴ In acidic media (sulfuric, hydrochloric and nitric acid), the concentrate can be leached under atmospheric or elevated pressures.⁸⁻¹⁴ Among them, the most promising one is sulfuric acid, since it can be readily produced at lower costs as the ore and, hence, the concentrate contain considerable amounts of sulphur.
The kinetics of leaching in sulfuric-acid media have been extensively analysed as chalcopyrite can be dissolved in a strong sulfuric acid. The main reactions are as follows:

\[ \text{CuFeS}_2 + 2\text{H}_2\text{SO}_4 \rightarrow \text{CuSO}_4 + \text{FeSO}_4 + 2\text{H}_2\text{S} \]  
\[ \text{H}_2\text{S} + \text{H}_2\text{SO}_4 \rightarrow \text{S}^0 + \text{SO}_2 + 2\text{H}_2\text{O} \]  

and

\[ 2\text{FeSO}_4 + 2\text{H}_2\text{SO}_4 \rightarrow \text{Fe}_3(\text{SO}_4)_2 + \text{SO}_2 + 2\text{H}_2\text{O} \]  

These reactions are very slow at atmospheric pressure because of the formation of a sulphide passivation layer on the chalcopyrite surface during the leaching process. There have been many efforts at the lab scale to overcome the difficulty of the concentrate being oxidized in the presence of several species (ferric ions, cupric ions, bacteria, pyrite oxygen, etc.). Additionally, some research has been done to increase the leach rate by favouring anodic reaction between the passivation layer and pyrite using the following redox reactions (4, 5, 6):  

\[ \text{CuFeS}_2 + 4\text{Fe}^{3+} \rightarrow \text{Cu}^{2+} + 5\text{Fe}^{2+} + 2\text{S}^0 \]  
\[ \text{CuFeS}_2 \rightarrow \text{Cu}^{2+} + \text{Fe}^{2+} + 2\text{S}^0 + 4\text{e}^- \]  
\[ 4\text{Fe}^{2+} + 4\text{e}^- \rightarrow 4\text{Fe}^{3+} \]  

Another process for increasing the leach rate can be roasting before leaching due to the fact that an oxidized and/or sulphatized concentrate has a higher dissolution rate because of the altered sulphide passivation layer. So, the pre-heating of chalcopyrite concentrate before the leaching in a different ambient would be an important step for the extraction of copper. As the passivation layer is composed of sulphur, it requires a sulphatization or oxidation process at elevated temperatures from 500–1000 K in an oxidizing ambient. Probable reactions during this treatment are extensively analysed in the literature. Among them, the most important ones for low temperatures are briefly given, in light of the relevant literature. L. Meunier et al. studied the behaviour of chalcopyrite in a stream of air in a lower-temperature regime of 573–823 K. The following reactions were mainly formed:

\[ 2\text{CuFeS}_2 \rightarrow \text{Cu}_2\text{S} + 2\text{FeS} + \text{S} \]  
\[ \text{FeS} \rightarrow \text{Fe}_2\text{O}_4 + \text{Fe}_3\text{O}_4 \]  
\[ \text{Cu}_2\text{S} \rightarrow \text{Cu}_2\text{O} \rightarrow \text{CuSO}_4 \rightarrow \text{CuO-CuSO}_4 \rightarrow \text{CuO} \rightarrow \text{CuFe}_2\text{O}_4 \]  

The first reaction (7) describes direct oxidation of chalcopyrite with released sulphur dioxide, occurring preferentially at temperatures higher than 900 K. At lower temperatures, sulphides are transformed into sulphates, as shown with reactions (8), (9) and (10). In the first step, Cu and Fe are separated form chalcopyrite in the form of sulphide and elemental sulphur. And then, these sulphides are oxidized as iron sulphide can be gradually transformed to magnetite and then hematite, as given in (9); copper sulphide can be easily converted into copper sulphate at low temperatures, and sequentially into tenorite and copper iron oxide at higher temperatures, as given in (10). Here, the produced copper iron oxide is reported to have a higher intrinsic resistance to an acidic attack than tenorite. As a result, chalcopyrite can be transformed into sulphates and oxides, which can be dissolved in sulphuric acid in line with the following reaction:

\[ \text{CuO} + \text{H}_2\text{SO}_4 \rightarrow \text{CuSO}_4 + \text{H}_2\text{O} \]  

The main aim of this work is to improve the leaching rate of chalcopyrite concentrate in a sulphuric-acid solution under atmospheric pressure in order to have a high Cu recovery at a low process cost. So, only sulphuric-acid solution without any catalysis was used for the leaching experiments since it can be produced from the same concentrate. In the first stage, direct leaching experiments were performed to disclose the low dissolution rate of chalcopyrite. This rate resulted in a limited extraction of Cu due to the existence of a sulphide layer on chalcopyrite and was, therefore, too far from establishing an industrial process. In the second stage, the concentrate was pre-heated at 500 °C to increase the extraction of Cu by converting sulphides into sulphates in the concentrate. In this case, the dissolution rate was found to increase enormously and a Cu extraction higher than 99.8 % was obtained in a leaching time of 180 min. Finally, a compromise between Cu and Fe extractions was made to have the pregnant solution free of Fe as much as possible.

