Talc Flotation

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Talc is a naturally hydrophobic gangue mineral in most sulfide ores. However, talc has vast applications in the cosmetics, paper, and paint industries due to its high chemical stability, and its demand continues to grow. Since flotation is the most effective beneficiation technique for upgrading sulfides, the high hydrophobicity of talc has made its selective separation challenging.

surface properties talc froth flotation talc depression talc hydrophobicity

1. Introduction

In nature, talc is commonly associated with different minerals such as carbonates (serpentine shear zones), silicates, and sulfides [1][2][3][4][5][6][7]. Talc has a different chemical composition and crystal morphology based on the origin, which may impact its mineral specifications and features (whiteness, particle shape, oil absorption, and amphiphilic specification) ^[2]. According to these precise features, talc has specific properties (platy, softness, hydrophobicity, organophilicity, and inertness), which can provide different functions in many varied applications (**Table 1**). Talc has high chemical stability, high surface affinity, and heat resistance compared to other silicate minerals $^{[8]}$. Talc world production is documented and shown in the graph (**Figure 1**).

Table 1. Common talc applications.

Talc with a theoretical chemical composition, Mg $_3$ Si $_4$ O $_{10}$ (OH) $_2$, is the softest mineral on Earth, mainly attributed to its chemical structure. The magnesium-based octahedral layer is in between silica rings through shared oxygen atoms. This exposes oxide surfaces that make talc only held together by weak van der Waals forces. Furthermore, these oxide surfaces make talc naturally hydrophobic. When hydrolyzed in water, the magnesia edges will show some hydrophilicity ^[10]. Talc has two faces: the hydrophobic basal surface and the hydrophilic edge surface ^[11]. The majority, around 90% of its total surface area, is a basal surface that accounts for the natural hydrophobicity or

floatability of talc [5][6][7]. Ergo, preventing talc (as a typical gangue for sulfide ores) from being recovered to the flotation concentrate was the primary focus of the earlier studies.

As mentioned, valuable sulfide minerals such as chalcopyrite are commonly intergrown with gangue minerals such as talc [1][2]. This is generally the challenge in the selective flotation of such minerals. In other words, the talc separation from other valuable sulfide minerals can be difficult since talc has good natural floatability. The existence of talc in the sulfide flotation concentration as a MgO-containing gangue can cause problems in the downstream metallurgical processes ^{[<u>8][10][9][11</u>]. In detail, talc has several negative effects on furnace operation in} pyro-metallurgical processes to extract various metals, especially from sulfide minerals. In the smelters, MgO can change the properties of the primary slag and reduce the strengths of the sinter and pellets. The primary slag with high MgO being sticky, and the softening-melting properties of the burden in the blast furnace becoming deteriorated. In other words, increasing talc/MgO in the feed of smelters can increase liquidus temperature and slag viscosity, which requires operating the furnace at higher temperatures, thus shortening the campaign life of the furnace. To avoid these negative effects, talc content has to be significantly reduced in the ore concentrate from mineral processing plants as the feed to smelters [2][10][12][13].

These problems result in a cost increase and recovery reduction of subsequent smelting operations [10][14]. Thus, it is of great interest to remove talc beforehand from the processing plant concentrate. Different kinds of depressants have been reported to inhibit talc floatability, and polysaccharides are commonly used. However, these are expensive and can cost relatively more than other reagents used in the flotation. Although these depressants are able to depress talc, they have poor selectivity. For example, Beattie et al. reported that CMC, as a talc depressant, depressed not only talc but also chalcopyrite ^[15]. Such depressants also impair the flotation performance of other valuable minerals ^[16]. Some investigations have found that using ultrasonic treatment can further improve the depression of talc. It was shown that it had improved the flotation performance, specifically in terms of flotation rate, overall recovery, and selectivity of sulfides [16][17].

All these issues indicated that it would be worthwhile to conduct a comprehensive review that analyzes the various conditions of talc flotation. The purpose of this work is to highlight and classify the investigations that were conducted on talc flotation, defining sub-areas of studies, and highlighting the areas that require further exploration for better understanding and optimizing of talc flotation. To the best of our knowledge, this is the first review to offer an in-depth analysis and critique of talc flotation studies, including experimental works and detailed analysis procedures suggested in the literature. The literature analysis showed that while talc depression through flotation is extensively studied, insufficient attention is paid to its flotation kinetics and application of column flotation. It is also highlighted that the literature available on the talc column flotation is rare. The lack of fundamental understanding of pretreatment on the flotation efficiency is discussed, which is essential for the separation process.

