Intermediates Transformation of Bornite Bioleaching by *Leptospirillum ferriphilum* and *Acidithiobacillus caldus*

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**Abstract:** Bioleaching experiments, electrochemical tests, X-ray diffraction (XRD), and X-ray photoelectron spectroscopy (XPS) were conducted to investigate the intermediates transformation of bornite by *Leptospirillum ferriphilum* and *Acidithiobacillus caldus*. The bioleaching experimental results showed that the presence of *L. ferriphilum* and *A. caldus* significantly accelerated the bornite bioleaching. In addition, the intermediate species of bornite bioleaching with these two kinds of bacteria were similar. Electrochemical analysis indicated that the dissolution of bornite was an acid-consuming process. The results of XRD showed that intermediate species, namely covellite (CuS), mooihoekit (Cu$_9$Fe$_9$S$_{16}$) and isocubanite (CuFe$_2$S$_3$), were formed during bornite bioleaching, and a mass of elemental sulfur was formed in the late stage of bioleaching. The Cu 2p photoelectron spectrum revealed that Cu was present in the form of Cu (I) during the bornite bioleaching. Additionally, the S 2p$_{3/2}$ photoelectron spectrum suggested that S$^{2-}$ and S$_2^{2-}$ were gradually converted to S$^0$, and the formation of elemental sulfur hindered the further dissolution of the bornite.

**Keywords:** bornite; *A. caldus*; *L. ferriphilum*; bioleaching; XRD; XPS

1. Introduction

Currently, the continuous consumption of high-grade copper resources is causing the grade of copper ore to decrease rapidly [1]. High energy consumption and environmental problems pose challenges to traditional pyrometallurgy and hydrometallurgy. Bioleaching has been widely used in leaching low grade copper ores and complex ores, with the advantages of a simple process, lower energy consumption, as well as being eco-friendly [2,3]. Nowadays, the main research focus of copper ore bioleaching is chalcopyrite, for the reserves of chalcopyrite are at the top of copper ore resource importance.

Bornite is also an important source of copper, following chalcopyrite and chalcocite in the list of copper resources [4]. Some researchers have studied the traditional hydrometallurgical process of bornite. The dissolution of bornite was divided into two steps in which bornite was leached in acidic ferric sulfate solution [5,6]. First, non-stoichiometric bornite was formed and second converted into chalcopyrite and elemental sulfur. Pesic [4] also had the same conclusion that the dissolution of bornite was divided into two steps during bornite leaching by acidic FeCl$_3$. They suggested that the first step reaction was an electrochemical reaction controlled by a mixing kinetics process. The potential
of the surface reaction was controlled by the Cu$^+$ in bornite crystal lattices and the first step was terminated when the Cu$^+$ in the bornite was consumed. Thereafter, the Fe$^{3+}$ passed through the solution layer to reach the mineral surface and was reduced on the mineral surface, thereby Cu$^{2+}$ was released. Buckley [7,8] studied the surface changes of natural copper ore in long-term exposure to the environment by using X-ray photoelectron spectroscopy (XPS) and found that the surface of minerals oxidized quickly with hydrated iron oxide and copper-sulfur compounds forming on the surface. As exposure time increased, iron migrated to the surface to form ferrites, leaving copper-containing sulfides behind. During the surface rusting, ferrites separated on the mineral surface from the outside and formed a rugged coating on the surface of the copper-sulfur compounds. It was found that bornite transformed into CuS and Cu$_2$S, and the iron element transformed into Fe(OH)$_3$ or FeS on studying the electrochemical dissolution of bornite [9,10]. Moreover, during oxidative leaching of bornite with oxygen in H$_2$SO$_4$, CuS and Cu$_3$FeS$_4$ were formed [11]. Many researchers suggested that the chemical formula of the bornite can be expressed as (Cu$_3$FeS$_4$)$_2$·2Cu$^+$. During the dissolution process, Cu$^+$ firstly transforms into Cu$^{2+}$ and forms Cu$_3$FeS$_4$, although there is no direct evidence to prove the existence of Cu$_3$FeS$_4$.

