Suppression of pyrite oxidation by carrier microencapsulation using Si and Catechol

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ABSTRACT

Acid mine drainage (AMD) is formed from the natural oxidation of sulfide minerals such as pyrite, FeS2. Prevention of acid mine drainage is very important and several techniques are currently being investigated for the treatment and abatement of AMD. This paper proposes carrier-microencapsulation (CME) using Si-catechol complex- Si(cat)₃²⁻ for preventing pyrite oxidation. In CME, the water soluble organic carrier-catechol and metal ion-Si makes a complex e.g., tris-catecholato complex of Si⁴⁺, which oxidatively decomposes on pyrite surface and forms a stable oxide or hydroxide e.g., Si(OH)₄ or SiO₂ layer on the pyrite surface as a protective coating against pyrite oxidation.

To demonstrate the effect of CME using Si(cat)₃²⁻ on pyrite oxidation, shaking-flask leaching experiments of pyrite without and with CME treatment were performed. Significant halt in pH drop was observed after the CME treatment of pyrite. The CME coating was found very effective even at low Si(cat)₃²⁻ concentration of 1 mol m⁻³ and 5 mol m⁻³. The amounts of Fe and S leached were lower from the pyrite with CME treatment than without CME treatment, indicating that CME using Si and catechol is effective in suppressing pyrite oxidation.

The effect of pH and presence of microorganism was also evaluated. The CME coating was found very effective even in acidic pH range. The CME treatment suppressed pyrite oxidation even in the presence of iron oxidizing microorganism. Presence of Si and O on the residue pyrite surface in SEM-EDX analysis supported the sustainability of CME coating at acidic pH.

KEYWORDS: Acid mine drainage; pyrite oxidation; carrier micro encapsulation

INTRODUCTION

Exposure of the sulfide bearing wastes such as pyrite (FeS₂) to the oxidizing environment results to a severe environmental problem called acid mine drainage (AMD): it is defined as a very low pH leachate formed by the oxidation of sulfide minerals contained in the mining and mineral processing waste [Kafui and Egiebor, 1995]. The problem becomes more severe when the toxic low pH leachate, from the mine site, contaminates downstream water bodies [Fillion et.al. 1990].

Pyrite oxidation occurs in the presence of oxygen and water and it is catalyzed by the iron oxidizing bacteria: Acidithiobacillus ferrooxidans. The Fe²⁺ ion formed as oxidation product can be further oxidized to Fe³⁺ that enhances the further oxidation of pyrite and lowers the pH at the same time [Temple and Colmer 1951]. Since AMD is a naturally “ongoing” process, once the process starts acidic water is continually formed even after mine closure and the release can be for centuries or even millennia [Logsdon, 2002]. The extent of the problem can be imagined from its total worldwide liability, which is likely to be in excess of 10000 millions of US dollars. In US alone the mining industries spend more than US$1 million every day to treat AMD water [Brown et.al. 2002].
A number of methods reported in the past tried to inhibit the pyrite oxidation by physical, chemical, or biological means. Some of the reported approaches for remedies of AMD include: adding low cost lime to neutralize the acid and precipitate heavy metals as hydroxides, underwater disposal of sulfide waste [Robertson, 1994], use of nonpermeable liners such as plastic, clay, and asphalt to provide a safe anoxic setting for storing sulfide waste [Swanson et al. 1997], the consumption of bactericides to reduce the population of catalyzing bacteria and hence diminish the rate of pyrite oxidation [Dugan, 1987], complexation of ferric ions in solution or at the pyrite surface with hydroxyl functional group to make pyrite surface unavailable for oxidation [Chander and Zhou 1992], phosphating in low and high temperature [Kafui and Egiebor, 1995], and coating of the pyrite surface with phospholipids or ferric-phosphate to inhibit oxygen diffusion to pyrite surface [Kargbo et al. 2004, Evangelou, 2001]. However, problems associated with the mentioned approaches made them difficult to be widely accepted as AMD remediation techniques. For instance, the traditional approach for AMD treatment, addition of low cost lime, has big issue on sludge disposal, as the sludge produced by lime neutralization process is voluminous with low solid content and low chemical stability, has no economic value, and requires specially designed landfill for disposal [Murdock et al. 1994, Mc Donald et al. 2006]. Liner technology- by using clay or other nonpermeable liners to design safe disposal site for sulfide waste can be cost-prohibitive but ineffective well due to the liners’ deterioration with time.

