Chalcopyrite Leaching in Acidic Chloride Solution without Sulphates

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Abstract: Chalcopyrite leaching in acidic chloride solution without sulphates. The effect of dissolved oxygen and ferrous/ferric/cupric ions in solution on the leaching behavior of chalcopyrite and the phenomenology of the passivating layer formation on the chalcopyrite surface in an aqueous acidic chlorinated medium with dissolved oxygen and without sulphate are studied. The research was performed using pure chalcopyrite samples with particle sizes of +45 to -75 microns (-200 + 325 mesh). It was found that the formation of hydrogen sulfide (H2S) at room temperature and atmospheric pressure is a possible precursor of passivating species that prevent chalcopyrite dissolution.

Keywords: Chalcopyrite; Chloride; Ferrous; Leaching; Passivation.

1. Introduction

Most of the world’s copper production has been obtained from secondary sulfides, which can be oxidized, as well as from other mixed sulfides. However, the exploitation of these sulfides during centuries has led to their scarcity, leaving as an alternative the most abundant copper sulfide in nature, but also the most refractory in leaching processes: chalcopyrite, [1-4].

Hydrometallurgical technology offers the possibility of economically processing some refractory mineral species, [5]. Processes involving chlorides have been of great interest because of their ability to increase the leaching kinetics in acidic media at atmospheric pressure, [6,7]. The mechanism of chalcopyrite leaching is an issue currently under debate; however, since much of the research has been carried out under different experimental conditions, it is not always comparable, [8-11].

Third, [12], reported that in acidic sulphate solutions, in the voltage range of 450 to 650 mV/SHE, there is an increase of the leaching rates with dissolved iron ions. This low voltage effect is attributed to the release of ferrous ions, which increase the extraction rate of copper from chalcopyrite above a critical voltage (greater than 500 mV/SHE); below that voltage, the extraction rate drops. The results show that the kinetics of chalcopyrite leaching increases in ferrous sulphate solutions with dissolved oxygen, and diminishes in aerated solutions of ferric sulphate, [13], where the amount of copper recovered with 0.1 M ferrous sulphate (pH 1) is five times greater than that obtained with ferric sulphate.

The objective of the present research is to contribute new knowledge on copper leaching from chalcopyrite ore, in a chlorinated acid medium without sulphate, under weakly oxidizing conditions, with the coexistence of ferrous/cupric ions. Given that the precursors of the passivating layer and its formation, in the absence of sulphates, have not been previously established, this will be investigated under ambient conditions.

2. Experimental

The ore used corresponds to a massive chalcopyrite from Durango, Mexico. The ore was crushed in an agate mortar, in order to avoid sample contamination. The ore was then classified granulometrically and the particle size range selected for in the present study was +45 to -75 microns (-200 + 325 mesh). The mineralogical composition of this size fraction was determined by X-ray fluorescence (Siemens® SRS-3000 Sequential X-Ray Spectrometer), X-ray diffraction (Siemens® D5000 X-ray diffractometer) and SEM micrograph (TESCAN®, model Vega-3, electron-scanning microscope).

The solutions were prepared with double distilled water and analytical reagent grade chemicals (Merck®): sodium chloride, ferric chloride, potassium chloride, cupric chloride, ferrous chloride, etc. The pH was adjusted by adding concentrated hydrochloric acid. Iron and copper ions were added as chlorides and 1 M NaCl as NaCl. The additional amount of chloride required to reach 1.5 M [Cl] was added as KCl.
To determine the effect of dissolved oxygen and its participation in the reactions that take place on the surface of the chalcopyrite, tests were performed in solutions with and without dissolved oxygen. Aerobic tests were carried out by introducing oxygen into the system to maintain a dissolved oxygen concentration of 8 ppm. For the tests without dissolved oxygen, an inert gas (nitrogen) was bubbled previously into the solution to remove the majority of dissolved oxygen and lower its residual concentration to less than 1 ppm.