2 EXPERIMENTAL WORK

2.1 Materials, analysis and characterization of samples

The agitated acid leaching method was used at atmospheric pressure for the Cu extraction from the chalcopyrite concentrate obtained from a newly discovered copper mine in the Kastamonu-Hanönü region of Turkey. The concentrate was produced by means of beneficiation of the sulphide copper ore via the flotation technique in a pilot-scale plant. In the first step, mineralogical/elemental analyses of the concentrate were obtained and summarized in Table 1. We can briefly say that the concentrate consists of 65 % chalcopyrite, 30 % pyrite and 5 % sphalerite.

In this work, mineralogical analyses of the samples were investigated with an X-ray diffractometer (Rigaku Primus IV) and elemental analyses were carried out with an inductively coupled plasma-atomic adsorption spectrometer (ICP-OES, Perkin Elmer Optima 2100DV) and an atomic absorption spectrometer (AAS, Perkin Elmer AAAnalyst 400). Additionally, a particle-distribution analysis was performed by means of an optical analysis (Nikon EPIPHOT 200); thus, the sizes of the chalcopyrite particles were found to be in the range of 1–60 μm.
and 80% of the particles were below 45 μm. During the leaching operations, sulphuric acid with a purity of 96% and a density of 1.84 g/cm³, obtained from the MERCK Company, and distilled water were used. The pre-heating and drying of the samples was done using a furnace (Protherm PL442T) and a stove (drying oven, MAS DT104), respectively. Finally, Cu and Fe extraction percentages were determined with the weighting method, using an ordinary lab scale with a 1 mg resolution.

### Table 1: Mineralogical/elemental analysis of chalcopyrite concentrate

<table>
<thead>
<tr>
<th>Mineral/Element</th>
<th>SiO₂</th>
<th>S</th>
<th>Cu</th>
<th>Zn</th>
<th>Au (g/l)</th>
<th>Fe</th>
</tr>
</thead>
<tbody>
<tr>
<td>%</td>
<td>1.12</td>
<td>36.44</td>
<td>21.10</td>
<td>21.26</td>
<td>0.39</td>
<td>33.34</td>
</tr>
</tbody>
</table>

#### 2.2 Experimental methods

Leaching experiments were performed in an 800 mL pyrex reactor, using the batch method. The reactor was placed in a bath, whose temperature was controlled by a temperature controller via a thermocouple within an error of ± 1 °C. The agitation of the leach solution was continuously maintained by an external propeller, rotated at a constant speed of 350 min⁻¹ throughout the processes. The complete system is shown in Figure 1a. Throughout this work, a constant volume of 600 cc of the leaching solution was used and its temperature was increased to the preselected value prior to the addition of 60 ± 0.5 g of the concentrate, which maintained a constant solid-content weight-to-volume ratio (w/v) of 1/10 in the solution. With the addition of the concentrate, the leaching process was commenced. After the process was finished, or a sample was taken out of the reactor, the residue was obtained using white-band filter paper. Upon determining its weight and carrying out elemental analyses, the amounts of Cu and Fe were determined and used for the calculation of the extraction percentage.

The process flowchart of this work is given in Figures 1b and 1c, for direct and pre-heated leaching processes, respectively. Initially, direct-leaching experiments without any pre-treatment of the concentrate were performed and very low extraction percentages were obtained. Secondly, the leaching process halted at the end of every 60-min period and the pregnant solution was replaced with a fresh solution in order to see the effect of acid consumption. Finally, the concentrate was pre-heated in air ambient at 773 K for 2 h in order to achieve an alteration of sulphide minerals.