Recently, an approach to coat the pyrite surface by titanium or silicon oxides known as carrier micro encapsulation (CME) of pyrite was reported by the present authors [Satur et al. 2007, Jha et al. 2008]. In CME, pyrite is treated with aqueous solutions of metal ion (Ti$^{4+}$ or Si$^{4+}$) complex with organic carrier such as catechol (cat), i.e. [Si(cat)$_3^{2-}$] or [Ti(cat)$_3^{2-}$]. The complex is adsorbed on the pyrite surface and oxidatively decomposes on the surface, releasing metal ions that are hydrolyzed to make a thin layer of metal oxide or hydroxide. It is found that the CME coating on pyrite surface suppresses floatability of pyrite during coal-processing. CME, in fact, seems highly promising to reduce the SO$_2$ emission (due to combustion of pyrite with coal) because of its effective separation of pyrite from coal [Jha et al. 2008].

In the present work, the authors had evaluated the effect of CME on pyrite oxidation using tris-catecholato complex of Si$^{4+}$ [Si(cat)$_3^{2-}$]. Shaking flask leaching experiments were carried out to verify the effect of CME on pyrite oxidation with pH and amounts of Fe and S leached as indicators. The result confirms that CME coating is able to slow down the pyrite oxidation rate successfully.

MATERIALS AND METHODS

Material:

A ground pyrite sample from Peru was used in this experiment. The pyrite sample was crushed in jaw crusher and grinded in ball mill followed by sieving to obtain the 53 to 75 µm size fractions. The sieved pyrite was washed following the procedures of Sasaki et al. 1995: washing with ethanol with ultrasonic vibration initially applied on the solution for 30 s, then washing with 1 kmol m$^{-3}$ HNO$_3$ to create fresh pyrite surfaces, distilled water to wash off the nitric acid, and acetone to facilitate drying, and finally vacuum drying the washed pyrite for 24 h [Sasaki et al, 1995]. The tris-catecholato complex of Si$^{4+}$ [Si(cat)$_3^{2-}$] for CME treatment was prepared by mixing 0.142 g of Na$_2$SiO$_3$.9H$_2$O and 0.165 g of catechol in 10 cm$^3$ distilled water. This solution contains 5 mol m$^{-3}$ Si(cat)$_3^{2-}$. The solution pHs were maintained by adding HCl or NaOH solution.

Methods:

Carrier micro encapsulation treatment
1 g of pyrite samples are shaken with the different known concentration of 10 cm$^3$ of Si(cat)$_3^{2-}$ solution inside water bath shaker for desired time period. The shaking amplitude of water bath shaker is 4 cm; and the frequency is 120min$^{-1}$. The pyrite sample was filtered and washed for shaking flask experiments.

**Long and short time shaking flask experiments**

The effect of CME on pyrite oxidation was investigated with different concentrations of Si(cat)$_3^{2-}$ by shaking flask leaching experiments at long and short time periods. For reference, the same experiment was carried out with the pyrite without any CME treatment.

The long time shaking flask experiment (one month) was carried out on 2g of CME treated pyrite shaken in 20cm$^3$ of distilled water under aerobic condition using a water bath shaker at 25°C. The shaking amplitude of the water bath shaker was 4 cm; and the frequency was 120min$^{-1}$. At fix intervals of 1 week, the 2 cm$^3$ of leachate were sampled from the flask to measure the pH and ORP and the Fe and S contents of the leachate by ICP-AES.