The experimental conditions that varied were dissolved oxygen concentration (for test sets with and without dissolved oxygen), the Fe$^{2+}$ and Fe$^{3+}$ concentrations (0, 0.025 and 0.05 M) and the Cu$^{2+}$ concentration (0 and 0.01 M). The potassium ion concentration varied according to the stoichiometric requirements to reach 1.5 M Cl$^{-}$ and 1 M Na$.^{+}$ Table I shows the experimental leaching conditions for the tests with and without dissolved oxygen, for the different combinations of initial ferrous/ferric/cupric ions. The parameters that remained constant during the tests were leaching time (400 h), pH (1), chlorine concentration (1.5 M), Na$^{+}$ concentration (1 M), solution volume (100 mL) and chalcopyrite ore leached (0.5 g).

In addition to these tests, a preliminary test 1-O2N2 (molar concentrations: Fe$^{2+}$ 0.50/ Cu$^{2+}$ 0.01/ Cl$^{-}$ 1.5 M at pH 1) was carried out, leaching during for 400 h in an aqueous solution with dissolved oxygen, after which the reactor was hermetically sealed so that no more oxygen could enter the system. The purpose to initially have oxygen in the solution was to initiate the H$_2$S generating reactions. The reactor was later hermetically sealed to prevent the H$_2$S produced from escaping, allowing it to act as a precursor for the passivating species on the chalcopyrite surface.

The residue from each leaching test was recovered from the aqueous system by vacuum filtration, using a 0.2 μm micro filter that retained the solid residues; these were washed with abundant water acidified to pH 1.6 to recover the solids adhered to the reactor walls and to avoid the precipitation of iron hydroxides. The residues were analyzed by SEM and X-ray fluorescence.

### 2.1. Experimental methodology

A volume of 100 mL of solution and 0.5 g of chalcopyrite were combined in a glass reactor, adjusting to pH 1 with concentrated hydrochloric acid. The slurry was placed in an orbital mechanical stirrer at 215 rpm (LabTech® shaking incubator model LSI-3016R) during 400 h at room temperature. The reactors were used for both the aerobic and anaerobic tests. In the anaerobic experiments, a screw cap, hermetically closed with parafilm, was employed to avoid the access of oxygen into the system. In the tests with dissolved oxygen, the same reactors were used, but with a cap that allowed oxygen introduction into the solution, diverting the exit flow of air into a flask containing silver nitrate (indirect method to qualitatively determine the formation of hydrogen sulfide, see Fig. 2). Of the total H$_2$S formed during the acid leaching of chalcopyrite, only a part was carried into the silver nitrate solution by the injected air, some of which was lost to the atmosphere; the rest stayed in the leaching solution. For this reason, the experimental set-up and the results obtained are only intended to confirm H$_2$S generation, not to quantify its production.

It is known that the rate of dissolution in chlorinated or sulphated solutions does not increase with the stirring velocity, [14], because the rate of dissolution is not controlled by the mass transfer in the solution, especially considering the lengthy experimental times. Therefore, agitation does not need to be excessive, rather just sufficient to ensure that the ore does not become agglomerated and the particles remain suspended. This guarantees that the mixture remains homogeneous over the entire leaching reactor, [15]. To avoid the solution loss due to sampling, the volumes removed (3 mL aliquots) were replaced by the same volume of solution of the same initial concentration, [16]. The total volume of solution was monitored and maintained constant with double distilled water to compensate for loss by evaporation.

Samples were taken every 100 h for chemical analysis, keeping the temperature in the range of 25 to 30 °C. The analysis of total copper and iron in the aqueous samples was made by atomic absorption spectrometry (GBC Scientific® model SenSAA). The concentration of ferrous ions was determined by the phenanthroline spectrophotometric method (Rayleigh® model UV1601 UV/VIS spectrophotometer), [17].

### 3. Results and discussion

#### 3.1. Results

Table I and Fig. 1 show the different experimental conditions and the results of the tests made on the solutions, with
and without dissolved oxygen, after 400 h of leaching. The tests without dissolved oxygen are from 1-N2 to 6-N2, without added sulphates, at pH 1, constant 1 M sodium and 1.5 M chloride concentrations; test 4S-N2 was performed under the same above conditions, but with the addition of iron and copper ions as sulphates. The tests with dissolved oxygen are from 1-O2 to 6-O2, without added sulphates, at pH 1, constant 1 M Na\(^+\) and 1.5 M chloride concentrations; test 4S-O2 was performed under the same above conditions, but with the addition of iron and copper ions as sulphates.

