Study of the dissolution of chalcopyrite in solutions of different ammonium salts

by T. Moyo* and J. Petersen*

Synopsis

The oxidative leaching of chalcopyrite in ammoniacal solutions has been evaluated using electro-analytical techniques and controlled bulk leaching studies. The anodic dissolution process has been established to be a seven-electron transfer process under nitrogen in ammonia-ammonium sulphate solutions and ammonia-ammonium carbonate solutions, which suggests that the sulphur is oxidized to thiosulphate and the copper and iron released as Cu⁺ and Fe²⁺ in these systems. The departure of Fe²⁺ and S₄O₆²⁻ is affected by choice of the ammonium salt used in the leaching process. In the perchlorate salt, only five electrons are transferred, supporting formation of different sulphur species as proposed for the sulphate and carbonate salts. Scanning electron microscopy and energy-dispersive spectroscopic analysis of the mineral surface after leaching indicate presence of an iron–sulphur surface layer completely free of copper in the sulphate system, an iron-rich surface layer in the perchlorate solutions, and absence of surface layer build-up in carbonate solutions. XRD analysis of a bulk leach residue from leaching in ammonia-ammonium sulphate solutions showed the surface layer to be mostly amorphous (90%). The crystalline content (10%) is composed of 95% polymorphs of anhydrous iron oxide hydroxide FeO(OH). Choice of ammonium salt and the hydrodynamic environment of leaching have been shown to influence the presence or absence of the surface product, as well as its nature.

Keywords
chalcopyrite, ammonia, surface deposits, coulometry, iron oxy-hydroxide.

Introduction

Hydrometallurgical processes for copper extraction provide a viable route to recover the metal from mixed and low-grade ores, as well as overcome the environmental challenges faced by the traditional pyrometallurgical processes. Chalcopyrite is not only the most abundant of the copper sulphides, but also the most stable, making it recalcitrant to hydrometallurgical processes. Hence, hydrometallurgical processing of chalcopyrite continues to be an attractive area of research due to the vaguely understood surface chemistry of the mineral in different aqueous media. Different routes for hydrometallurgical treatment of chalcopyrite can be followed. These include thermal treatment prior to leaching, direct leaching and direct electrochemical leaching (Venkatachalam, 1991). Direct leaching of chalcopyrite can be carried out in various solution systems, as reviewed by Roman and Benner (1973) and Venkatachalam (1991). Ammoniacal solutions are attractive and effective lixiviants that form stable amine complexes with some base metal cations while rejecting iron. Leaching of chalcopyrite in ammoniacal solutions in the presence of an oxidant is possible due to the stabilization of copper(I) and copper(II) by ammonia at elevated pH levels. In oxygenated ammonia solutions, it has been suggested that chalcopyrite dissolves according to Equation [1] (Beckstead and Miller., 1977a):

\[ \text{CuFeS}_2 + 4\text{NH}_3 + 17/4\text{O}_2 + 2\text{OH}^- \rightarrow \text{Cu(NH}_3\text{)}^{2+} + \frac{1}{2}\text{Fe}_2\text{O}_3 + 2\text{SO}_3^{2-} + \text{H}_2\text{O} \]  

Chalcopyrite is characterized by very slow leaching kinetics and this has been strongly linked to the formation of a passive film on its surface. Researchers have not reached a consensus on the actual composition or degree of stability of this passive surface film. Fe₂O₃ or its hydrated form Fe₂O₃.xH₂O (Beckstead and Miller, 1977b; Yin et al., 1995; Forward and Mackiw, 1955; Feng and Van Deventer, 2002) and Fe(OH)₃ (Yin et al., 1995; Warren and Wadsworth, 1984) have been reported to be the surface products of chalcopyrite oxidation. Warren and Wadsworth (1984) studied the electrochemical oxidation of chalcopyrite in ammonia–ammonium sulphate solutions and reported significant amounts of ferrous iron (5–20%) in the product film. The authors postulated that a ferrous iron intermediate was formed, which can be readily oxidized to ferric, and the oxidation can be achieved even if only traces of dissolved oxygen are present. If ferrous iron is formed from chalcopyrite dissolution, it is expected that its further reactions would be affected by the solution conditions, hence the different surface effects as observed under the

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carbonate and sulphate ammonium salts. Asselin (2011) reported that Fe(II) ammines are thermodynamically stable only under reducing conditions and concluded that they were unlikely to be formed if oxygen is present. He presented quasi-equilibrium Pourbaix diagrams for the Fe–NH$_2$–H$_2$O system. According to the diagram, Fe(OH)$_3$ is the species present at noble potentials across all pH ranges, while Fe(OH)$_2$ is present only above pH 11.