In addition, some researches have been conducted on bornite bioleaching in recent years. Bevilaqua et al. [12,13] mainly used electrochemical testing methods to study bornite bioleaching. It was found that the presence of Acidithiobacillus ferrooxidans can accelerate the leaching of bornite. The fitting data of electrochemical impedance spectroscopy (EIS) displayed that a passivation film was formed on the surface of the bornite which was confirmed by cyclic voltammetry (CV) data. Wang [14] found that after 30 days of leaching with A. ferrooxidans, the copper extraction rate of bornite was 72.35% at pH 2.0, initial Fe$^{2+}$ concentration at 9 g/L, and pulp density at 5%. Zhao et al. [15–17] mainly studied the mixed bioleaching process of bornite and chalcopyrite. The results showed that the presence of bornite could maintain the solution potential within a suitable range and thus accelerated the dissolution of chalcopyrite. At the same time, the galvanic effect between these two minerals also accelerated the dissolution process. Wang et al. [18] studied the bioleaching of mixed ore of bornite and pyrite by L. ferriphilum. It was found that the addition of pyrite in the mixed ore increased the copper extraction of bornite, which was related to the galvanic effect between them. Yang et al. [19] used XRD and XPS to characterize the intermediary species during bornite bioleaching by mixed moderately thermophilic culture, which suggested that bornite can be transformed into chalcocite, chalcopyrite, and covellite. Furthermore, Zhao [20] found that bornite tended to be directly oxidized to CuS, FeOOH, and S$^0$ on the surface in bioleaching by using moderately thermophilic culture.

Overall, it is obvious that the dissolution mechanism of bornite is not fully resolved. For example, whether CuS, CuFeS$_2$, Cu$_3$FeS$_4$, and/or Cu$_2$S are intermediate species during bornite bioleaching, whether different kinds of bacteria have an influence on the production of different intermediates in the bornite bioleaching, and whether these intermediates are present at the same time are all open questions. In addition, no previous studies have further investigated bornite bioleaching by L. ferriphilum or A. caldus and the intermediate phase transformation during the process remains unclear. Therefore, this paper further investigates the above problems to improve knowledge on the dissolution mechanism of bornite.

2. Materials and Methods

2.1. Minerals

The bornite was collected from the Yushui copper mine in Meizhou, Guangdong, China. After grinding and sieving, the 200 mesh (0.074 mm) mineral was obtained for experiments. Chemical analysis conducted by X-ray fluorescence (parameters given in Section 2.5) showed that the bornite sample contained (w/w) 62.30% Cu, 10.03% Fe, and 21.76% S (Table 1). XRD results indicated that bornite was the main mineralogical component so the ore samples used in the study were of high purity (Figure 1).
Electrochemical experiments were conducted with a conventional three-electrode cell. Graphite rods were used as the counter electrode and an Ag/AgCl (3.0 M KCl) electrode as the reference electrode. Carbon paste electrodes were used as the working electrode, which was made by mixing 0.7 g minerals, 0.2 g graphite, and 0.1 g solid paraffin. The mixture was heated and stirred in a 50 mL beaker till the paraffin melted. After that the mixture was transferred rapidly into a tablet model for tableting under a pressure of 150 kg/cm² for three minutes, and then taken out for air drying. The electrodes were polished using 1200-grit silicon carbide paper to obtain a smooth surface before electrochemical measurements.

Table 1. Chemical composition of the bornite.

<table>
<thead>
<tr>
<th>Elements</th>
<th>O</th>
<th>Si</th>
<th>S</th>
<th>Fe</th>
<th>Cu</th>
<th>Zn</th>
<th>As</th>
<th>Ag</th>
<th>Pb</th>
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<td>Wt %</td>
<td>2.91</td>
<td>0.63</td>
<td>21.76</td>
<td>10.03</td>
<td>62.30</td>
<td>0.15</td>
<td>0.15</td>
<td>1.35</td>
<td>0.47</td>
</tr>
</tbody>
</table>