2. Talc Surface Properties

2.1. Relative Humidity

Talc, the oldest known solid lubricant, has a low shear strength ^[6]. Different factors influencing the hydrophobicity of talc are elongation, flatness, roundness, and relative width. A talc particle with higher elongation and flatness and lower roundness and relative width will tend to be more hydrophobic since, in these cases, the adhesion force is stronger due to more areas and lines for contact ^[18]. These different surface properties of talc can be attributed to its unique molecular chain in the faces and edges (**Table 2** and **Table 3**). Its crystal has three layers, with the two outer layers being composed of Si-O tetrahedra, while the layer in between is a brucite Mg(OH)2 layer or hydroxyl octahedral, bonded by van der Waals forces [4][5][6][7][8]. These two outer layers have zero residual charges, which means no interlayer cation is present ^[19], while the octahedral layer has charged ions Mg2+, OH- that draw many bonding possibilities with the water molecules^{[5][7][10]}. Moreover, -OH groups of the octahedral layer are known to be influencing the surface behavior of talc ^[20]. In its natural state, talc adsorbs organic and inorganic species on its surface. The highly energetic sites on its surface are mostly blocked by nitrogen and water molecules ^[21]. **Table 2.** Effect of the molecular chain in the talc surface and surface properties.

Table 3. Properties of surfaces formed after cleavage.

Relative Humidity (RH) can be considered an essential factor for measuring the talc surface's hydrophobicity. Strong binding sites on the talc surface lead to water adsorption at low RH [34,35]. However, at the point of saturation, the talc surface is dominated by water-water interaction (cohesive interaction), which consequently reduced the water and the strong binding sites interaction (adhesive interaction), rendering the surface hydrophobic. On the other hand, at high RH, hydrophobic faces consisting of Si-O-Si groups show low interaction with water (cohesive interaction > adhesive interaction). Therefore, the electrokinetic properties of the talc could be affected by Hydropholic Sites ratio [32]. This can be translated as the surface face ratio has a direct relationship with the level of talc hydrophobicity edge (Figure 2) [20,36] and has a positive trend with the surface wettability of talc [23,37]. A smaller edge-to-face surface ratio would result in a lower measured zeta potential since fewer charged surface groups per unit surface area [35,38].

Figure 2. (**A**) The relation between the size of particles and surface/edge ratio (**B**) force of cohesion, work of cohesion, and hence surface tension.

Thus, due to talc's delamination that produces more surface faces, the hydrophobicity increases by decreasing the particle size ^[28]. Beyond a certain particle size reduction limit (i.e. after subjecting a 10 microns-sized talc in an hour of high-intensity planetary ball milling), the hydrophobicity was seen to decrease (inferred to the loss of crystallinity in the talc and structural degradation) ^[29]. As examined, the hydrophobic face and hydrophilic edges constitute around 90% and 10% of the total surface area, respectively ^[11]. These properties also made a significant surface roughness effect on the talc hydrophobicity. A cleaved talc surface is considered to have a very low root mean square roughness, which indicates a very smooth surface. This cleaved surface becomes very hydrophobic, as manifested in the contact angle measurement [12][30][31][26][32][33].

2.2. Zeta Potential

Exploring electrical charge on the talc surface in various conditions indicated that its zeta potential (ZP) magnitude is negative at wide pH ranges (pH 2–12) $\frac{[18][34][35][36][11][21][37]}{[11][21][37]}$. This may be attributed to the Mg-OH and Si-OH groups ionization, passing of H+ into the water, and OH ions in water adsorbed on Si2+ and Mg+ ^[38]. Different studies reported the isoelectric point (IEP) of talc occurred at around pH 2–3 (**Table 4**). Furthermore, in phyllosilicate minerals such as talc, the IEP may depend on the mineral side where the measurement is made (the edge vs. the face) ^[14]. The hydrophobic faces consist of –Si–O–Si– groups, have a very slight electrical charge and are nonpolar in water ^{[29][39]}. On the other hand, the edges (constructed by breaking ionic/covalent bonds and composed of silicon, hydroxyl ions, oxygen, and magnesium ions ^[24]) are polar in water. They indicated a relatively high electrical charge, depending on pH [11][39].

Table 4. The talc IEP of the absence and presence of reagents.