Forward and Mackiw (1955) suggested that sulphide minerals, chalcopyrite in this case, react with oxygen, water, and ammonia to produce soluble salts in such a way that the iron present in each mineral particle is converted to hydrated iron oxide in situ, with the result that the particles, when leaching is complete, consist of hydrated iron oxide pseudomorphic with the original mineral. Beckstead and Miller (1977b) discuss the nucleation and growth of the haematite passivating layer (in ammonia–ammonium sulphate solutions), stating that this occurred on the anodic sites. Thus, limiting the overall reaction by limiting the anodic reaction. Agitation at speeds up to 3000 r/min increased the rate of reaction, possibly by abrading the haematite layer and exposing more of the anodic sites. The contradiction between these two studies stems from the fact that Beckstead and Miller looked at the nucleation and growth of the Fe product and identified it as haematite, while Forward and Mackiw looked at ‘an ion substitution mechanism, resulting in a Fe product atypical of chalcopyrite’. In a study on the surface oxidation of chalcopyrite in alkaline solutions, Yin et al. (2000) reported that the iron in the top layer of chalcopyrite oxidized, forming a monolayer of Fe(OH)$_3$ and Fe$_3$O$_4$, while the copper and sulphur remained unoxidized in the original chalcopyrite crystal structure, forming a phase they designated as CuS$_2^*$, which together with the Fe(OH)$_3$ and Fe$_3$O$_4$ retarded the oxidation of the mineral. The abrasion of the passivating layer, although not widely researched in ammoniacal media, has been demonstrated to have potential to increase copper recoveries and a process, the FLSmidth® Rapid Oxidative Leach (ROL), has been patented (Eyzaquiirre et al., 2015). This process uses very low-energy interstage attrition to enhance dissolution of copper-bearing minerals, primarily chalcopyrite, through a mechano-chemical process that is based on the removal of the passivating product layers. The application of such a process agrees well with results from Beckstead and Miller’s (1977b), in which the chalcopyrite surface was observed to be free of surface deposit at high agitation speeds, possibly due to increased abrasion.

Literature on iron chemistry in ammonia–ammonium carbonate solutions is limited for the chalcopyrite leaching process, thus reference will be made to the Caron process. Studies have been carried out on the leaching of nickel from pre-reduced laterite ores (nickel and iron are present as reduced metallic grains and metal alloys) (Das and Anand, 1995; Jandová and Pedlík, 1994, Kim et al., 1991; Lee, Osseo-Asare, and Pickering, 1985; Nikoloski, 2002; Nicol, Nikoloski, and Fittock, 2004). At a pH of about 9.8 in ammoniacal solutions, the dominant dissolved iron species is the ferrous tetra-ammine ion and the dissolution reaction has been postulated to occur according to Equation [2] (Nikoloski, 2002; D’Aloya and Nikoloski, 2012; Subrata, 2010; Nikoloski and Nicol, 2006; Osseo-Asare and Ashihene, 1979):

\[
\text{Fe} + 4 \text{NH}_3 \rightarrow \text{Fe(NH}_3)_2^2+ + 2e
\]  
[2]

Fe$_n$Ni$_m + 8\text{NH}_3 \rightarrow \text{Fe(NH}_3)_2^{2+} + \text{Ni(NH}_3)_2^{2+}
\]  
[3]

The ferrous ammine is oxidized in aerated solutions to form ferric, which does not form ammines and is thus precipitated as Fe(OH)$_3$ releasing the ammonia:

\[
\text{Fe(NH}_3)_2^{2+} + 0.25 \text{O}_2 + 2\text{OH}^- + 0.5 \text{H}_2\text{O} \rightarrow \text{Fe(OH)}_3 + 2\text{NH}_3
\]  
[4]

Nikoloski (2002) showed that iron is prone to passivation in solutions typical of those used in practice, and confirmed its occurrence in leaching reactors of a commercial scale in the Caron process. Passivation was shown to occur due to the formation of an oxide layer at the potentials that can be attained in the presence of high concentrations of dissolved oxygen.