2.2. Microorganisms

*L. ferriphilum* and *A. caldus* are both moderately thermophilic bacteria which are more suitable to relatively high temperature than mesophiles and to high pulp density than extreme thermophilic bacteria [21]. *L. ferriphilum* and *A. caldus* used in this work were provided by the Key Lab of Biohydrometallurgy of Ministry of Education, Central South University, Changsha, China. *L. ferriphilum* and *A. caldus* were cultured both in 9K medium containing (NH₄)₂SO₄ (3.00 g/L), KCl (0.10 g/L), K₂HPO₄ (0.50 g/L), MgSO₄·7H₂O (0.50 g/L), Ca(NO₃)₂ (0.01 g/L) [22] at 45 °C, 170 rpm, and pH 1.70. The energy resources of *L. ferriphilum* and *A. caldus* were FeSO₄·7H₂O (44.70 g/L) and S (10 g/L) respectively. The cell concentration was observed under a microscope every two days until it grew to the exponential growth stage with a cell concentration higher than 1.0 × 10⁷ cells/mL. Then suspension containing cells were collected by centrifugation (Beckman Coulter, Inc. Avanti j-E) at 10,976 × g and 4 °C for 20 min, bacteria adhering on the centrifuge bottle wall were rinsed with 1 mL sterilized distilled water which was acidified by 20% (w/v) dilute sulfuric acid to pH 2.0 and transferred into 10 mL centrifuge tubes. The cell concentration of the concentrated bacterial solution was higher than 1.0 × 10⁸ cells/mL.

2.3. Electrochemical Experiments

Electrochemical experiments were conducted with a conventional three-electrode cell. Graphite rods were used as the counter electrode and an Ag/AgCl (3.0 M KCl) electrode as the reference electrode. Carbon paste electrodes were used as the working electrode, which was made by mixing 0.7 g minerals, 0.2 g graphite, and 0.1 g solid paraffin. The mixture was heated and stirred in a 50 mL beaker till the paraffin melted. After that the mixture was transferred rapidly into a tablet model for tableting under a pressure of 150 kg/cm² for three minutes, and then taken out for air drying. The electrodes were polished using 1200-grit silicon carbide paper to obtain a smooth surface before electrochemical measurements.
The electrolyte used in the experiments was 9 K medium. The pH of the solution was adjusted to 1.70 with 20% (w/w) dilute sulfuric acid. Nitrogen gas was sparged 15 min before and during the electrochemical measurements to remove dissolved oxygen from the electrolyte.

All electrochemical experiments in this paper were performed with a Princeton Model 283 potentiostat (EG & G of Princeton Applied Research) coupled to a personal computer. Cyclic voltammetry (CV) tests were conducted at a sweep rate of 20 mV/s and potentiostatic polarization tests were performed for 1200 s. All potentials in this paper are against an Ag/AgCl electrode (3.0 M KCl).

2.4. Bioleaching Experiments

Bioleaching experiments were carried out in 250 mL Erlenmeyer flasks containing 100 mL of 9K medium and 2 g of bornite. The initial cell concentration in the solution was controlled at about $1.0 \times 10^7$ cells/mL by adding 1 mL of the concentrated bacterial solution. The temperature and rpm were set at 45 °C and 170 r/min, respectively. The pH was adjusted to $1.70 \pm 0.02$ with 20% (w/w) dilute sulfuric acid when the pH of solution was higher than 1.70. The evaporation loss was compensated for periodically by adding distilled water. After the leaching experiments, the solution was filtered through a neutral filter paper, and the filter residues were washed 3 times with distilled water having a pH of 1.70. After drying in a vacuum drying oven at normal temperature for 24 h, the dried filter residues were subjected to the corresponding tests.

