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Suppression of floatability of pyrite in coal processing by carrier microencapsulation

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ABSTRACT

Carrier microencapsulation, CME, is a technique to form a thin layer of metal oxide or hydroxide on pyrite surface using a water soluble organic carrier combined with metal ions. The present study investigated the effect of CME using tris-catecholato complex of Si⁴⁺, Si(cat)₃²⁻ on pyrite-coal separation by dynamic bubble pick-up experiments and Hallimond tube flotation experiments using coal, pyrite, and a coal-pyrite mixture. The mineral samples were treated in 0-5 mol m⁻³ Si(cat)₃²⁻ solutions at pH 4-9 at treatment times of 1-24 h. Dynamic bubble pick-up experiments showed that CME treatment
converted the pyrite surface from hydrophobic to hydrophilic but did not affect coal’s hydrophobic surface. The results of the Hallimond tube flotation experiments of a coal-pyrite mixture at pH 7-9 in the presence of kerosene as a collector showed that pyrite floatability was selectively suppressed after 1 h CME treatment with 0.5 mol m$^{-3}$ Si(cat)$_3$$^{2-}$ while both coal and pyrite were floated without the treatment. This indicates that CME treatment is effective in suppressing pyrite floatability in coal-pyrite flotation.

*Keywords*: Pyrite; Flotation; Microencapsulation; Surface property; Coal processing

**Introduction**

Pyrite, FeS$_2$, is one of the most abundant minerals, yet economically insignificant and less desirable. It is a common gangue mineral in coal, and causes SO$_2$ emissions during coal combustion. SO$_2$ is one of the main constituents for acid rain formation therefore the separation of pyrite from coal before combustion is environmentally important [1, 2].

Dense media separation, cyclones, jigs, and froth flotation are some of the existing techniques for coal processing [3]. Gravity separation methods e.g. dense media separation, cyclones, and jigs are quite efficient in separating pyrite from coal because pyrite density ($\sim$ 5 g cm$^{-3}$) is much higher than that of coal (1.3-1.5 g cm$^{-3}$). However, the efficiency of gravity separation falls rapidly for particles, finer than 100 µm [1]; therefore froth flotation is used for washing of fine coal [4]. The amount of coal processed by froth flotation is about 13.6% of the total coal preparation [3].
In froth flotation, air bubbles are introduced into a mineral suspension. Hydrophobic minerals attach to the air bubbles and are recovered as the froth product but hydrophilic minerals remain in the suspension and are discarded to the tailing pond as the tailing. Coal is hydrophobic and is separated from hydrophilic gangue minerals e.g., pyrite in the flotation. However, the natural hydrophobic property of pyrite makes coal-pyrite separation difficult as significant amounts of pyrite are recovered in the froth along with the coal [5, 6].

The other problem associated with pyrite in coal is the formation of acid mine drainage or AMD in the tailing dam when pyrite is disposed as tailing from the flotation process. The disposed pyrite comes in contact with oxygen and water and is oxidized to produce sulfuric acid, which can dissolve toxic heavy metals from other minerals [7].

Relating to the above two problems (recovery of entrapped pyrite in the froth during flotation and AMD formation due to pyrite oxidation), many studies have been done in the past. To suppress pyrite floatability in flotation, the use of depressants such as sodium sulfide, potassium dichromate, potassium permanganate, ferrous/ferric sulfate, aluminum and ferric chlorides, and sodium cyanide [6,9], bacteria like Acidithiobacillus ferrooxidans [8, 10], as well as several preconditioning techniques [11-12] have been reported. For controlling AMD formation, application of nonpermeable liners (e.g., clay or plastic liner in the tailing dam) and techniques to make a protective coating on pyrite
to suppress oxidation were reported [13-16]. However, a single technique to overcome both problems simultaneously has not yet been developed.