An earlier study by the authors (Moyo et al., 2015) investigated chalcopyrite dissolution in ammonia–ammonium sulphate solutions. Coulometric measurements on a chalcopyrite electrode resulted in postulating a seven-electron transfer mechanism, with the potential formation of thiosulphate as the product of sulphur oxidation. Copper(II), rather than oxygen, was found to be the primary oxidant on the mineral surface. Rapid formation of an iron oxide layer on the electrode surface was observed, which did not inhibit, but slowed down, the rate of anodic dissolution over time. No iron was found in solution, whereas no copper was found in the surface layer.

The present study aims to elucidate the nature of the surface deposits and to what extent anions in the ammoniacal solution might have an effect on their formation. Tests were done in ammonium sulphate, carbonate, and perchlorate solutions. Coulometric tests were done in each case to ascertain if the dissolution mechanism remained the same in the different solutions before investigating the surface layers formed on the chalcopyrite electrode.

Experimental

Chalcopyrite electrodes

Samples of natural chalcopyrite from Durango, Mexico (supplied by Wards Natural Science) were mounted on brass stubs with conductive silver epoxy and the assembly then imbedded in non-conductive epoxy resin. X-ray diffraction (XRD) analysis of the mineral sample indicated it to be 100% chalcopyrite, while quantitative electron microscopy (QEMSCAN™) indicated it to be 95% chalcopyrite, with 3% sphalerite and 1% calcite. The electrode surface was polished on 1200-grit SiC abrasive paper, then on 1 µm, 0.3 µm, and 0.05 µm aluminium oxide prior to each experiment, except where stated. Only one electrode was used for the purposes of this study and the exposed surface area of the electrode was measured to be 0.36 cm$^2$ using imageJ™ for image analysis. The surface area was measured intermittently during the course of the study and was found not to change significantly.

Electrochemical measurements

A standard three-electrode cell with a chalcopyrite rotating-disk working electrode was used for all electrochemical
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measurements. A saturated calomel reference electrode and platinum wire auxiliary electrode were used. A freshly polished chalcopyrite electrode was placed into an electrolyte of desired composition and rotated at 1600 r/min. Rest potentials were measured for 30 minutes. A Gamry Series G 300/750 potentiostat was used. Coulometric experiments were carried out by fixing the potential at a value measured during the rest potential measurements described above (after 30 minutes) in a solution of similar composition but in the absence of copper(II) ions. Total charge was obtained by integration of the current–time curves. The current response of the electrode was then measured for 2 hours. Additional coulometric experiments were carried out for 5 hours or 22 hours at the rest potential or the vicinity of the rest potential in a minimum volume of solution. At the end of the experiment, the solutions were analysed for copper using inductively coupled plasma optical emission spectroscopy (ICP-OES). All potentials are reported against the standard hydrogen electrode (SHE) unless stated otherwise. Experiments were conducted at 25°C in a thermostatted cell containing 50 mL solution. The total Cu concentration in solution at the end of the coulometric experiments was established by ICP-OES.

Controlled leaching experiments

Three tests were carried out, two in which blocks of chalcopyrite 5 mm × 5 mm × 5 mm were leached (reactors A and B) and a third one in which micronized chalcopyrite (81% passing -75 μm) was leached in 3 M total ammonia-ammonium sulphate solutions at 1% solids (reactor C). Reactor A had glass beads added, which allowed for the abrasion of the surface product from the chalcopyrite blocks, while reactor B had no beads in it. The leaching experiments went on for a period of 5 days, allowing for significant amounts of surface product to be generated. Magnetic stirrers were used; this was adequate to keep the sample in swirling motion but could not suspend the sample in solution for motion but could not suspend the sample in solution for

samples were taken intermittently and analysed for copper using atomic absorption spectrophotometry (AAS).