To determine the effectiveness of the CME treatment at acidic conditions, shaking flask experiment for short time (one week) was done on 1g of pyrite shaken along with Si(cat)$_3^{2-}$ and 10 cm$^3$ of distilled water adjusted at different pH conditions from 0-6 by adding HCl.

Another long time shaking flask experiment (4 weeks) was carried out on 1g of CME treated pyrite along with 5 mol m$^{-3}$ of Si(cat)$_3^{2-}$ and acidithiobacillus ferrooxidans acidophilic bacteria. At fixed intervals of 1 week, the flask was removed from the shaker to measure the pH and ORP of the leachate.

After the desired time of interval of leaching experiment, solid-liquid filtration was carried out and the filtrate thus obtained was analyzed by Inductively Coupled Plasma Atomic Emission Spectrometry (ICP-AES) to measure the Fe, and S content in it. The surface morphologies of the untreated and CME treated pyrite were analyzed by SEM- EDX.

RESULT AND DISCUSSION

Catechol is a week diprotic acid (pKa1=9.2, pKa2 =13) and therefore can remain in the undissociated form over a wide pH range [Rodriguez et. al. 1996]. It can form a triscatecholato silicon(IV) complex with Si$^{4+}$ - Si(cat)$_3^{2-}$ - [Kansal and Laine, 1994, Caulfield et.al. 2001]. In the present work, triscatecholato silicon (IV) complex -Si(cat)$_3^{2-}$ - was used for making a thin stable layer of SiO$_2$ or Si(OH)$_4$ on the pyrite surface to inhibit pyrite oxidation. In carrier micro encapsulation (CME), Si(cat)$_3^{2-}$ complex is oxidatively decomposed on the pyrite surface and the released Si$^{4+}$ ion hydrolyses to form a thin layer of SiO$_2$ or Si(OH)$_4$ on the pyrite surface [Jha et.al. 2008].

The reaction involved in AMD can be illustrated as:

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\text{FeS}_2+7/2 \text{O}_2+\text{H}_2\text{O} \rightarrow \text{Fe}^{2+} + 2\text{SO}_4^{2-} + 2\text{H}^+ \ldots \ldots \ldots \ldots (I)
\]

\[
\text{FeS}_2 + 2\text{H}^+ + 1/2 \text{O}_2 \rightarrow \text{Fe}^{2+} + 2\text{S} + \text{H}_2\text{O} \ldots \ldots \ldots \ldots (II)
\]

The Fe$^{2+}$ produced can be further oxidized by O$_2$ into Fe$^{3+}$, which in turn will precipitate as Fe(OH)$_3$ and lower the pH at the same time [Evangelou,1995]. Thus, the pH and the amounts of Fe and S present in the solution are the main indicators of AMD. Therefore, in this paper the effect of CME on pH drop and Fe and S leachability was analyzed.

The effect of CME on pyrite oxidation was investigated with different concentrations of Si(cat)$_3^{2-}$ by shaking flask leaching experiments. For reference, the same experiment was carried out without CME treatment.
Figure 1(A)(B)(C) shows the result of the one month shaking flask experiment as indicated by the final pH of leachate, amount of Fe and S present in leachate after four weeks as a function of concentration of Si(cat)\textsuperscript{3-}. The initial pH of the leachate was around 3.5 and after four weeks the pH of leachate for untreated pyrite dropped down to 1.7 while the pH for the CME-treated pyrite (Si(cat)\textsuperscript{3-} concentrations of 1 to 10 mol m\textsuperscript{-3}) was higher than 2.7. CME treatment with 5-10 m mol\textsuperscript{-3} of Si(cat)\textsuperscript{3-} demonstrated same capacity to halt the pH drop. In Figure 1(B) and 1(C) the amount of Fe and S present in leachate after the four week of leaching is presented and it shows that the amount of Fe and S present in the leachate of CME treated, (Si(cat)\textsuperscript{3-} concentrations of 1 to 10 mol m\textsuperscript{-3}), pyrite is very low in comparison of that for the non treated pyrite leachate. For the succeeding experiments, the concentration of Si(cat)\textsuperscript{3-} are decided to be kept at 5 mol m\textsuperscript{-3}.