In the tests with dissolved oxygen, the analysis of the precipitate formed by the reaction of silver nitrate with the H\(_2\)S coming from the reactor showed that it was the same in all tests. Fig. 3 shows test with SEM micrograph of the sediments from the silver nitrate solution of test with oxygen, that there is evidence for the formation of Ag\(_2\)S\(_{(s)}\) in all tests, but cannot be quantified due to the co-precipitation of AgCl\(_{(s)}\) caused by the entrainment of HCl\(_{(v)}\) and its introduction to the flask.

A series of preliminary tests were performed using the hermetically capped leaching reactors, containing ambient oxygen, where no additional oxygen was introduced. Fig. 4 shows the SEM micrograph and X-ray diffraction of the altered chalcopyrite leaching residue from the test with a large quantity of ferrous ion (1-O2N2, molar concentrations: 0.50 Fe\(^{2+}\)/0.01 Cu\(^{2+}\)/1.5 Cl\(^-\)) at pH 1, after 400 h of leaching.

Fig. 5 shows a comparison of tests 4S-O2 (molar concentrations: 0.05 Fe\(^{2+}\)/0.01 Cu\(^{2+}\)/1.5 Cl\(^-\), with dissolved oxygen) and 4S-N2 (molar concentrations: 0.05 Fe\(^{2+}\)/0.01 Cu\(^{2+}\)/1.5 Cl\(^-\), without dissolved oxygen), in a sulphate environment, after 400 h of leaching.

### 3.2. Discussion

After performing the leaching tests as described above, the influence of dissolved oxygen on the solution and of the ferrous/cupric/ferric ions in an acidic chlorinated system can be analyzed.

#### 3.2.1. Effect of dissolved oxygen on acidic chlorinated solutions

The results of dissolving copper during 400 h, given in Table I and Fig. 1, show that in all cases, more copper dissolves in
solutions with dissolved oxygen. Both with and without oxygen in acidic chloride media, chalcopyrite can react to form stable cuprous ion and solids (Equation 1, [10]).

$$2 \text{CuFeS}_2 + 4 \text{H}^+ \rightarrow \text{Cu}_2\text{S}_2 + 2 \text{Fe}^{2+} + 2 \text{H}_2\text{S}$$ (1)

Reaction 1 represents a non-oxidative dissolution, documented by Nicol, et al., 2010, [10], in which covellite ($\text{Cu}_2\text{S}_2$) is formed at low solution potentials (550 mV/SHE). The partially vacant valence bonds of $\text{Cu}_2\text{S}_2$ has strong contributions from the 3p orbital of the sulfur, which lead to a more correct description of covellite as ($\text{Cu}^+$)$_2$($\text{S}^2-$)$(_2\text{S}_2$; the oxidation state of copper in the majority of chalcogenides is $\text{Cu}^+$, [18].

Oxygen, in addition to being an oxidant, is a catalyst for the further oxidation of ferrous and cuprous ions on the surface of the chalcopyrite. A high concentration of dissolved oxygen usually leads to a more positive solution potential; similarly, a more acidic solution (pH close to 1.0) also favors the oxidation of ferrous to ferric ions by the dissolved oxygen, consequently, higher potentials are found under these conditions, [19]. However, it has been found that the ferrous ion tends to prevail in highly acidic solutions, [20], and furthermore, in the present study, the difference in solution potential in solutions without and with dissolved oxygen was only 10 mV on the average.

Therefore, the catalytic action of dissolved oxygen may be explained by an adsorption of the gas on the surface of the sulphated ore, causing polarization of the covalent bonds in the crystal lattice and forming a negatively charged superficial layer; this perhaps would attract positive ions when in contact with the solution, [21].

This effect destabilizes the structure, which in turn, increases the diffusion rate through the passivating layer; a destabilizing effect caused by the interaction of the unpaired electrons of the adsorbed oxygen with the electrons in the ore is possible. The localized weakening of the ore’s structure due to the adsorption of oxygen would imply a decrease of the activation energy of the dissolving process. The catalytic action of the dissolved oxygen would be due to its high electronegativity and paramagnetism, which gives it great oxidizing power from the thermodynamic standpoint, but because of its low kinetic reactivity, the chemical oxidation reaction occurs slowly, allowing other oxidizing agents present (in this case, cupric ions coexisting with ferrous ions) to react, favoring the oxygen-ore interaction.