Solution preparation

Deionized water and reagent-grade CuSO₄·5H₂O, NH₃OH, (NH₄)₂SO₄, (NH₄)₂CO₃, NH₄ClO₄, H₂SO₄, and NaOH were used. All solutions were prepared by mixing 1:1 molar ratio ammonium hydroxide to ammonium salt based on the ammonium ion, i.e., a 1 M (NH₄⁺,NH₃) solution was made by mixing 1 M NH₃OH with 0.5 M (NH₄)₂SO₄. Rest potentials that were later used for the coulometric experiments, were measured in the presence of 5 g/L initial copper(II). Either oxygen (99.99%) or nitrogen was bubbled into the electrolyte for 10 minutes prior to starting the experiment, and bubbling was continued throughout the experiment, taking care to ensure that gas bubbles did not accumulate on the chalcopyrite electrode surface (in electrochemical experiments). This was established to be sufficient time to either saturate the solution with dissolved oxygen or remove it from solution.

Surface measurements

The surface of the chalcopyrite was examined by scanning electron microscopy (SEM) and QEMSCAN. An elemental analysis was done on the surface using energy-dispersive X-ray spectroscopy (EDS) in an attempt to characterize the surface effects and products under the conditions of this study. The leach residue was further analysed using X-ray diffraction (XRD), Brunauer-Emmett-Teller (BET) surface analysis, as well as digested and subjected to chemical analysis using atomic absorption spectroscopy.

Results and discussion

Coulometry

Coulometry studies were done in the different salt media, as described in the methodology. The stoichiometry of the anodic dissolution reaction was determined by this method. The coulometric results shown in Table 1 include data from our previous study (Moyo et al., 2015) on the sulphate system. Table 1 shows the number of electrons transferred per molecule of chalcopyrite (measured as copper released into solution) at potentials corresponding to rest potentials measured in 1 M (NH₄⁺,NH₃) with 5 g/L Cu(II) at 25°C, but here in the absence of initial Cu in solution. All measurements were repeated at least three times and the calculated number of electrons transferred was found not to vary by more than 5%.

<table>
<thead>
<tr>
<th>Ammonium salt</th>
<th>E₀ (V vs SHE)</th>
<th>Total Cu from ICP (mol/L)</th>
<th>Total charge Q (C)</th>
<th>n(e)=Q/F</th>
<th>No of electrons/mole Cu</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbonate N₂</td>
<td>0.231</td>
<td>9.47E-07</td>
<td>0.635</td>
<td>6.58E-06</td>
<td>7</td>
</tr>
<tr>
<td>Sulphate* N₂</td>
<td>0.26</td>
<td>3.96E-07</td>
<td>0.261</td>
<td>2.71E-06</td>
<td>6.8</td>
</tr>
<tr>
<td>Perchlorate N₂</td>
<td>0.013</td>
<td>1.01E-06</td>
<td>0.473</td>
<td>4.90E-06</td>
<td>4.8</td>
</tr>
</tbody>
</table>

*Data from Moyo et al. (2015)
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In the carbonate system, as previously found for the sulphate system, approximately seven electrons are transferred per mole of copper. This suggests the formation of a thiosulphate intermediate and that the copper and iron are in the cuprous and ferrous states. The cuprous and ferrous ions would then be subsequently oxidized in solution, in non-faradaic reactions, to the cupric and ferric state. The stoichiometry of the reaction would be approximated by Equation [5]. In perchlorate solution, on the other hand, non-faradaic reactions, to the cupric and ferric state. The cuprous and ferrous take place as faradaic reactions:

\[
\begin{align*}
\text{CuFeS}_2 + 4 \text{NH}_3 + 6 \text{OH}^- & \rightarrow \text{Cu(NH}_3)_2^{+} + \\
\text{Fe}^{2+} + \text{S}_2 \text{O}_3^{2-} + 3 \text{H}_2\text{O} & + 7\text{e}^{-}
\end{align*}
\]