Figure 2 shows the pH, Fe and S contents in the leachate obtained at one week intervals for one month shaking flask experiment. In Figure 2(A) the pH trend with leaching time is presented. The result shows that the pH of the leachate of the pyrite with CME treatment is higher than that of the leachate of the untreated pyrite. The pH drop for un-treated pyrite was significant and after four weeks it was recorded 1.7. on the other hand, the pH of the leachate of the pyrite with CME treatment was 3.2 over the one month experiment. Figures 2 (B) and (C) show the Fe and S concentrations (in ppm) in the leachate, respectively. The amount of Fe and S leached from the CME treated pyrite is significantly less in comparison to that of from un-treated pyrite after one month leaching. Figure 2(B) depicts that after 30 days of leaching 160ppm of Fe ion was present in the leachate of non-treated pyrite, on the other hand only 60ppm of Fe ion was present in the leachate of CME treated pyrite, indicating that suppression in Fe leaching behavior is achieved by CME treatment of pyrite. Suppression of sulphur leaching is also observed for CME treated pyrite: Figure 2(C) depicts that after 30 days leaching 20ppm of S ion was present in the leachate of non-treated pyrite, while the leached amount of S ion for CME treated pyrite leachate sample was only 2ppm. In addition, the higher mass ratio of Fe and S in the leachate could be explained based on the equation (II) i.e. the formation of elemental sulfur on pyrite oxidation. These results imply that the SiO\textsubscript{2}/Si(OH)\textsubscript{4} layer formed on the pyrite surface by the CME treatment confirmed itself as a barrier layer against pyrite oxidation by bringing a halt in pH drop and by controlling the leaching of Fe and S.

The effectiveness of the CME treatment at acidic conditions was investigated next. After the CME treatment with 5 mol m\textsuperscript{-3} Si(cat)\textsuperscript{3-}, the pyrite sample was shaken in the leaching solution of different pH ranging from highly acidic to natural (0-6) for a week. The leached amounts of Fe and S in leachate of pyrite without and with CME treatment after one week leaching experiment is shown in Figure 3(A) and (B) as a function of final pH.

The results show that for the CME treated pyrite, the highest amount of leached Fe was around 5 ppm in the investigated pH region. For the un-treated pyrite, the leached amount of Fe was high at the low pH region: 35-40ppm, but the amount decreased at around pH 5. This may be due to the precipitation of Fe\textsuperscript{3+} as Fe(OH)\textsubscript{3}. In Figure 3(B), the amounts of S ion present in the leachates after one week show that the S leaching is insignificant (<1ppm) for the CME treated pyrite. The amount of S ion leached from the un-treated pyrite was higher (6ppm) than from that of CME treated pyrite in the investigated pH range (0-6). The results shown in Figure 3 indicate that even in the acidic pH range, pyrite oxidation was suppressed successfully by CME treatment.

After one week leaching in different pH, pyrite residue were collected and dried for analysis of the surface morphology analysis by SEM-EDX. The SEM image and elemental mapping for the pyrite residues pyrite without and with CME treatment are shown in Figure 4. The homogeneous existence of Si on the surface of CME treated pyrite leached at ph 2.98, supports the formation of SiO\textsubscript{2}/Si(OH)\textsubscript{4} layer on pyrite by CME and the stability of the layer at low pH. Comparatively weak signals of Si for non treated pyrite sample are obtained, which may be due to the Si contained in the pyrite itself. The coating formed by CME endured itself and inhibited the Fe and S leaching at very low pH of 2.98. Sustainability
of CME coating at highly acidic pH condition verifies that CME is a suitable technique for suppression of pyrite oxidation.

The presence of iron oxidizing bacteria (*Acidithiobacillus ferrooxidans*) may also enhance the spontaneous pyrite oxidation [Temple and Colmer, 1951]. In this context, the effectiveness of CME coating was investigated in the presence of iron oxidizing bacteria. One month shaking flask experiment on the CME treated (with 5 mol m\(^{-3}\)Si(cat)\(_3\)^2\(^{-}\)) and un-treated pyrite was done in presence of *Acidithiobacillus ferrooxidans*; the results of the pH profile, amounts of Fe and S leached at 1 week intervals are presented in Figures 5 (A) (B) (C).