![SEM micrograph of the silver nitrate solution sample sediments of test 3-O2 (molar concentrations: 0.05 Fe$^{2+}$/1.5 Cl$^-$/M), pH 1, with dissolved oxygen, without addition of sulphates, after 400 h of shaking.](image)

<table>
<thead>
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<th>Norm. mass percent (%)</th>
<th>Spectrum</th>
<th>S</th>
<th>Cl</th>
<th>Ag</th>
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<td>16.54</td>
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<tr>
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<td>0.70</td>
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<tr>
<td>Sigma:</td>
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<td>2.63</td>
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<tr>
<td>Sigma mean:</td>
<td>0.15</td>
<td>1.21</td>
<td>1.31</td>
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</table>
In the tests with dissolved oxygen there was more reaction of the ore, even though on the surface there was no detectable product layer. The explanation for this is that all of the hydrogen sulfide (H$_2$S) generated was carried by the air stream without the formation of a product layer, or, conversely, that the dissolved oxygen and the hydrogen sulfide react to form SO$_2$, although this is unlikely. In the tests without dissolved oxygen there was very little reaction, as reflected in the low level of copper dissolution and in the absence of a product layer on the surface of the ore.

### 3.2.2. Effect of cupric and ferrous ions in acidic solution

The analysis of the residues from all the experiments does not show the formation of chlorine-copper species; furthermore, there is no evidence of covellite (Cu$_2$S$_2$) formation, as would be expected, probably because its quantity was too little to detect.

The improvement in the leaching rate of chalcopyrite at low voltages (below a critical voltage of about 550 mV/SHE) is possible only when the ferrous and cupric ions coexist in the solution due to a synergistic effect between them, [22]. A two-stage reductive/oxidative dissolving model is proposed to

<table>
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<th>Component species</th>
<th>Composition</th>
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<td>Akaganeite</td>
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<td>Iron hydroxide sulphate</td>
<td>2Fe(OH)SO$_4$</td>
<td>Cu</td>
<td>11.9%</td>
</tr>
<tr>
<td>Ramsbeckite</td>
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<td>8.57</td>
<td>6.43</td>
<td>9.28</td>
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<td>Sigma mean:</td>
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<td>4.13</td>
<td>4.29</td>
<td>3.21</td>
<td>4.64</td>
<td>5.40</td>
</tr>
</tbody>
</table>

**Fig. 4:** SEM micrograph and X-ray diffraction and fluorescence of altered chalcopyrite leaching residue of test 1-O2N2 (molar concentrations: 0.50 Fe$^{2+}$/0.01 Cu$^{2+}$/1.5 Cl$^-$ M), pH 1, with dissolved oxygen, without added sulphates, after 400 h of leaching.
explain this phenomenon, assuming the intermediate formation of chalcocite (Cu$_2$S) during the oxidation of the chalcopyrite, Equation 2, and the subsequent rapid oxidation of the chalcocite according to Equation 3 or Equation 4, [23].

\[
\text{CuFeS}_2 + 3 \text{Cu}^{2+} + 3 \text{Fe}^{2+} \rightarrow 2 \text{Cu}_2\text{S} + 4 \text{Fe}^{3+} \quad (2)
\]
\[
\text{Cu}_2\text{S} + 4 \text{Fe}^{3+} \rightarrow 2 \text{Cu}^{2+} + \text{S} + 4 \text{Fe}^{2+} \quad (3)
\]
\[
\text{Cu}_2\text{S} + 4 \text{H}^+ + \text{O}_2 \rightarrow 2 \text{Cu}^{2+} + \text{S} + 2 \text{H}_2\text{O} \quad (4)
\]

The model predicts the formation of an intermediate chalcocite product, which is oxidized more easily than chalcopyrite, and this leads to the increased extraction of copper at low solution potentials, in the presence of the cupric and ferrous ions.