[5]

\[
\begin{align*}
\text{CuFeS}_2 + 4 \text{NH}_3 + 3 \text{OH}^- & \rightarrow \\
\text{Cu}^{2+} + \text{Fe(OH)}_3^{3+} + 2 \text{S} + 5\text{e}^{-}
\end{align*}
\]

[6]

**Topological effects**

The visual differences between the chalcopyrite surfaces after polarization were apparent even to the naked eye, with those treated in the sulphate and perchlorate solutions having lost their lustre and exhibiting a reddish-brown colouration, while that treated in the carbonate solution had lost its lustre but did not exhibit any colour changes. Figure 1A shows optical microscope images of a freshly polished chalcopyrite surface and 1B shows chalcopyrite after oxidation in ammonia–ammonium sulphate solutions – the difference is quite clear. Freshly polished chalcopyrite prior to leaching was analysed by Warren and Wadsworth (1984), who upon scraping off and dissolving the surface film from their experiments identified some orthorhombic sulphur, confirming that elemental sulphur was present in both the surface film and leachate. The researchers attributed this to the disproportionation of some of the intermediate sulphur species expected to be found in the leachate. Kuhn, Arbiter, and Kling (1974) presented the possibility of the recrystallization of sulphur from the sulphide to form rhombic or monoclinic sulphur, which would be expected to be found in localized sites within the mineral, while cations diffuse out of the mineral. Therefore, the presence of sulphur on the mineral surface, as observed in this study, is possible although not widely reported.

The morphology of the surface layer suggests that it may have been formed by precipitation, which is in contradiction to the ion-substitution mechanism proposed by Forward and Mackiw (1955). This is more apparent when looking at residue on the surface of a chalcopyrite block leached for 5 days in the absence of glass beads (reactor B) (Figure 5), which shows what appears to be an agglomerate of small particles. This surface product readily dissolved upon short contact with concentrated sulphuric acid solutions, leaving behind an etched but pure chalcopyrite surface. The acid solution used to wash off the surface layer was analysed using ICP and found to contain 16.05 mg/L iron but only 0.095 mg/L copper, confirming the SEM results which indicated it to be an iron-rich surface layer.

As stated in the introduction, the presence of a relatively small amount of sulphur in the surface deposit prompted the investigation of chalcopyrite dissolution in non-sulphate media. Figure 3 shows the mineral surface after 22 hours of oxidation in ammonia–ammonium carbonate solution. All other solution conditions were maintained the same. The bulk EDS results indicates 23.78% Fe, 44.12% S, 20.82% Cu, 25.67% Fe and 49.71% S, measured by bulk EDS, are consistent with those of pure chalcopyrite. This provides a basis of comparison for the samples analysed post-leaching. Figure 2 shows a SEM image of a chalcopyrite surface after 22 hours of oxidation in ammonia–ammonium sulphate solutions under nitrogen. It is copper-free and has significant shift from the 1:2 ratio of pure chalcopyrite. The Fe:S ratio is 1.9:1, which represents a significant shift from the 1:2 ratio of pure chalcopyrite. The presence of sulphur on the surface was previously reported by Warren and Wadsworth (1984), who upon scraping off and dissolving the surface film from their experiments identified some orthorhombic sulphur, confirming that elemental sulphur was present in both the surface film and leachate.
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Figure 3—SEM image of chalcopyrite immediately after leaching in ammonia–ammonium carbonate solution. Bulk EDS (in atomic per cent) 23.78% Fe, 44.12% S, 20.82% Cu, 11.28% O, and a spot EDS within the grain boundary 26.19% Fe, 10.51% S, 3.87% Cu, 58.67% O, 0.8% Si

would be present. The visible cracks were observed to be more copper-depleted. EDS analysis of the crack areas showed only 4–6% to be copper, suggesting these to be areas of preferential chalcopyrite dissolution in the carbonate medium. This is further supported by the visibility of polishing marks on the unleached areas in Figure 3. The differences in surface effects between the sulphate and carbonate solutions suggest that the surface reaction mechanism varies, although the same number of electrons is transferred. Iron carbonate complexes potentially form in this system, and tend to stay in solution, whereas the formation of a surface precipitate in the sulphate system is more likely to be linked to the formation of iron sulphate complexes that precipitate.

