Interactions between Kaolinite and Molybdenite in Fresh and Seawater

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Abstract
Scarcity of fresh water, particularly for operations situated in desert areas, has forced the copper-molybdenum mining companies to look for new sources of water such as seawater. The use of this saline aqueous medium in the process of flotation leads to some important effects on the efficiency of this process. Besides, copper grades of large copper deposits have constantly decreased during the last decades, which is typically associated to high concentrations of clay minerals in the ores which is known to negatively affect the flotation process. The objective of this research was to evaluate the effect of kaolinite on the floatability of molybdenite at different pH conditions, using two types of solutions (0.01 M NaCl solution and seawater). Floatability of molybdenite particles was assessed through micro-flotation tests in a 150cm³ Partridge-Smith glass cell and using nitrogen (N₂) at a flowrate of 20cm³/min, for a total flotation time of 10 minutes. Flotation feed was prepared mixing known amounts of molybdenite with particles of kaolinite. The results of this study indicate that molybdenite floatability slightly decreased under the presence of fine particles of kaolinite when conventional water was used in the tests. However, in seawater the negative effect of kaolinite was more relevant, in particular at pH>9, which suggests that the presence of cations in solution intensify the phenomenon of heterocoagulation between fine kaolinite particles and molybdenite (slime coating). It is assumed that the stronger depressing effect of kaolinite at pH>9 could be related to the formation of hydroxy-complexes of magnesium and calcium.

Introduction
Molybdenite (MoS₂) is the primary source of molybdenum. From a structural point of view, molybdenite particles are formed by crystal layers of S-Mo-S which are held together by the action of strong covalent bonds. The interaction between each of these S-Mo-S crystals occurs through the S-S planes (Figure 1), and the process of breakage of the molybdenite particles occurs along these planes. As a result, a non-polar and highly hydrophobic surface which is usually referred to as face, is created. Consequently, molybdenite particles are usually described as anisotropic entities with hydrophobic faces and hydrophilic edges [2]. It is known that the zeta potential of molybdenite particles is the lowest at pH values ranging between 5.5 and 6.5, which is in agreement with the range of maximum floatability [2]. Regarding the interaction between molybdenite and gangue particles, previous studies showed that the presence of Ca²⁺ ions in solution induces heterocoagulation between silica and molybdenite which explains the low molybdenum recoveries observed under these conditions [3,4]. The negative effect of silica on the floatability of molybdenite was assumed to be related to a phenomenon of slime coatings [4].

Figure 1: Crystalline structure of molybdenite Castro et al. [1].
Molybdenite is commonly associated with copper. The grades of copper-molybdenum deposits in the world have declined, which is typically associated with high concentrations of phyllosilicates, with the clay minerals being the most abundant group. These types of mineral species are usually dispersed in mineral slurries as fine particles, which generates the conditions for non-Newtonian behavior [5]. These characteristics affect the entire mineral processing chain, i.e., milling, froth flotation, thickening, and dewatering [6-12]. The efficiency of the froth flotation process can be severely affected by the presence of high contents of phyllosilicates, and in many cases this problem can be dealt with successfully by diluting the slurry, blending different ore types, and sometimes even disposing of the problematic ores which inevitably leads to losses of valuable minerals [13]. In particular, clay minerals are the most common phyllosilicates found in copper ores and kaolinite is one of the most abundant species [14]. From the surface chemistry point of view, kaolinite has two different surfaces generated during particle breakage, i.e., the basal silica-like and the alumina-like faces [15,16]. The silica-like faces carry a negative charge while the alumina-like displays a pH dependent surface charge resulting from protonation/deprotonation of exposed hydroxyl groups with IEP values around 7.0 [17]. Some studies also suggested that the silica-like faces display hydrophobicity and that dispersion of kaolinite particles in water is explained by the presence of surface imperfections on the faces which create polar micro-edges [18,19]. The anisotropic behavior of clay minerals promotes the attachment of these fine particles to minerals (slime coating), and their interactions with reagent and bubbles. All these phenomena reduce the copper and molybdenum floatability and hence recovery [6,13,20,21], and cause major changes in the rheological behavior of mineral slurries [22].