The authors have proposed carrier microencapsulation (CME) as a new coating technique for pyrite. In CME, pyrite is immersed in aqueous solutions of an organic carrier (e.g., catechol) combined with a metal ion (e.g., Ti^{4+} or Si^{4+}). The complex adsorbs on the pyrite surface and the organic carrier is oxidatively decomposed on pyrite. The metal ion released from the complex hydrolyzes to form a thin layer of metal oxide or hydroxide on the pyrite surface [17, 18]. In the previous papers, it was reported that the layer formed by CME treatment is hydrophilic and protective against oxidation; therefore CME can suppress the floatability and oxidation of pyrite [17, 18]. If CME can form a hydrophilic metal oxide/hydroxide layer on pyrite but not on coal, then it can be used as a depressant for pyrite in coal-pyrite flotation. Moreover, the proposed "Carrier micro encapsulation (CME) using tris-catecholato complex of Si_{4}^{+}\)" is industrially viable, easy to handle and economical. In the case of CME treatment, it is also expected that the pyrite tailing would not be oxidized to produce AMD after disposal in the tailing dam since pyrite is encapsulated with a protective coating of metal oxide or hydroxide. Therefore, as the proposed CME treatment is equally effective in suppressing pyrite oxidation in the tailing dam, the proposed technique will cut the huge amount of expanses of aftercare / treatment for AMD-acid mine drainage problem created by the air
oxidation of discarded pyrite.

In this study, effect of CME treatment (using tris-catecholato complex of \(\text{Si}^{4+}\)) on coal–pyrite separation is investigated by performing lab scale flotation test.

**Material and Methods**

*Minerals and solution preparation*

A specimen-grade pyrite (origin: Peru) was crushed and in a jaw crusher (Retsch BB51) and grounded in a ball mill, and sieved to obtain a 100-150 µm size fraction. Just before the experiments, the pyrite was washed following the procedures reported in the literature [19]: 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.

The raw coal (origin: Kushiro Coal Mine, Japan) was crushed to small pieces by a hammer and the impurities were removed by hand-picking. The coal particles were further crushed in a jaw crusher (Retsch BB51) and sieved to obtain a 100-150 µm size fraction. The coal samples were stored inside polyethylene plastic bags filled with nitrogen. The proximate and ultimate analysis of the coal sample was carried out with the equipment MICRO CORDER JM10, the result is shown in table 1(A) and 1(B).
The tris-catecholato complex of Si$^{4+}$ [Si(cat)$\text{3}^{2-}$] 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)$\text{3}^{2-}$. Solutions containing known concentrations of Si(cat)$\text{3}^{2-}$ were prepared by diluting the 5 mol m$^{-3}$ Si(cat)$\text{3}^{2-}$ solution with distilled water. The solution pHs were maintained by adding HCl or NaOH solution.

**CME treatment**

CME treatment was done in a 50 cm$^3$ Erlenmeyer flask containing 1 g of the mineral sample (1 g pyrite, 1 g coal, or 0.5 g pyrite-0.5 g coal mixture) and 10 cm$^3$ of Si(cat)$\text{3}^{2-}$. The flask was shaken in a water bath shaker under aerobic conditions at 25$^\circ$ C for a required treatment time. The shaking amplitude of the water bath shaker was 4 cm and the frequency was 120 min$^{-1}$. After the required time of shaking, the flask was removed from the shaker and the contents used for the dynamic bubble pick-up experiments and flotation experiments.

**Dynamic bubble pick-up experiment**

Dynamic bubble pick-up experiments [20] for the CME-treated pyrite or coal were conducted using an electronic induction timer (MCT-100). The CME-treated mineral along with the solution was transferred from the Erlenmeyer flask to a rectangular optical
glass cell to form a bed of particles. Inside the cell, an air bubble was formed at the tip of the capillary glass, and the bubble was brought into contact with the mineral particles at various contact times. The experiments were carried out at constant amplitude of the capillary motion for bubble pick-up, and the contact time between particles and air bubble was 1,000 ms unless specified. Ten bubble pick-up trials were conducted on different spots of the bed. The attachment of the mineral particles to the air bubble was observed with the help of a microscope connected to a computer display. The probability of attachment of the particles to an air bubble was calculated as the number of times the particle attached to the air bubble over the total number of bubble pick-up trials. The effect of pH, CME treatment time, and Si(cat)\(_3\)^{2-} concentration on the attachment of the mineral particles to an air bubble was investigated.