2.5. Analytical Methods

Chemical analysis was conducted by X-ray fluorescence, using an X-ray fluorescence S4 Explorer spectrometer (Bruker AXS, Germany). The X-ray tube (Rh tube with a 75 µm window) was set at 40 kV and 15 mA and the sample powder was scanned over the range of 5–85°. The copper concentrations were analyzed by inductively coupled plasma-atomic emission spectrometry (ICP-AES) (America Baird Co. PS-6). The XRD analysis of mineral samples was performed by a Bruker D8 Advance X-Ray diffractometer (Cu Kα, λ = 1.5406 Å) with an operating voltage of 40 kV, temperature of 25 °C, current of 40 mA, steps of 0.004°, and a step time of 28.5 s. The XPS analyses of the mineral samples were performed by a Thermo Fisher Scientific ESCALAB 250Xi photoelectron spectrometer. The light source was a monochromatic Al excitation source (1486.6 eV) with a power of 200 W, a pass energy of 20 eV, a step size of 0.1 eV, and a vacuum of less than $1 \times 10^{-9}$ mbar. Calibration was using the C1s binding energy at 284.6 eV. Due to spin-orbit splitting, the 2p peak of S existed in the form of a double peak (S 2p$_{3/2}$ and S 2p$_{1/2}$), and the peak intensity of S 2p$_{3/2}$ was twice the intensity of the S 2p$_{1/2}$ peak. The binding energy of S 2p$_{1/2}$ peak is 1.19 eV higher than that of the S 2p$_{3/2}$ peak. The Gaussian–Lorentzian (SGL) function was used for fitting with a 50% Gaussian function and a 50% Lorentzian function. The background fitting was performed using the Shirley method. To simplify the graph, only the S 2p$_{3/2}$ peak is shown in the figures.

2.6. Calculation of Acid Consumption

The total consumption of acid during the leaching of bornite was calculated by Equation (1) [23].

$$N = \left(10^{-a} - 10^{-b}\right) \times 1000 \times V \div M$$  \hspace{1cm} (1)

where N is the consumption of H$^+$ for per gram of ore, with units of mmol/g, a is the pH value measured on the previous day (measured after adding sulfuric acid), b is the pH value measured on the following day (measured before adding sulfuric acid), V is the volume of leaching solution, with units of L and M is the weight of added minerals, with units of g.
3. Results and Discussion

3.1. Electrochemical Experiments

3.1.1. Potentiostatic Polarization Experiments

The potentiostatic polarization tests were mainly used to describe the strength of the surface reaction of the electrode and the redox properties of the surface reaction. Figure 2 shows the relationship between total charge (calculated based on the integrals from the current-time curves of bornite electrode) and the applied potential at pH at 1.0, 1.5, and 3.0. It can be seen that the oxidation region and reduction region of bornite at the above three pH conditions are similar, with the same critical point at 200 mV. This suggests that the difference of pH cannot change the principle reaction of bornite during bioleaching. The bornite tends to be directly oxidized when the redox potential is higher than 200 mV and to be reduced when the redox potential is less than 200 mV. In addition, the slopes of the curves in the oxidation region and reduction region increase as the pH increases, which suggests that bornite is easier to be oxidized at lower pH. Therefore, the pH of the solution is the main step to limit the leaching of bornite.

![Figure 2. Relationship between total charge and applied potential at pHs of 1.0, 1.5, and 3.0.](image)

3.1.2. Cyclic Voltammetry Measurements

Cyclic voltammetry measurements of bornite electrodes at different pH in 9 K media were carried out by the positive scan route. The positive scan route, scanning twice in total, started scanning from 200 mV to 800 mV, then reversed to −800 mV, and finally switched towards the anodic direction again. The result is shown in Figure 3. Few anodic and cathodic current peaks are observed with the bornite electrode at pH 3.0, however, five distinct peaks are detected in the other two curves. It is obvious that bornite hardly reacts at this pH. This is consistent with the conclusion from Figure 2. The reaction indicated by the anodic peak a, is related to the oxidation of copper as reported in the literature [15]. Peak b appears in the potential range of 600 mV to 800 mV, which is an indication of the formation of covellite [24,25]. The first cathodic peak c represents the reduction of covellite formed during the oxidation of bornite [9,15]. The cathodic peaks d and peak e are considered to be the reduction of covellite or bornite and copper (I) to metallic copper, respectively [13].