Nowadays, water is a critical and scarce resource for the mining industry and the use of seawater to process copper-molybdenum ores by flotation seems to be a sustainable solution in zones with restricted resources of fresh water such as northern Chile [23]. Seawater contains significant concentrations of different ions (Na+, K+, Mg2+, Ca2+, SO42-, Cl−) which can potentially adsorb on the solid surfaces and induce heterocoagulation among molybdenite and clay minerals. Earlier publications from the CRHIAM Center of the University of Concepcion, on flotation of Cu-Mo sulfide ores in seawater [23,24] indicated that molybdenite flotation is strongly affected in seawater when pH is raised with lime to depress pyrite, which is related to the depressing effect of Mg2+ and Ca2+ hydrolysis products on the hydrophobicity of molybdenite. These studies showed that molybdenite floatability can be severely depressed by the action of the products of magnesium hydrolysis, in particular Mg(OH)2 and that in the case of Ca2+ the depressing action can be related to the action of Ca(OH)+. Recently Uribe et al. [22] showed that the depressing effect of kaolinite on chalcopyrite may be related to the formation of hydrolyzed species of calcium and magnesium which can induce heterocoagulation between kaolinite and chalcopyrite (slime coating). Liu et al. [10] also showed that calcium enhanced heterocoagulation between montmorillonite and bitumen can lead to low bitumen recoveries. The objective of this investigation was to evaluate the depressing effect of kaolinite on the floatability of molybdenite using two types of solutions, i.e., a 0.01M NaCl solution, seawater, and solutions containing Ca2+, Mg2+, Na+ and K+ at the concentrations at which they appear in seawater, over the pH range from 8 to 11.

Figure 2: Particle size distribution of molybdenite samples.

Methodology

Samples and reagents

The kaolinite sample was provided by the Clay Minerals Society Source clays. XRD analyses (Bruker D4-Endeavor operated with Ni-filtered Cu radiation) indicated that the sample had 96% kaolinite, 3% anatase, and 1% of other traces. Cation exchange capacity (CEC) and Brunauer-Emmett-Teller (BET) surface area were 0.7 cmol/kg and 180 m²/g respectively. The molybdenite sample used in this work was obtained from an industrial molybdenum concentrate, and further purified through 3 stages of flotation without collector. Samples of two particle size distributions were used in this research, i.e., coarse and fine samples. The coarse sample with a P80 of 100 microns corresponded to the original molybdenite samples obtained from the process of purification, while the fine sample with a P80 of 50 microns was obtained by hand grinding the...
coarse sample. Figure 2 shows the particle size distributions. The samples were then washed and cleaned following the procedure described by Uribe et al. [22] which considers 3 stages of washing with sodium hydrosulphide (NaSH) and acetone in order to remove any organic reagents such as collectors present in the sample. Table 1 shows the chemical analysis of the samples. In order to keep molybdenite oxidation at minimum, the samples were stored in a freezer in sealed plastic bags previously blown with nitrogen, at a temperature of about 1 °C [25-27].

Table 1: Chemical analyses of molybdenite samples.

| Sample | Cu(%) | Fe(%) | S(%) | Mo(%) | Ins(%) |
|--------|-------|-------|------|-------|--------|
| Coarse | 0.55  | 0.49  | 37.9 | 55.4  | 5.3    |
| Fine   | 0.53  | 0.39  | 38.0 | 55.8  | 5.1    |

Methyl Isobutyl Carbinol (MIBC) of analytical grade was used as frother at 15ppm, and potassium amyl xanthate (PAX) as collector at 25ppm for the micro-flotation tests. It has to be noted that PAX was purified by dissolution in acetone, and precipitated with ether. Analytical grade NaCl, KCl, MgCl₂ and CaCl₂ (Merck) were used to prepare the solutions. Distilled water with an electrical conductivity of 21.5μS/cm was used in all the experiments. Seawater samples were taken from the Lenga Coast in the VIII Region of Bio in Chile. The pH was adjusted with sodium hydroxide and hydrochloric acid (Merck).