**Flotation experiments**

Lab-scale flotation experiments for pyrite, coal, and a coal-pyrite mixture were done using a Hallimond tube [21,22]. CME treatment was applied to the mineral using 0.5 mol m\(^{-3}\) Si(cat)\(_3\)^{2-} solution at natural pH for 1 h. The CME-treated mineral with solution was transferred to a beaker containing 100 cm\(^3\) distilled water and stirred for 2 min using a magnetic stirrer. Afterwards, 0.278 cm\(^3\) of a kerosene-water emulsion (kerosene content: 625 ppm) was added as a flotation collector, and stirring was continued for 5 min. The
mixture was transferred to the Hallimond tube, and flotation was conducted for 5 min by supplying air at a flow rate of 100 cm$^3$ min$^{-1}$. The froth and tailings were recovered by filtration, washed with distilled water, transferred into drying bottles, and dried at 105°C. From the weight of the dried froth and tailing products, the percentage recovery of the mineral in the froth was calculated. In the coal-pyrite mixture flotation experiments, pyrite and coal in the froth and tailing products were separated by using a dense medium (41 wt% sodium tungstate solution of specific gravity 1.5).

RESULTS AND DISCUSSION

Catechol (1, 2-dihydroxybenzene) is a bidentate chelating agent and has a remarkable affinity for metal ions in high oxidation states or with high charge/radius ratios. The studies involves catechol forming coordination compounds with silicon, tris-catecholato silicate complex ion or Si(cat)$_3$$^{2-}$ are well-documented in the literature [23, 24]. In the present study, catechol was selected as the organic carrier and Si as the metal ion to form tris-catecholato silicate complex ion or Si(cat)$_3$$^{2-}$ to form a SiO$_2$/Si(OH)$_4$ coating on the pyrite surface to prevent pyrite floatability and oxidation.

The dynamic bubble pick-up experiments for pyrite and coal with and without CME treatment was carried out to evaluate the effects of CME on the surface wettability
of these minerals. To optimize the conditions for CME treatment, the effect of the concentration of Si(cat)$_3^{2-}$, pH, and treatment time were investigated.

Pyrite and coal particles were treated in different concentrations of Si(cat)$_3^{2-}$ solution for 1 h at pH 7~9, and the effect of the concentration on the attachment probability of the mineral particle to an air bubble was investigated. As shown in Fig. 1, the attachment probability of pyrite to an air bubble decreased drastically with increasing Si(cat)$_3^{2-}$ concentration, and it became almost zero when the concentration was over 0.5 mol m$^{-3}$. The decrease in the probability of attachment was only observed for the coal at Si(cat)$_3^{2-}$ concentrations greater than 0.5 mol m$^{-3}$ (probabilities of attachment were over 0.9 at 0-0.05 mol m$^{-3}$ concentrations). At the highest concentration of 5 mol m$^{-3}$, the probability of attachment of coal to an air bubble decreased to 0.7. Based on this result, the concentration of Si(cat)$_3^{2-}$ for further experiments was kept at 0.5 mol m$^{-3}$.

The effect of CME treatment time on the attachment probabilities of pyrite and coal with and without CME treatment using 0.5 mol m$^{-3}$ Si(cat)$_3^{2-}$ was evaluated (Fig. 2). The attachment probabilities of pyrite particles shaken in distilled water (pyrite without CME) were over 0.8 at all times tested, showing that pyrite is hydrophobic. When pyrite was treated with 0.5 mol m$^{-3}$ Si(cat)$_3^{2-}$, the attachment probability became less than 0.1 after 1 h treatment, indicating that Si(cat)$_3^{2-}$ effectively converts the pyrite surface from hydrophobic to hydrophilic within a short time. The attachment probabilities of coal
without CME treatment were above 0.9 at all times tested. With CME treatment, the attachment probabilities of coal decreased with time and remained at 0.6 even after 24 h. This indicates that with long time CME treatment, the coal wettability becomes affected by the CME treatment but the hydrophilic effect is not as great as for pyrite.