The behaviour of iron in ammonium carbonate solutions has been explored by some researchers, with the focus on improving the leaching kinetics in the Caron process. Kim et al. (1991) studied the active-passive behaviour of sintered iron in ammoniacal ammonium carbonate solutions at pH 9.7 and reported that surface films were formed on the bulk iron during exposure to air or immersion in ammoniacal solutions. The authors characterized the surface films using X-ray photoelectron spectroscopy (XPS) and generated cathodic reactivation transients; they reported that in the presence of air Fe$_3$O$_4$ was formed. Caldeira, Ciminelli, and Osseo-Asare (2008) investigated the effect of carbonate on pyrite oxidation in alkane solutions and identified (using diffuse reflectance infrared spectroscopy) iron carbonate compounds as one of the products of pyrite oxidation. The authors explain that the increased oxidation rate typically observed on pyrite in carbonate solutions is possibly due to the formation of Fe(II)–CO$_3$ complexes, the buffering effect of the carbonate and the fact that complexation with bicarbonate/carbonate provides a stronger Fe(III)/Fe(II) redox couple, increasing the Fe(III) solubility. This could be an explanation for the apparent lack of significant surface layer formation on chalcopyrite in the carbonate system, through the possible formation of soluble Fe(II)–CO$_3$ complexes which are then transported away from the vicinity of the mineral surface. It should be noted that, although other studies of the carbonate system reported the presence of an oxide layer, the obvious lack of any significant layer in this study could also be attributed to the hydrodynamics around the electrode area. Rotating at 1600 r/min may be sufficient to allow the transport of the metastable iron carbonates away from the mineral surface before they have a chance to nucleate and precipitate. The effect of electrode rotation speed on the extent of surface layer build-up and its morphology in the sulphate system have been reported by Beckstead and Miller (1977b), who found that chalcopyrite surfaces were much cleaner and relatively free of product at agitation speeds higher than 1100 r/min. They postulated that increased agitation speeds improved leaching by increasing turbulence, which in turn allowed for the abrasion of the surface product, thereby exposing a fresh mineral surface for leaching.

To provide further assessment, especially with regard to the sulphur deportment, tests were done in ammonia–ammonium perchlorate solutions, as the perchlorate ions are not expected to complex with the oxidation products of chalcopyrite.

Figure 4 shows an SEM image of a chalcopyrite sample after 22 hours of leaching in ammonia–ammonium perchlorate solutions. Visually, the surface appears not dissimilar to that observed in the sulphate system. However, the bulk surface EDS gave an Fe:S atomic ratio of 1:1.1 and copper is present at a S:Cu of 1:0.38. Analysis of isolated flaky areas on the sample showed these areas to be more iron-rich: Fe:S is 2:1 whereas the S:Cu remains at 1:0.38. The results suggest the presence of an iron-rich surface together with a sulphur species. The atomic ratios of the sulphur and iron are not consistent with the stoichiometric ratios suggested by coulometry (Equation [6]), which would require all sulphur to accumulate on the surface as elemental sulphur and hence there should be twice as much sulphur as iron. However, tests done so far cannot be used to make any conclusive deductions as to whether the sulphur observed to be present on the mineral surface is elemental sulphur or some form of Fe–S complex. Furthermore, the fact that copper

Figure 4—SEM image of chalcopyrite immediately after leaching in ammonia–ammonium perchlorate solution. Bulk EDS (in atomic per cent) 16.25% Fe, 17.33% S, 6.77% Cu, 53.92% O, 4.03% Si. Analysis of the highlighted area 25.56% Fe, 13.08% S, 5.11% Cu, and 56.24% O
was retained on the surface could have affected the copper balance during the coulometric study of the perchlorate system, thus leading to an under-estimate of electrons transferred.