Additionally, it is clear that each peak of the bornite electrode at pH 1.0 is higher than at pH 1.5, which means the high pH value of solution will hinder the oxidation of bornite. On this basis, considering the optimum pH range of the bacteria growth, the following bioleaching experiments were carried out.
3.2. Bioleaching Experiments

Figure 4 shows the variations of pH during bornite bioleaching by *L. ferrophilum*, *A. caldus*, and the sterile control. The pH of the sterile control solution rises every day for the first 14 days of leaching. The pH begins to decrease from day 14 to day 21, and then the previous changes are restored. Variation of pH in the presence of *L. ferrophilum* is similar to the sterile control, except that the degree of change is more intense. It indicates that the presence of *L. ferrophilum* cannot change the oxidation principle of bornite. However, the variation of pH in the presence of *A. caldus* is significantly different from the behavior at the above two conditions. The pH rises only in the first three days then is lowered to a very low level at about pH 1.0 in the initial stage of leaching. It may relate to the oxidation of sulfur-containing compounds that cause a rapid decrease in pH of the solution. Therefore the presence of *A. caldus* may accelerate the leaching of bornite more in the initial stage of bornite leaching than that in the cases of *L. ferrophilum* and sterile control. After that, the rise of pH is slow and after 20 days, the variation is similar to the former two conditions.
The total acid consumption of *L. ferriphilum*, *A. caldus* and the sterile control during bornite bioleaching is shown in Figure 5, which indicates that the dissolution of bornite is acid-consuming and the reaction is shown in Equation (2) [26]:

\[
4\text{Cu}_5\text{FeS}_4 + 37\text{O}_2 + 20\text{H}^+ \rightarrow 20\text{Cu}^{2+} + 4\text{Fe}^{3+} + 16\text{SO}_4^{2-} + 10\text{H}_2\text{O}
\]  

(2)

**Figure 5.** Total acid consumption of *L. ferriphilum*, *A. caldus* and sterile control during bornite bioleaching.

The result of total acid consumption of *L. ferriphilum* is close to the sterile control while the total acid consumption of *A. caldus* is slightly higher than the sterile control.

Figure 6 shows the variations in the copper extraction rate during bornite bioleaching. During the whole leaching process, the copper extraction of bornite by *L. ferriphilum* and *A. caldus* is significantly higher than that by the sterile control. After leaching for 25 days, the copper extraction rate of the bacteria group reaches a maximum of about 70%, while the sterile control is only 36% after 30 days. It can be seen that the presence of *L. ferriphilum* and *A. caldus* significantly accelerates the bornite leaching process. The copper extraction rate of the sterile control increases rapidly in the first five days, reaching 15%, and then the growth becomes slow. However, the copper extraction rate of the bacteria group increases rapidly until around the 15th day. Although the final copper extraction rate of the bornite is similar in the presence of *L. ferriphilum* and *A. caldus*, in the latter it rises faster in the early stage of leaching. This is in line with the result of pH variation in the bornite bioleaching at the early stage by *A. caldus* (Figure 4).

*L. ferriphilum* mainly plays the role of oxidizing Fe\(^{2+}\) and producing Fe\(^{3+}\) during the leaching process. The Fe\(^{3+}\) as an efficient oxidant accelerates the leaching process of the bornite, as shown in the following equation [4,13].

\[
\text{Cu}_5\text{FeS}_4 + 12\text{Fe}^{3+} \rightarrow 5\text{Cu}^{2+} + 13\text{Fe}^{2+} + 4\text{S}^0
\]  

(3)

\[
\text{Fe}^{2+} \xrightarrow{\text{L. ferriphilum}} \text{Fe}^{3+}
\]  

(4)

The main role of *A. caldus* is to convert sulfur to sulfate as reported by Gu et al. [27] and the equation is shown below.

\[
\text{S} + \frac{3}{2}\text{O}_2 + \text{H}_2\text{O} \xrightarrow{\text{A. caldus}} \text{H}_2\text{SO}_4
\]  