Figure 5 (A) shows the pH profiles of the leachates of pyrite with and without CME in the presence of iron oxidizing bacteria-*Acidithiobacillus ferrooxidans*. It shows that CME could control the pH drop even in the presence of iron-oxidizing bacteria. Comparing the results of with and without CME treatment, the pH values of the leachate of CME treated pyrite were over 2.3 throughout the investigated time period, where the pH of the leachate of the un-treated pyrite were lower and drops down to 1.8 at the fourth week.

In Figure 5(B) and (C) show the amount of Fe and S leached from the pyrite with and without CME treatment as a function of time respectively. In Figure 5(B) the amount of Fe present in the leachate for non treated pyrite after four weeks is almost twice as of in the CME treated pyrite leachate: the leached amount of Fe from the un-treated pyrite increased with passing weeks and was 750ppm, but only 350ppm from the CME treated pyrite after four weeks. In Figure 5(C) the behavior of S leaching from pyrite with and without CME treatment is depicted. The sulfur leaching for non treated pyrite is almost double than that of CME treated pyrite after four weeks. Continuous extraction of S was observed from the un-treated pyrite and S concentration in the leachate reached 24ppm at five weeks contrary to less than 10ppm in the leachate of the CME treated pyrite. These results show that the Fe and S leaching was successfully suppressed by the CME treatment even in the presence of ample amount of iron oxidizing bacteria. Therefore it can be stated that the presence of bacteria would not hinder suppression of pyrite oxidation by the CME.

**CONCLUSION**

In this study, carrier-microencapsulation (CME) using Si and catechol [Si(cat)\(_3\)^2\(^{-}\)] was proposed for preventing pyrite oxidation and was demonstrated by shaking flask leaching experiments. The Si(OH)\(_4\) or SiO\(_2\) coating made by CME treatment was found to be effective in controlling pH drop over one month experiment. The Fe and S leachability was also suppressed by the CME treatment. The suppressive effect of CME coating was effective even in the acidic pH range (pH 0-6) and even in the presence of iron oxidizing bacteria.

**REFERENCES**


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Figure 1: (A) Final pH of the leachate, (B) amount of Fe (in ppm) present in the leachate, and (C) amount of S (in ppm) present in the leachate, obtained after four week of shaking flask experiment as function of Si(cat)$_3^{2-}$ concentration. CME treatment time was 1h.
Figure 2: (A) pH profile of the leachate, (B) amount of Fe (in ppm) and (C) amount of S (in ppm) in the leachate of pyrite without and with CME treatment, obtained from shaking flask experiments for one month at one week interval.
Figure 3: (A) Amount of Fe (in ppm) and (B) amount of S (in ppm) present in the leachates of pyrite without and with CME treatment after 1 week shaking flask leaching experiment at acidic to natural pH. The Si(cat)$_3^{2-}$ concentration was 5 mol m$^{-3}$. 

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\text{Si(cat)}_3^{2-} \text{ concentration was } 5 \text{ mol m}^{-3}. 
\]
Figure 4: SEM image and EDX mapping for elements on the pyrite particle leached in low pH for 1 week in aerobic condition. (A) Pyrite without CME treatment, leachate pH was 2.34. (B) Pyrite with CME treatment, leachate pH was 2.98. The Si(cat)$_3^{2-}$ concentration was 5 mol m$^{-3}$. 
Figure 5: (A) pH profile of leachate of pyrite with and without CME (B) amount of Fe and (C) amount of S present in the leachate of pyrite with and without CME treatment shaken in the presence of iron oxidizing bacteria for four weeks. Si(cat)$_3^{2-}$ concentration was 5 mol m$^{-3}$. Bacteria density was: 1.0 x10$^8$cells/ml.