Micro flotation

The floatability of molybdenite was evaluated through micro-flotation tests that were carried out in a 150mL Patridge and Smith glass cell using nitrogen (N₂) at a flowrate of 80mL/min (Figure 3). The flotation feed was prepared by mixing 1g of molybdenite (coarse and fine) with 0.15g of kaolinite to then prepare a suspension using 100mL of solution (0.01M NaCl or seawater). After 3 minutes of conditioning at the required pH, MIBC and PAX were added at a concentration of 15ppm and 25ppm respectively (based on the 150mL of the micro-flotation cell) and conditioned for 5 more minutes at the previously established pH value. Additional solution was then added to complete the 150mL required for the micro-flotation glass cell. Finally, the gas valve was opened to start the process of flotation, which was carried out for 3 minutes, pulling froth every 10 seconds. Pulp level in the micro-flotation cell was kept constant by adding a background solution prepared at the same composition and pH of the original solution. Finally, in order to separate molybdenite and kaolinite particles, the concentrates and tailings were sieved using a 400# screen (38 microns) and dried in an oven at 105 °C for 5 hours. This way it was possible to measure the mass of molybdenite in the concentrate, and the solids flotation yield was calculated as the ratio between the mass of molybdenite in the concentrate and the mass of molybdenite in the concentrate tailings. All testing was done in duplicate.

Induction time measurements

Induction time measurements were conducted using the procedure described by Castro et al. [24] and the induction time meter built at the University of Alberta. According to this procedure, a fresh bubble held on a capillary is pushed downward through a solution so that it is kept in contact with a bed of particles for a measured and controlled contact time span. Then, the bubble is lifted upwards to its initial position and observed through a microscope in order to determine whether attachment occurred during the contact period or not. The process is repeated 10 times at different positions on the particle bed and the number/percentage of successful contacts is determined. Then, the contact time is changed to different values and the procedure is repeated again in each case. The final result of the test is a plot of percentages of successful contacts (Nsc) versus the measured contact time (tc).

In this work, the induction time was obtained by determining the contact time at which 50% of the contacts were successful. The experimental data were fitted using Equation 1.

The beds of particles were prepared by mixing 3 grams of molybdenite (for size) with 30mL of solution containing 40ppm of PAX. Immediately after, fine particles of kaolinite were added at a concentration of 13% (solid base). Then, the pH was adjusted to the required value and the slurry was conditioned for 12 minutes, after which it was transferred to a 15mL cell in order to start the induction time measurements. It has to be pointed out that no frother was added in these experiments and that air was used to generate the bubbles. All these experiments were conducted in duplicate thus the results presented in this paper are average values. It has to be noted that these types of measurements were previously used to evaluate the effect of slime coating on the floatability of molybdenite in the presence of clay minerals [28] and different ions.

Results and Discussion

Micro flotation

Figure 3 shows the recovery of coarse molybdenite in 0.01M NaCl solution and in seawater, in the presence and absence of kaolinite. The results indicate that the effect of kaolinite on the floatability of molybdenite at the whole range of pH is minor when the process of flotation is carried out using a 0.01M NaCl solution. In contrast, in seawater the effect of kaolinite on molybdenite recovery seems to be stronger, in particular at pH>9. In other words, the depressing effect of kaolinite seems to be important only in an aqueous media of high ionic strength such as seawater. The theory of stability of colloidal systems developed by Derjaguin-Landau-Verwey-Overbeek (DLVO) can be used to better understand the interactions between kaolinite and molybdenite. The DLVO theory considers the existence of two forces that affect the process of aggregation of colloidal particles, i.e., van der Waals forces of attraction and repulsive forces generated by the interactions of the electrical double layers. Thus, to explain the results presented in Figure 3, it is necessary to analyze the effect of pH on the surface charge of both minerals. The surfaces of the faces and edges of molybdenite particles are expected to be negatively charged at pH values between 7 and 11 [2]. Besides, previous studies reported that the surface charge of the alumina-like faces of kaolinite becomes close to zero at pH values around 7-8 [15,16]. Then, according
to this information heterocoagulation between kaolinite and molybdenite should be more likely at lower pH, however the data presented in Figure 3 indicate that there is not a clear correlation between pH and the depressing effect of kaolinite on the floatability of molybdenite which suggests no heterocoagulation.