(5)
3.3. X-ray Powder Diffraction Analysis

Figure 7 shows the X-ray powder diffraction analysis of bornite leached residues by L. ferrophilum. In the early stage of leaching, the peaks of covellite (CuS, PDF#78-2121) and mooihoekit (Cu9Fe9S16, PDF#27-0165) are detected in the XRD pattern. Actually, the diffraction peaks of covellite and mooihoekit both fitted well to the leached residues because their crystal parameters are very similar. The results suggest that the dissolution of bornite is easily converted into a non-stoichiometric ratio of copper iron sulfide in the early stage of bioleaching. Some previous studies reported that covellite coating was also found in the early stage of leaching during a hydrometallurgical process of bornite [11]. However, some CuS peaks can still be observed in the late stage of leaching, which may relate to the mutual conversion of CuS and other copper iron sulfide intermediates. Then mooihoekit transforms into isocubanite (CuFe2S3, PDF#27-0166), as peaks of isocubanite are observed from the 12th day as the reaction proceeds. Wang et al. [18] also obtained a similar conclusion when studying the effect of pyrite on bornite bioleaching. Then the characteristic peaks of elemental sulfur (S8, PDF#74-1465) are clearly observed on the 16th day. The number of elemental sulfur peaks gradually increases until the end of leaching. In combination with the phenomenon that the copper extraction rate of bornite in the presence of L. ferrophilum becomes slow after the 15th day (Figure 6), it can be considered that the occurrence of elemental sulfur may cause passivation of the bornite. The phenomenon of passivation in bioleaching is widespread, not only in laboratory experiments but also in industrial production such as in recovering the metals from a complex waste matrix or solid wastes using bioleaching [28,29]. It was generally believed that elemental sulfur formed and covered the surface of the minerals or secondary resources, which stopped further reaction. Taking chalcopyrite leaching as an example, it was observed that an impervious sulfur layer formed on the surface of chalcopyrite which reduced the contact of bacteria or reactants with minerals [30–32]. Moreover, the passivation layer functioned as a capacitor, which made the diffusion of molecules and ions to and from the mineral surface difficult [33]. In a word, the formation of a passivation layer will reduce the extraction. Therefore, some researchers made studies on depassivation [34,35], in order to improve the metallic extraction. In any case, the problem, including the specific passivation mechanism and the method of depassivation, is worthy of further consideration.
It is notable that the intermediate species of bornite during bioleaching in the presence of *L. ferriphilum* are the same as in the presence of *A. caldus*. The difference between them is that the elemental sulfur of the latter one appears earlier, and the peaks of elemental sulfur are observed on the fourth day. The result is consistent with the conclusion of a faster leaching speed of bornite dissolution by *A. caldus* in the early stage of bioleaching (Figure 6).

Therefore, according to the results of the XRD patterns (Figures 7 and 8), it is considered that the dissolution process of the bornite is: \( \text{Cu}_9\text{Fe}_4\text{S}_8\rightarrow \text{Cu}_9\text{Fe}_2\text{S}_{16}\rightarrow \text{CuFe}_2\text{S}_3\rightarrow \text{CuS}\rightarrow \text{Cu}^{2+} \), which agrees with the conclusion of Wang [18]. Considering that these two kinds of bacteria have similar
intermediate species during bornite bioleaching, the following XPS analysis of bornite residues leached by *L. ferriphilum* was performed.

3.4. X-ray Photoelectron Spectroscopy Analysis

The Cu 2p3/2 peak of the bornite is located at 932.4 eV, the Cu 2p1/2 peak is at 952.3 eV, and there is no satellite peak between 940 and 950 eV (Figure 9a), which indicates that Cu in the bornite exists in the form of Cu (I) not Cu (II) [15, 36]. On the 7th day and the 21st day of leaching, the Cu 2p peak position and the bimodal spacing in the leached residues remain substantially unchanged, which suggests that Cu is also present in the form of Cu (I) during leaching. The Cu 2p3/2 peak at 932.4 eV represents CuS [37], which means Cu 2p3/2 Peaks in the leached residues on day 7 and 21 are all derived from CuS, which is consistent with the CuS appearing in the XRD pattern. The Cu LMM peak of the ore is located at 569.2 eV (Figure 9b). On the 7th and 21st day, the Cu LMM peaks of the leached residues are located at 568.9 eV and 568.5 eV, respectively, which is also consistent with the binding energy value (568.5 eV) of the previously reported CuS LMM peak [20, 38, 39].