### 3 RESULTS AND DISCUSSION

#### 3.1 Direct atmospheric acid leaching

Experiments of direct leaching were performed with the parameters of temperature and sulphuric-acid concentration under atmospheric pressure for 360 min. The parameters and corresponding Cu % extraction results are reported in Table 2. Although the results of the Cu % extraction given in Table 2 are somehow scattered, a net increase by the temperature is observed. Thus, the effect of the acid concentration seems to be lower than that of the temperature as the temperature increases. More importantly, the Cu % extraction stays below 17%, which is in agreement with the results reported in the literature.³¹ This behaviour contradicts the one explained above, in which galvanic reactions are reported to dominate the process due to the addition of pyrite, for the redox potential to be between 400 and 500 mV.¹⁵ In our case, as the concentrate has 30% pyrite, galvanic effects may also be expected for the reactions without any electric field. However, a probable built-in redox potential seems to stay below 400 mV, since pyrite stays undissolved. Otherwise, pyrite should be dissolved at the redox potential of 500 mV.³²–³³ Another evidence of this is the XRD analysis of the leach residue, given in Figure 2, where the pyrite and chalcopyrite peaks are clearly seen. This confirms that the dissolution processes of chalcopyrite and pyrite in sulphuric acid are very slow when there is no control of the redox potential of the leach solution.

![Figure 1: Experimental setup: a) flowchart of direct, b) pre-heated and c) leaching processes](image)

**Figure 1:** Experimental setup: a) flowchart of direct, b) pre-heated and c) leaching processes

**Slika 1:** Eksperimentalni sestav: a) potek neposrednega, b) toplotno predobdelanega in c) izluževanja za procesa

![Figure 2: XRD spectra of: a) chalcopyrite concentrate and b) residue of direct leaching process with 105 g of acid content at 85 °C for 360 min (chalcopyrite: ▲; pyrite: ■; sphalerite: ●; jarosite: ○)](image)

**Figure 2:** XRD spectra of: a) chalcopyrite concentrate and b) residue of direct leaching process with 105 g of acid content at 85 °C for 360 min (chalcopyrite: ▲; pyrite: ■; sphalerite: ●; jarosite: ○)

**Slika 2:**Rentgenograma: a) koncentrat halkopirita in b) ostanek neposrednega 360 min izluževanja pri 85 °C v vsebnosti kisline 105 g (halkopirit: ▲; piritt: ■; sfalerit: ●; jarozit: ○)
Table 2: Results of direct-leaching experiments lasting for 360 min

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Acid concentration (g/L)</th>
<th>% Cu extraction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Room Temperature (25)</td>
<td>15</td>
<td>8.99</td>
</tr>
<tr>
<td></td>
<td>45</td>
<td>15.83</td>
</tr>
<tr>
<td></td>
<td>75</td>
<td>12.97</td>
</tr>
<tr>
<td>45</td>
<td>15</td>
<td>14.46</td>
</tr>
<tr>
<td></td>
<td>45</td>
<td>12.10</td>
</tr>
<tr>
<td></td>
<td>75</td>
<td>13.55</td>
</tr>
<tr>
<td>85</td>
<td>15</td>
<td>15.01</td>
</tr>
<tr>
<td></td>
<td>45</td>
<td>16.42</td>
</tr>
<tr>
<td></td>
<td>75</td>
<td>14.06</td>
</tr>
</tbody>
</table>

On the other hand, an increase in the acid concentration from 45 g/L to 75 g/L at room temperature and 85 °C seems to cause a decrease in Cu recoveries of 2.86 % and 2.36 %, respectively. This might be explained with the rapid formation of the sulphur layer, reported in the literature.6

3.2 Effect of acid consuming

The proposed reactions for the chalcopyrite leaching, given with Equations (1) to (3), are acid-consuming processes. The most proton-consuming (acid-consuming) reaction is given below:2,3,4

\[
\text{CuFeS}_2(s) + 4\text{H}^+ (aq) = \text{Cu}^{2+} (aq) + \text{Fe}^{2+} (aq) + 2\text{H}_2\text{S}(aq) \quad (11)
\]

It is hypothesized that it is governed by two steps: (i) a rapid dissolution to establish the equilibrium (11) between the soluble species at the chalcopyrite surface and the bulk solid and (ii) a rate-determining diffusion of the soluble species away from the surface.

In order to test the effect of the above reaction, the influence of the acid consumption on the copper extraction was investigated with a solid-liquid ratio of 1/20 g/L in a H_2SO_4 concentration of 105 g/L at 60 °C. Before the process was started, a fresh acidic solution was prepared and held at a process temperature of 60 °C. At the end of every 60-minute period of leaching, agitation was stopped for the solid species to precipitate in the leaching bath. Five minutes after the completed precipitation, 560 mL of the solid-free leach liquor was drained out of the reactor. At this step, a 1-mg residue was taken out of the reactor to analyse the copper content via AAS. Then, the process was continued with an addition of 560 mL of the fresh solution of 105 g/L H_2SO_4 concentration into the reactor at 60 °C. This procedure was repeated every 60 min throughout the process.