Figure 3: Recovery of coarse molybdenite using conventional and seawater with and without kaolinite.

When looking at the results for seawater in Figure 3, it can be seen that the effect of kaolinite is more significant in seawater than in fresh water at the whole range of pH, but at pH>9 the depressing effect of kaolinite becomes more important. The fact that the pH value at which the trend changes is 9, suggests that the stronger depressing effect of kaolinite observed at this pH range is related to the formation of hydroxy-complexes of magnesium and calcium and their influence on the interactions between kaolinite and molybdenite. Previous studies showed that the first hydroxy-complex of magnesium Mg(OH)\(^+\) starts forming at pH 9, reaching a maximum concentration at pH 10.5. At this pH, positively charged precipitates of Mg(OH)\(_{2}\)(s) can also form which can affect the properties of the air/liquid and solid/liquid interfaces. In the case of calcium, the first hydroxy-complex Ca(OH)\(^+\) forms around pH 9.5-10 with the maximum concentration obtained at pH values around 13-13.5. The presence of these hydroxy-complexes seems to enhance the interactions between clay particles and molybdenite, phenomenon referred to as "slime coating". The induction time is the time required to break the liquid film existing between a bubble and a particle after the two collide. Therefore, high induction time values indicate that the particle-bubble adhesion is favorable and therefore, better molybdenite recoveries should be expected.

Figure 4 shows the induction time data obtained using coarse molybdenite in conventional and seawater, with and without kaolinite. The results of induction time are in good agreement with the micro-flotation data. Firstly, it can be seen that in general the induction time values are higher in the presence of kaolinite in both fresh and seawater. Additionally, the depressing effect of kaolinite is more important in seawater, in particular at pH>9, which is again in perfect agreement with the data of micro-flotation. Therefore, the induction time results validate the conclusions obtained in the micro-flotation testing.

Figure 4: Induction time of fine molybdenite using conventional and seawater with and without kaolinite.

Figure 5 shows the recovery of fine molybdenite using conventional and seawater, with and without kaolinite. The first observation that can be made is that in general the recovery values obtained in a 0.01 NaCl solution using the fine molybdenite sample tend to be lower than those obtained with the coarse sample. This is also observed in the results obtained using seawater at pH 7-9. This can be related to the fact that the face/edge ratio decreases with particle size, which has an effect on hydrophobicity leading to low values of recoveries. However, when comparing the recovery of molybdenite at pH>9, it can be seen that the
The depressing effect of kaolinite on the floatability of molybdenite at pH values between 7 and 11 is less important when the process of flotation is carried out using a 0.01M NaCl. In seawater the depressing effect of kaolinite seems to be stronger, in particular at pH>9. The fact that the pH value at which the trend changes is 9, suggests that the stronger depressing effect of kaolinite observed at this pH range can be related to the formation of hydroxy-complexes of magnesium and calcium and their influence on the interactions between kaolinite and molybdenite. Hydroxy-complexes of magnesium and calcium can induce attachment of kaolinite to the faces and edges of molybdenite through a mechanism of surface charge neutralization [1,29]. As the face/edge ratio increases with particle size, heterocoagulation between kaolinite and molybdenite faces is more likely for coarse sizes, and that could explain why kaolinite affects more the recovery of coarse molybdenite at pH>9. These conclusions were validated through induction time measurements.

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