![Figure 9. Cu 2p XPS spectra (a) and Cu LMM XPS spectra (b) of bornite residues after bioleaching for 7 days and 21 days.](image)

The S 2p3/2 XPS spectra of original bornite, and of leached residues at 7 days and 21 days are shown in Figure 10. The strongest S 2p3/2 peak in the initial bornite is located at 161.4 eV, representing S2− in the bornite bulk, and the location of the S2− peak is slightly different from the results reported at 161.8 eV by Harmer and Acres et al. [40, 41]. The fitting results of Harmer and Acres et al. showed a small S2− 2p3/2 peak at 160.1 eV at the bornite surface. It can be seen that the fitting is not good when fitting the surface layer S2− separately. So two kinds of S2− are not distinguished in this paper, which is consistent with the work of Harmer and Acres et al. [40, 42]. The elemental sulfur 2p3/2 peak is over a wide range (163.05–164.7 eV), and the polysulfide was also reported over a wide range (163.0–163.9 eV) [37]. Moreover, polysulfides have a variety of species with different chain lengths, and it is difficult to achieve a differentiating fit [36], therefore, the elemental sulfur and polysulfide were combined in the range (163.0–164.7 eV) for fitting.

Figure 11 shows the compositions of sulfur species of bornite leached by *L. ferriphilum* for 7 days, and 21 days. The content of S0/S0 in the original bornite is 13.53%, and the content of S0/S0 after leaching for 7 days and 21 days is 44.88% and 50.00%, respectively. The substance with a peak at 162.3–162.4 eV is S2− [37]. It can be found that the original bornite contains 23.37% of S2−. After 7 days of leaching, the content of S2− is 30.97% and 29.78% after 21 days of leaching. It can be seen from the variation of sulfur species contents that S2− and S2− are gradually transformed into S0 during bornite bioleaching. S2O3− is not found in the fitting results, indicating that the sulfur metabolism pathway of bornite in the presence of *L. ferriphilum* is mainly the polysulfide pathway. The leached residues were washed three times with distilled water with a pH of 1.70, and SO42− was mainly derived from jarosite [43]. On the 7th day and the 21st day of leaching, SO42− in the leached
residues is only about 5%, indicating that there is only a small amount of jarosite formed. However, few peaks of jarosite are observed in the XRD pattern, perhaps because of the relatively low content of jarosite. Therefore, the formation of elemental sulfur is the main reason for passivation in the last stage of bioleaching.

![Figure 10. S 2p₃/₂ XPS spectra of original bornite (a) and leached residues: (b) 7 days and (c) 21 days.](image)

![Figure 11. Compositions of sulfur species of bornite leached by L. ferriphilum at 7 days and 21 days.](image)
4. Conclusions

The results of bioleaching experiments and electrochemical experiments indicated that the dissolution of bornite was an acid-consuming process. Compared with the sterile control, the presence of \textit{L. ferriphilum} and \textit{A. caldus} significantly accelerated the leaching of bornite. The maximum copper extraction rate of the bornite bioleaching by \textit{L. ferriphilum} and \textit{A. caldus} was similar at about 70%. XRD results showed that with the bornite bioleaching by \textit{L. ferriphilum} or \textit{A. caldus}, a large amount of mooihoekit (Cu$_9$Fe$_9$S$_{16}$) and covellite (CuS) were produced in the early stage of bioleaching. Moreover, isocubanite (CuFe$_2$S$_3$) that might be converted from mooihoekit (Cu$_9$Fe$_9$S$_{16}$) was observed on the 12th day. The X-ray photoelectron spectrum of the leached residues revealed that Cu was present in the form of Cu(I) during the bornite bioleaching. Additionally, the S 2p$_3/2$ photoelectron spectrum suggested that S$_2^-$ and S$_2^{2-}$ were gradually converted to S$_n^{2-}$/S$^0$, so the formation of elemental sulfur may hinder the further dissolution of the bornite.

Author Contributions: The experiments were conceived and designed by M.H., G.Q. and J.W.; The experiments were performed by M.H., R.L. and X.H.; The data was analyzed by M.H. and X.W.; J.W. and G.Q. contributed reagents/materials/analysis tools; The paper was written by M.H.; M.H., X.W., J.W., L.W. and C.F. improved and revised the paper.

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