The effect of the acid consumption was tested by determining the Cu content of the residue for each sampling. The results are given in Table 3, where a smooth decrease in the copper grade of the leach residue from 17.70 % to 17.10 % is clearly seen at 60 min and 300 min., respectively. However, the dissolution rate of Cu decreases in time and may tend to stop at around 17 %. This blockage would be caused by the passivation layer grown on the surface of chalcopyrite minerals to resist the leaching kinetics, as reported in the literature.15,35 It may be concluded that the acid consumption has little effect on direct leaching of the chalcopyrite concentration since the results are not scattered as in the previous case.

3.3 Atmospheric acid leaching of pre-heated chalcopyrite concentrate

The experiments above showed that the leaching of a chalcopyrite concentrate in a sulphuric-acid solution is hardly possible for industrial applications without an alteration of the passivation layer formed around chalcopyrite minerals.

Table 3: Copper concentration of the residue for each sampling when 560 mL of leach liquor was replaced with a new one

<table>
<thead>
<tr>
<th>Leaching time (min)</th>
<th>Added fresh solution (mL)</th>
<th>Cu % of leach residue</th>
</tr>
</thead>
<tbody>
<tr>
<td>60</td>
<td>560</td>
<td>17.70</td>
</tr>
<tr>
<td>120</td>
<td>560</td>
<td>17.40</td>
</tr>
<tr>
<td>180</td>
<td>560</td>
<td>17.23</td>
</tr>
<tr>
<td>240</td>
<td>560</td>
<td>17.10</td>
</tr>
<tr>
<td>300</td>
<td>560</td>
<td>17.10</td>
</tr>
</tbody>
</table>

Therefore, the concentrate was subjected to pre-heating at 773 K for 2 h in an atmospheric ambient before the leaching process. The change in the mineralogical property of the concentrate was determined with an XRD analysis whose spectra before and after the treatment are given in Figure 3. Dominant peaks of chalcopyrite and pyrite of the unprocessed concentrate are seen in Figure E. UZUN et al.: IMPROVEMENT OF SELECTIVE COPPER EXTRACTION FROM A HEAT-TREATED CHALCOPYRITE ...

![Figure 3: XRD spectra of: a) chalcopyrite concentrate, b) heated concentrate and c) residue of leaching process with 150 g acid content at 85 °C for 360 min (chalcopyrite: ▲; pyrite: ■; sphalerite: ●; iron sulphate: ○; hematite: □; chalcocyanide: △; tenorite: ▽)](image-url)
3a. But, in Figure 3b, the chalcopyrite peaks are disappeared whereas the pyrite peaks are still seen for the heated concentrate; here, hematite, iron sulphate and chalcocyanide peaks appear as dominant peaks.\textsuperscript{36,37} Moreover, a trace of tenorite is found to have little peaks in Figure 3b. It can be concluded from the results that the chalcopyrite in the concentrate is almost converted to CuSO\textsubscript{4} and CuO, which are soluble in a sulphide-acid solution, and to those compounds (hematite and iron sulphide) resistant to an acid solution.\textsuperscript{1,35}

The Cu and Fe extraction percentages in the leaching experiments of the pre-heated chalcopyrite concentrate were determined with the weighting method, via an elemental AAS analysis of the residue. The Cu extraction results for three different acid concentrations and temperatures as a function of the leach duration are given in Figure 4. The first thing to be mentioned is the fact that the leaching process seems to be nearly finished within 60 min as the lowest Cu extraction obtained is almost over 90\% for the lowest acid content and the lowest temperature. Additionally, the time dependence on the Cu extraction becomes almost flat at 85 °C, depicted in Figure 4, so that the reactions are nearly finished and independent of the acid concentration. Moreover, only hematite peaks are clearly observed and any trace of Cu bearing species is hardly found on the XRD graph given in Figure 3c, obtained for the residue of the experiment with the highest Cu recovery of 99.8\%. This enormous increase in comparison with the former experiments, obtained with the so-called direct leaching method, is certainly caused by the alteration of the sulphides in the concentrate of the latter case.

The pre-heating of the concentrate in air ambient makes it soluble in sulphuric acid, i.e., the solubility of CuSO\textsubscript{4} is higher than for Cu\textsubscript{2}S, CuS and CuFeS\textsubscript{2}. In addition, an increase in the solubility of hematite in a strong acid is also reported in \textsuperscript{38}.