Experiments were done in which 5 mm blocks of the same sample of chalcopyrite were leached in ammonium sulphate, as described in the methodology. In both instances, the remaining pieces of the blocks, as well as any residue that accumulated in the reactor at the end of the experiment, were washed in distilled water, dried, and analysed under the microscope and by QEMSCAN. Figure 5 shows a SEM image of the residue collected from the reactor that contained the beads. Neither sulphur nor copper were found on the residue, and EDS of the residue gave a ratio of O:Fe 1.1:1, fairly close to the 1.5:1 ratio typical of haematite, although this is unlikely to form under the present experimental conditions.

XRD analysis of the residue indicated it to be 90% amorphous, and of the 10% crystalline content 95% comprised polymorphs of anhydrous iron oxide hydroxide FeO(OH) as shown in Table II. ICP and LECO analysis indicated the residue to contain 44% iron and less than 1% of FeO(OH) as shown in Table II. ICP and LECO analysis showed that a ratio of O:Fe 1.5:1, fairly close to the 1.5:1 ratio typical of haematite, although this is unlikely to form under the present experimental conditions.

It should be noted that the leach reaction proceeded much faster in the reactor containing glass beads, with copper recoveries at the end of the 5-day leaching period at 54.9%, whereas in the reactor without glass beads merely 14.7% was leached. It is apparent that the continuous removal of the surface deposits in the reactor with beads had a positive impact on the recoveries. This is in agreement with Beckstead and Miller (1977b), who reported that at low stirring speeds the rate of the reaction was significantly reduced: they related this to the growth and nucleation of a haematite phase. They also reported on the possible impact of stirring speed on the morphology of the haematite phase they observed. In the development of the Sherritt–Gordon process, critical agitation speeds were reported (Forward and Mackiw, 1955) and these are said to have allowed for the abrading of the ‘haematite’ layer, thus improving leaching rates. Also, in the Arbiter process (Kuhn et al., 1974), intense mixing is used to abrade the haematite phase from the Cu–Fe–S phase, thus exposing a fresh surface for reaction, as well as achieving good oxygen transfer rates.

Figure 6 shows QEMSCAN and SEM images of one of the sample blocks placed in the reactor without beads. An iron-rich surface deposit is visible at the edge on the mineral, and the morphology of this surface layer can be seen on the BSE image to be similar to that of the debris in Figure 5. The bulk phase, orange in colour, has been identified as chalcopyrite, while the yellow is an iron-rich phase containing small percentages of sulphur and the black phase is an iron- and oxygen-rich phase without any sulphur. An EDS analysis of the surface layer showed it to be 57.7% Fe, 32.12% O, 1.45% S, and the rest reported to be silica and calcium.

Leach residue from reactor C (finely ground material) was washed, dried, and analysed. Figure 7 shows a QEMSCAN and SEM image of part of the debris. It is apparent that an iron-sulphur-based layer formed around a chalcopyrite core. The layer also contained iron which is not associated with

**Table II**

XRD results of the crystalline phases found in the chalcopyrite debris recovered from reactor B

<table>
<thead>
<tr>
<th>Phase</th>
<th>Composition (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chalcopyrite (CuFeS₂)</td>
<td>4.89</td>
</tr>
<tr>
<td>Quartz (SiO₂)</td>
<td>0.48</td>
</tr>
<tr>
<td>Geothite (α-FeO(OH))</td>
<td>5.8</td>
</tr>
<tr>
<td>Lepidocrite (γ-FeO(OH))</td>
<td>51.5</td>
</tr>
<tr>
<td>Akaganeite (β-FeO(OH))</td>
<td>37.4</td>
</tr>
</tbody>
</table>

Figure 5—SEM image of chalcopyrite leach residue abraded off the mineral surface by glass beads in a 3 M equimolar ammonia–ammonium sulphate solutions at 25°C under oxygen. Sample taken after 5 days of leaching. EDS of the marked area (in atomic per cent) 45.41% Fe, 51.53% O, 3.06% Si

Figure 6—QEMSCAN and BSE images of a section of the chalcopyrite blocks after leaching in 3 M (NH₄)₂SO₄ at 25°C under oxygen

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sulphur, which was identified as an iron oxy-hydroxide species. It is worth mentioning that all species that contained iron and sulphur in Fe:S ratios higher than 0.5 reported to pyrite and pyrrhotite, but this classification does not in any way mean this phase is in actual fact pyrite or pyrrhotite.