The presence of ferrous ions coexisting with cupric ions was beneficial to increase the dissolution kinetics of copper. The action of cupric ions in solution improves the leaching rate in a sulphate environment and the presence of ferrous ions is important for that purpose, [24]. The average increase in copper
extractions in those experiments where ferrous and cupric ions coexist in chloride media, was 48.5%.

On the surface of the chalcopyrite, the oxidation of ferrous ions by the dissolved oxygen is faster than in the liquid phase (Equation 2). The ferric ions produced on the oxidized surface of the chalcopyrite (Equation 3) are reduced simultaneously to ferrous ions. In this model, ferrous and ferric ions adsorbed on the surface of the chalcopyrite are the mediators in the transfer of electrons from the chalcopyrite to the dissolved oxygen, [25].

The effects of ferrous ions in sulfuric acid solutions with ferric ions in the absence of dissolved oxygen were studied, and it was concluded that in the presence of cupric ions, the oxidation of chalcopyrite was improved due to the high concentration of ferrous ions, [25].

It is proposed that cupric chloride is reduced much faster than ferric chloride on the surface of chalcopyrite; cuprous chloride is oxidized faster than ferrous chloride, [26]. In general, the Cu\(^{2+}/\text{Cu}^+\) couple is faster and more reversible than the Fe\(^{3+}/\text{Fe}^{2+}\) couple in the corrosion of chalcopyrite, because the mixed voltage, the corrosion current, and the oxidation rate are higher in the presence of cupric ions than of ferric ions.

These improvements in the leaching rate of chalcopyrite at low voltages (below a critical voltage of around 550 mV/SHE) are possible only when ferrous and cupric ions coexist in the solution, due to a synergistic effect between these two ions, [22].

3.2.3. Generation of hydrogen sulfide

It is proposed that chalcopyrite dissolution can occur in the absence of any reagent, in the formation of cupric ions and soluble sulfur species, which presumably is H\(_2\)S, [27]. Studies of the anodic dissolution of chalcopyrite in deoxygenated sulfuric acid solutions, free of ferrous ions, at pH 1.7 and 70 °C were performed, varying the applied voltage over the 550 to 900 mV/SHE, [28]. The authors concluded that for voltages below 700 mV/SHE, there is an imbalance between the total electronic charge (Q\(_e\)) passed to the solution and the ionic charge (Q\(_i\)) associated with the Cu\(^{2+}, \text{Fe}^{3+}\) and Fe\(^{3+}\) ions released into the solution, so that Q\(_e\) < Q\(_i\). This imbalance of positive and negative charges indicates that during the dissolving at low potentials, there is some additional positive charge that is being incorporated to the chalcopyrite structure. These results are interpreted as a mechanism that involves a reaction with protons with the formation of an intermediate polysulfide compound. In this case chalcopyrite dissolution, can be represented by Equation 5.

\[
\text{CuFeS}_2 + 2 \text{H}^+ \rightarrow \text{Cu}^{2+} + \text{Fe}^{3+} + 2 \text{HS}^- + 2 \text{e}^- \quad (5)
\]

In this reaction the chalcopyrite is dissolved anodically as it reacts with the protons, which remove the sulfide ions from the crystal lattice of chalcopyrite, first as HS\(^{-}\) and later, as it continues to react with H\(^+\), in the form of H\(_2\)S\(^{+}\), which can remain on the surface of the chalcopyrite, giving rise to a wide range of polysulfides, [29]. This passive layer can hinder the diffusion of the protons through that layer, slowing or preventing the formation of H\(_2\)S.