Fig. 3 illustrates the effect of pH on the attachment probability of CME-treated pyrite and coal to an air bubble. The conditions were 0.5 mol m\(^{-3}\) Si(cat)\(_3\)\(^{2-}\) and 1h treatment time. The attachment probabilities of pyrite were less than 0.3 but the attachment probabilities of coal were higher than 0.8 over a wide pH range of 2-9. Especially in the pH range 4-9, the attachment probabilities of pyrite were zero and that of coal were over 0.9, indicating that CME treatment can be applied for separating pyrite from coal over a wide pH range.

Considering the results of the dynamic bubble pick-up experiments, experimental conditions of pH 7-9, treatment time of 1 hr, and Si(cat)\(_3\)\(^{2-}\) concentration of 0.5 mol m\(^{-3}\) were applied in the Hallimond tube flotation experiments. Figs. 4, 5, and 6 show the results for coal, pyrite, and the coal-pyrite mixture with and without CME treatment, respectively. Kerosene, a common flotation collector for coal, was used in the flotation experiments to evaluate the effect of CME treatment on pyrite and coal floatability in the presence of a collector.
Fig. 4 shows the effect of CME on coal recovery in the froth in the Hallimond tube flotation experiments. Without CME treatment, the coal recovery was 82% in the absence of kerosene the addition of kerosene increased the recovery and coal recovery was 89%. After applying CME treatment, more than 85% of coal was recovered in the presence and absence of kerosene. This shows that CME treatment induces no suppressive effect on coal flotation.

Fig. 5 shows the effect of CME treatment on pyrite recovery in the froth in the absence and presence of kerosene. Without CME treatment, pyrite recovery in the froth was 21% in the absence of kerosene and 72% in the presence of kerosene. With CME treatment, pyrite recovery in the froth was only 1% without kerosene and 8% with kerosene. This shows that CME treatment converts the pyrite surface from hydrophobic to hydrophilic, and the hydrophilic effect is strong as evidenced by the low recovery of pyrite in the froth even in the presence of kerosene.

Fig. 6 shows the effect of CME treatment on the flotation of coal-pyrite mixture in the presence of kerosene. Coal floatability was unaffected by the CME treatment while pyrite floatability was significantly suppressed: coal recovery in the froth was more than 90% with or without CME treatment while pyrite recovery in the froth was 60% without CME treatment and 18% with CME treatment. This indicates that CME treatment is able to suppress pyrite floatability selectively in coal-pyrite flotation.
The above results imply that a hydrophilic SiO$_2$ or Si(OH)$_4$ layer was formed on the pyrite but not on the coal. The selective layer formation is assumed to be due to the differences in the electrical conductivities of pyrite and coal: pyrite is a semiconductor but coal is an insulator. A hydrophilic SiO$_2$ or Si(OH)$_4$ coating is formed on the pyrite surface as follows [17, 18]: the Si(cat)$_3^{2-}$ complex adsorbs on the anodic site of the pyrite surface and decomposes into quinone and Si$^{4+}$ ion, according to:

$$
2
\begin{array}{c}
\text{Si} \\
\text{(cat)} \\
\text{3}^{2-}
\end{array}
= 2\text{Si}^{4+} + 6\text{O}_2 + 12\text{e}^{-}
$$ (1)

The electrons released are consumed by dissolved oxygen at the cathodic site of the pyrite:

$$
\text{O}_2 + 4\text{H}^+ + 4\text{e}^- = 2\text{H}_2\text{O}
$$ (2)