Thus, a pregnant solution may also have FeSO\textsubscript{4} species making it hard to produce pure Cu directly from the solution (liquor).

Therefore, an optimization between purity and Cu recovery is necessary for the copper production, since Fe incorporated into the Cu structure especially deteriorates the electrical characteristics of copper. In other words, iron incorporation into copper should be as low as possible.

The optimization of the copper production was carried out by selecting the leaching process between the Cu and Fe recoveries, which were determined from the residue with the AAS analysis. The selectivity was calculated by subtracting Fe \% from the Cu \% recovery as a measure of the Cu purity in the solution, depicted in Figure 5, where Cu recoveries of over 97\% are reported for 65 °C and 85 °C. For example, the highest Cu extraction of 99.82\% was obtained from the process with the parameters of the temperature, 85 °C, acid concentration, 150 g/L, and duration, 180 min, as seen in Figure 4c: the highest Fe extraction of 37.31\% was also obtained, and hence, the selectivity of the leaching process was reduced to 62.51\%. In the figure, the selectivity goes through a peak with the acid concentration at 65 °C, but it decreases and seems to reach saturation at 85 °C. This behaviour is the evidence of an increased solubility of hematite at high acid concentrations and temperatures. As a result, the optimum leaching condition can be chosen as the peak on Figure 5, for the parameters of 65 °C and the 120 g/L acid concentration. Weaker acid concentrations and lower temperatures seemingly result in a low Cu extraction, in which some Cu is disposed in the residue.

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{figure4.png}
\caption{Copper extraction versus leaching time at: a) 90 g/L H\textsubscript{2}SO\textsubscript{4}, b) 120 g/L H\textsubscript{2}SO\textsubscript{4}, c) 150 g/L H\textsubscript{2}SO\textsubscript{4} acid concentrations (\textbullet\ 85 °C; ▲ 65 °C; ◆ 45 °C)}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{figure5.png}
\caption{Selectivity of Cu in the solution with respect to the acid concentration for leaching processes of 180 min}
\end{figure}
4 CONCLUSION

In the course of developing a hydrometallurgical process as an alternative to the conventional smelting and refining method, the production of Cu directly from the leach solution of the chalcopyrite concentrate would be a cornerstone. The most important step in this process is to have a Fe-free solution since Fe can be incorporated into the Cu structure during the subsequent electrowinning step. An observation of the selectivity between Cu and Fe can be taken as a measure of the purity of the produced copper. For achieving this aim, the main aspects of the kinetics of chalcopyrite leaching in the sulphuric acid solution is briefly reviewed in the light of the relevant literature. Additionally, transformation reactions of chalcopyrite at high temperatures are studied because direct leaching of chalcopyrite is a very slow process.

Direct-leaching experiments were performed at three different temperatures of (25, 45 and 85) °C, and acid concentrations of (15, 45 and 75) g/L. They resulted in very low Cu recoveries, below 15 %, due to the existence of a sulphide layer around the chalcopyrite species. In addition, a slight decrease in the acid concentration was observed at 85 °C, indicating acid consumption during the process.

Acid consumption was checked with a solid-liquid ratio of 1/20 g/L of a 105 g/L H₂SO₄ concentration at 60 °C. Its effect was measured with the change in the Cu concentration in the leach residue over time. The Cu concentration was found to decrease initially and have a saturation tendency, confirming that the leaching process was limited by the motion of the species through the passivation layer around chalcopyrite.

Finally, the pre-heating of the concentrate before the leaching at 500 °C for 120 min was found to have converted sulphides into sulphates. The subsequent leaching experiments were performed as a function of the acid concentration in g/L, the temperature and the time, using a solid-to-liquid ratio of 1/10 w/v at a stirring speed of 350 min⁻¹ under atmospheric pressure. As a result, a high dissolution rate was obtained because of the Cu recovery of over 90 % in the leaching process of 60 min. Thus, very high Cu recoveries of up to 99.82 % were achieved. Besides, over 36 % of Fe was found to have dissolved in the concentrate during the reactions because of the increased dissolution rate of hematite due to the increase in the temperature and acid concentration of the solution. Finally, the optimum selectivity of Cu and Fe was determined for the process with the acid concentration of 120 g/L, the temperature of 65 °C and the leach duration of 180 min. These can be taken as the proposed process parameters for the Cu production, optimized for the balance between the purity and Cu recovery.

Acknowledgment

Authors wish to thank ARGETEST Co. for experimental facilities. This work was supported by a grant from the Karabuk University BAP Project and, in part, by ASYA MADEN Co.

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