The oxidation of chalcopyrite takes place through a progressive ore reduction leading to covellite. It is proposed that the oxidation of chalcopyrite forms a nonstoichiometric copper and iron sulfide, and the elemental sulfur is also transformed progressively from the ore, initially being released as HS\(^{-}\) from the chalcopyrite (Equation 5), becoming the precursor of the passivating species on the solid surface. This proposal is based on the fact that in the preliminary tests with limited oxygen, without the injection of air that can carry the H\(_2\)S, the small amount formed was transformed and deposited on the surface of the ore. Fig. 4 shows the SEM micrograph and the corresponding X-ray diffractogram and fluorescence of the leaching residue of experiment 1-O2N2 (molar concentrations: 0.50 Fe\(^{2+}/0.01 \text{Cu}^{2+}/1.5 \text{Cl}^-\), pH 1, with dissolved oxygen and no added sulphates, after 400 h of leaching. A surface with diminished amounts of sulfur and copper is observed, showing that there is more copper is dissolved and that there is a progressive transformation of the sulfur on the surface of the ore. In this media with no added sulphates, akaganeite [Fe\(_8\)(O,OH)\(_8\)Cl\(_{16}\)], iron and copper chlorides, and iron and copper sulphates are formed. It should be stressed that the associated sulphate originated from the sulfur contained in the chalcopyrite. At the beginning of the leaching experiments, the solution contained dissolved oxygen that was slowly consumed as the leaching proceeded. Since the reactors were hermetically closed in order to avoid the loss of the generated H\(_2\)S, the sulfur reacted with the iron and copper in solution.

To detect the formation of hydrogen sulfide, which is reported to be the possible precursor of passivating species, [23], an indirect method of determination was used. Air was injected into the solution throughout the aerobic leaching experiments (1-O2 to 6-O2), which carried the H\(_2\)S generated into a solution of 0.05 M AgNO\(_3\), causing the precipitation of silver sulfide, see Figs 2 and 3, Equation 1 (reactor) and Equation 6 (sample).

\[
\text{H}_2\text{S} + 2 \text{AgNO}_3 \rightarrow \text{Ag}_2\text{S}(s) + 2 \text{HNO}_3 \quad (6)
\]

Of all the hydrogen sulfide formed during the acidic leaching of chalcopyrite, it is assumed that part remains in the solution and the other part leaves the solution as a gas. For this reason, the implemented setup is meant to confirm, rather than quantify, the formation of hydrogen sulfide.

In the tests with dissolved oxygen (air flow), when the H\(_2\)S leaving the reactor is precipitated by the silver nitrate (equation 6) and the precipitates are analyzed, the presence of silver sulfide is confirmed (see Fig. 3). Furthermore, the solution is acidified.

As the H\(_2\)S produced is carried over, it cannot be involved in the formation of products, although the small dissolved part may cause the thin layer to be formed. It should be mentioned that the H\(_2\)S has been reported in sulphated acidic solutions at 70 °C, [23], without any certainty of whether that formation is due to the sulfur from the chalcopyrite or from the added sulphate; its formation at ambient temperature and in chlorinated...
solutions without the addition of sulphates has not been reported, as in the case of the experimental results presented in the present work.

The tests with dissolved oxygen produced a more quantitative reaction of the chalcopyrite. However, on the surface there was no detectable product layer. Although the dissolved oxygen, upon reacting with the chalcopyrite, formed hydrogen sulfide (H₂S), external factors did not allow a product layer to be formed on the solid surface, either because it was carried away by the air or due to its oxidation with the dissolved oxygen.

In an acidic chloride medium without added sulphates, in the presence of ferrous/cupric ions (0.05 M Fe²⁺ and 0.01 M Cu²⁺), there is a change in the product layer formed on the surface, different from that formed in an environment with added sulphates. This probably promotes an increase in the copper dissolution kinetics of copper. The following sequence of reactions is proposed:

First, chalcopyrite reacts slowly with the hydrogen ion present in the acidic solution to form H₂S (evidenced by the image in Fig. 3), equation 1. Simultaneously, chalcopyrite interacts with cupric and ferrous ions, reducing the former and oxidizing the latter to form chalcocite (equation 2).

In the presence of ferric ion, the sulfide of chalcocite is readily oxidized by ferric ion or simply oxygen to form elemental sulfur, releasing the copper as cuprous ions.

\[
\begin{align*}
\text{Cu}_2\text{S} + 2 \text{Fe}^{3+} + 2 \text{Cl}^- & \rightarrow 2 \text{CuCl} + \text{S} + 2 \text{Fe}^{2+} \quad (7) \\
2 \text{Cu}_2\text{S} + 4 \text{H}^+ + \text{O}_2 + 4 \text{Cl}^- & \rightarrow 4 \text{CuCl} + 2 \text{S} + 2 \text{H}_2\text{O} \quad (8)
\end{align*}
\]

Assuming that the reaction of Equation 1 is favored, a non-oxidative/oxidative model is proposed in which it is established that the addition of ferrous ions improves the dissolving rate when cupric ions are present. This model is similar to that proposed by Hiroyoshi et al., 2004, [22], but applied here for chloride media.