From Figures 5 to 7, the morphology of the surface product appears to be significantly different from that of the initial sample. The layer appears to be composed of an agglomeration of smaller particles as to it be relatively porous compared to chalcopyrite. Surface area and porosity were measured using the Brunauer-Emmett-Teller (BET) method on the micronized sample prior to and after leaching (Table III). It is apparent that the surface area and surface pore volume of leached particles increased significantly relative to that of the non-leached chalcopyrite. It is also notable that the percentage of particles less than 25 μm (diameter) increased significantly in the leach residue samples, from 67% in an unleached sample to about 90% in leached samples. This suggests that although oxidation of chalcopyrite may occur in situ as suggested by Forward and Mackiw (1955), this surface layer is in fact quite friable and would be easily abraded, hence the formation of smaller sized particles, also confirmed by the increased surface area per gram of the samples. These observations are consistent with findings by Guan and Han (1997), who identified a friable and porous surface product while oxidizing chalcopyrite in ammonia-ammonium sulphate and ammonia-ammonium carbonate solutions. The readily soluble nature of the surface layer observed when washing the electrode layer in acid solutions indicates that this surface layer is not stable.

Conclusions
1. The number of electrons (seven) transferred per mole copper during anodic oxidation is similar for the ammonia-ammonium sulphate and ammonia-ammonium carbonate solutions
2. Ammonia-ammonium perchlorate solutions promote a five-electron transfer/copper reaction, possibly forming elemental sulphur on the mineral surface
3. Ammonium sulphate leaching results in the formation of a Fe-oxyhydroxide layer with low sulphur on the mineral surface
4. Ammonia-ammonium carbonate solutions resulted in marginal accumulation of iron on the mineral surface, but no formation of a layer was observed
5. Ammonium perchlorate leaching results in the formation of a Fe-oxyhydroxide layer with moderate sulphur on the mineral surface
6. The surface product was largely amorphous (90%) and significantly more porous (9-12 times) than unleached chalcopyrite. The observed morphology of the surface product suggests that it is formed through secondary precipitation rather than as part of the chalcopyrite dissolution mechanism
7. Surface abrasion allows for the removal of the surface product, leading to improved leaching recoveries
8. The abraded surface product from the small particles leaching experiment contained no sulphur, while surface products found on the stationary block of mineral contained small quantities of sulphur.

Figure 7—QEMSCAN and SEM images of chalcopyrite leach residue after 15 days of leaching in ammonia-ammonium sulphate solutions 3 M (NH₄)₂SO₄, 25°C, pH 9.6±0.15 under oxygen

Table III
Surface analysis results of chalcopyrite prior to and post leaching at 1% solids, pH 9.6±0.15, 25°C in the presence of oxygen in solutions of different total ammonia concentration

<table>
<thead>
<tr>
<th>Sample</th>
<th>Surface area (m²/g)</th>
<th>Pore volume (m³/g)</th>
<th>% passing 25 μm</th>
</tr>
</thead>
<tbody>
<tr>
<td>chalcopyrite</td>
<td>1.275</td>
<td>0.0055</td>
<td>66.61</td>
</tr>
<tr>
<td>1M leached</td>
<td>57.346</td>
<td>0.0642</td>
<td>94.07</td>
</tr>
<tr>
<td>3M leached</td>
<td>57.734</td>
<td>0.0503</td>
<td>89.89</td>
</tr>
<tr>
<td>6M leached</td>
<td>59.526</td>
<td>0.0577</td>
<td>89.52</td>
</tr>
</tbody>
</table>
Study of the dissolution of chalcopyrite in solutions of different ammonium salts

It has thus been shown that a relatively unstable iron product forms on the surface of chalcopyrite through secondary reactions to the faradaic oxidation reaction. This product may contain small percentages of sulphur but, regardless of system, the majority of leached sulphur reports to the solution. Choice of ammonium salt and the hydrodynamic environment of leaching influence the presence or absence, as well as the nature, of the surface product. It appears that the formation of surface products in turn influences the reaction mechanism of chalcopyrite dissolution, and the two aspects need to be studied in conjunction.

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References