The Si$^{4+}$ ion released by the reaction in Eq. 1 hydrolyzes with water to form an SiO$_2$ or Si(OH)$_4$ layer on the pyrite surface:

$$
\text{Si}^{4+} + 4\text{H}_2\text{O} = \text{Si(OH)}_4 + 4\text{H}^+
$$ (3)

$$
\text{Si(OH)}_4 = \text{SiO}_2 + 2\text{H}_2\text{O}
$$ (4)
Summing up the anode, cathode, and hydrolysis reactions (Eqs. 1-4) give the overall reactions (Eqs. 5-6):

\[
\begin{align*}
\text{Si(cat)}_3^{2-} + 2\text{H}_2\text{O} + 4\text{H}^+ + 3\text{O}_2 &= 2\text{Si(OH)}_4 + 6\text{ }\text{benzoic acid} \\
\text{Si(cat)}_3^{2-} + 4\text{H}^+ + 3\text{O}_2 &= 2\text{SiO}_2 + 6\text{ }\text{benzoic acid} + 2\text{H}_2\text{O}
\end{align*}
\]

On the other hand, the Si(cat)$_3^{2-}$ complex does not decompose on the coal surface because coal is an insulator. As a result, no hydrophilic coating is formed on the coal surface thus coal floatability is not affected. Although the details of the above mechanism are not confirmed yet and further investigation is needed, the results in the present study show that CME using Si(cat)$_3^{2-}$ is effective in suppressing pyrite floatability in coal flotation.

**CONCLUSION**

The effect of carrier microencapsulation (CME) treatment using a Si-catechol
complex \((\text{Si(cat)}_3^{2-})\) on coal-pyrite wettability and floatability was investigated through bubble pick-up experiments and Hallimond tube flotation experiments. The results of bubble pick-up experiments showed that CME treatment changed the hydrophobic pyrite surface to hydrophilic but did not change coal’s hydrophobic surface over a wide range of conditions: pH 4-9, treatment times of 1-24 h, and \(\text{Si(cat)}_3^{2-}\) concentrations of 0.5-5 mol m\(^{-3}\). The Hallimond tube flotation experiments of a coal-pyrite mixture showed that pyrite floatability was selectively suppressed even in the presence of kerosene as collector.

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Figures and Captions:

Fig: 1 Effect of concentration of Si(cat)$_3^{2-}$ on attachment probability of pyrite and coal particle to air bubble in dynamic bubble pick up experiments. CME treatment time was 1 hr. pH of the Si (cat)$_3^{2-}$ was 7 to 9.
Fig: 2 Effect of CME treatment time on attachment probability of pyrite and coal particle to air bubble with/without CME in dynamic bubble pick up experiments. Si(cat)$_3^{2-}$ concentration was 0.5 mol m$^{-3}$, pH of the Si(cat)$_3^{2-}$ was 7 to 9.
Fig: 3 Effect of pH on attachment probability of pyrite and coal particle to air bubble in dynamic bubble pick up experiments. Si(cat)$_3^{2-}$ concentration was 0.5mol m$^{-3}$. CME treatment time was 1hour.

![Graph showing effect of CME treatment on coal recovery in froth in presence or absence of kerosene.]

Fig: 4 Effect of CME treatment on coal recovery in froth in presence or absence of 0.278 cm$^3$ of 625 ppm kerosene. The Si(cat)$_3^{2-}$ concentration: 0.5 mol m$^{-3}$, pH: 7-9, CME treatment time: 1 h.
Fig: 5 Effect of CME on recovery of pyrite in froth in the presence or absence of 0.278 cm$^3$ of 625 ppm kerosene. The Si(cat)$_3^{2-}$ concentration: 0.5 mol m$^{-3}$, pH: 7-9, CME treatment time: 1 h.
Fig: 6 Effect of CME treatment on flotation of the coal-pyrite mixture in the presence of 0.278 cm$^3$ of 625 ppm kerosene. The Si(cat)$_3$-conc: 0.5 mol m$^{-3}$, pH: 7-9, CME treatment time: 1 h.

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Table:1 (A) Proximate analysis of the Kushiro coal sample
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Table: 1 (B) Ultimate analysis of the Kushiro coal sample