### 3.2.4. Solutions with added sulphates

The SEM micrographs of the residue selected from each experimental condition (for tests both with and without oxygen) in all the samples consisted of primarily unreacted chalcopyrite and small amounts of product layer. The exceptions are tests 4S-O2 (molar concentrations: 0.05 Fe²⁺/0.01 Cu²⁺/1.5 Cl⁻ with dissolved oxygen) and 4S-N2 (molar concentrations: 0.05 Fe²⁺/0.01 Cu²⁺/1.5 Cl⁻, without dissolved oxygen), in a sulphated environment, where a product layer on the surface of the chalcopyrite can be distinctly observed (Fig. 5). After 400 h of leaching, the surface of the chalcopyrite in the solution without oxygen (4S-N2), there is a nonstoichiometric formation of sulfur and chalcopyrite crystals, while the aerobic tests show the formation of jarosite (4S-O2). The amount of jarosite produced decreases when the acidity is increased in the chloride, as well as sulphate medium, [30]. Equation 9 shows a precipitation reaction typical of a jarosite. It is believed that this reaction is not reversible in systems with chalcopyrite, since it is an insoluble product, [31].

\[
\text{Na}^+ + 3 \text{Fe}^{3+} + 2 \text{SO}_4^{2-} + 6 \text{H}_2\text{O} \leftrightarrow \text{NaFe}_3(\text{SO}_4)_2(\text{OH})_6 + 6 \text{H}^+ \quad (9)
\]

If the formation of jarosite causes the passivation on chalcopyrite, it can be assumed that pyrite would show a similar behavior. However, it is well known that pyrite does not undergo passivation. It can be seen that the formation of a product layer on the surface of the ore is not homogeneous, and this can be explained by the fact that on the surface of the chalcopyrite, Cu³⁺Fe³⁺(S₂)₂, on some plane orientations, the Cu is being oxidized and the Fe is being reduced to Fe²⁺, while the S can be easily oxidized or reduced; on the surface of the ore there are FeS₂ type compounds as the distances between the Fe and Cu atoms are rearranged; in other orientations there are significant changes in the atomic arrangement of S, Fe and Cu, [32]. Another possible explanation of the nonhomogeneous formation of the product layer on the surface of the ore is its low adherence.

### 4. Conclusions

From the chlorinated acid leaching in the presence of ferrous/cupric/ferric ions without the addition of sulphates, the following conclusions can be reached:

The presence of dissolved oxygen in an acid chloride environment is important for the leaching reaction of chalcopyrite to take place in a slightly oxidizing medium. Oxygen is a catalyst for a more quantitative oxidation of the ferrous ions on the surface of chalcopyrite, causing more release of ferrous and cuprous ions from the surface of the chalcopyrite. Therefore, there is a greater dissolution of copper in solutions with dissolved oxygen, suggesting an increase of the kinetics of the leaching process, improved the copper extraction and promotion of H₂S formation.

The presence of ferrous ions coexisting with cupric ions was beneficial to increase the dissolving kinetics of copper; the oxidation of chalcopyrite was improved due to the elevated concentrations of ferrous ions (0.05 M), at 560 mV/SHE. The extraction of copper was controlled by the ferrous/ferric concentration ratio or the redox potential of the solution. Ferrous ions suppress the oxidation of chalcopyrite when the cupric ion concentration is low.

In a chlorinated medium without the addition of sulphates with dissolved oxygen, there was generation of H₂S, which is the precursor of passivating species of the surface of chalcopyrite. However, on the surface of the samples there was no detectable product layer because the hydrogen sulfide (H₂S) did not have time to form species either because it was entrained by the air stream or it reacted with the dissolved oxygen in the solution. H₂S formation at ambient temperature and in chlorinated solutions without the addition of sulphates, as in the case
of the experimental tests of the present work, has not been previously reported.

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6. References

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