Different reagents have various impacts on the talc electrical surface charge (**Table 4**). It was found that the addition of depressants such as tragacanth gum, guar gum, and carrageenan slightly decreases the absolute value of talc's potential at a specific pH range [18][35][11][25][40]. Depressant adsorptions could consequently shift the slip

surface electrical double layer towards the outside and cause this decrease. On the other hand, the talc IEP in the presence of depressants just increased negligibly. At a very low and very high pH, the addition of some depressants has an insignificant effect. This is attributed to the protonation of, for example, carboxyl groups (for gum Arabic (GA) at pH 2–4), which decreases surface charge; thus, talc's ZP became less affected ^[41]. These carboxyl groups will mostly dissociate at a high pH (pH 10–12) and enhance the electrostatic repulsion between GA and talc. It was also found that the presence of chloride-based metal salts, such as NaCl, decreased the ZP of talc but did not have a significant change in IEP $^{[42]}$. With the presence of an electrolyte such as Ca2+ in addition to depressant CMC, zeta potential is highly affected by pH changes since the electric potential is almost closer to zero in Ca2+ solution [43]. For the cases of leached talc, ZP remains the same, while calcinated talc shows a significant shift. The latter has become more negative, attributed to the change of brucite to MgO and loss of structural water from talc ^[44].

2.3. Contact Angle

Measuring the talc's contact angle in air, water, and the number of organic solutions, indicated that its surface is not completely hydrophobic ^[45]. Various measurements and bulk/surface analysis had been applied to assess the talc contact angle and hydrophobicity for the gross content and cleavage specifications ^[46]. Assessment of these measurements demonstrated that the talc mono-crystal surface is hydrophobic; however, as mentioned, they indicated hydrophilic behavior at low relative humidity ^[44]. In other words, particle size could influence the contact angle. Since smaller particle sizes would have a larger edge-to-face ratio ^[32], the contact angle could also be smaller for these particles than larger ones. However, with the depressants' presence, the contact angle might be lower for larger particles as the depressant adsorption density could be higher compared to smaller particles (**Table** 5) ^[47]. Thus, the talc contact angle is correlated to the surface coverage, and its surface wettability is influenced by the effect of polymer adsorptions ^[48]. It would be decreased (hydrophobicity increased) when cationic surfactants and different depressants (dextrin, starch, humic acid, and combination of calcium chloride and sodium lignosulfonate) were presented at their sufficient concentrations [4][12][30][32]. The extent of aluminum substitution or impurity in the tetrahedral silica layer could decrease the talc contact angle [19][42][49]. The talc contact angle would not vary significantly with pH variations and instead remained high, owing to the talc's natural hydrophobicity ^[50]. It was documented that the static contact angle measurements might not be very reliable due to talc's possible stickslip behavior at the wetting front that could result in hysteresis in advancing and receding contact experiments ^[3] . [24]

Table 5. Contact angle measurement of talc in the different liquid mediums.

3. Talc Flotation

3.1. Collectors

Talc shows excellent floatability in a wide pH range $(3-11)$ $[16]$. Although talc can be considered a naturally hydrophobic mineral, it might need treatment to float. Investigation on pure talc flotation showed that it could float easily ^{[<u>18][33</u>] in a collectorless flotation system by adding frothers (alcohols) ^{[28][51][52][53]}. However, various} collectors could be considered for the talc selective flotation separation in complex ores ^[54]. When using xanthate as a collector, Atomic Force Microscope (AFM) and Fourier-transform infrared spectroscopy (FTIR) test results indicated that the collector was physically adsorbed on the talc surface, and adsorption would be negligible. For example, because the potassium amyl xanthate (PAX) did not adsorb onto talc, no competition was detected through the flotation in the adsorption of this collector depressant ^[39]. In addition, the absence and presence of potassium butyl xanthate (PBX) over the whole pH range did not affect the talc recovery ^[55]. Among anionic collectors, it was observed that increasing the oleic acid concentration created the multilayers of the collector on the talc surface and reduced its recovery ^[56]. Cationic collectors (Tallow amine acetate and Dodecylamine) could improve its floatability [5][57]. The adsorbed amount of collector affected the talc recovery, where a high concentration made the talc surface more hydrophobic ^[57]. It should be observed that adsorption capacity would be dependent on pH, particle size, temperature, adsorption time, and collector concentration ^[5]. It was reported that without a collector, the talc flotation does not follow the first-order kinetics model but follows the Kelsall model. With collector, it follows both models since the talc morphology had less effect on the flotation performance on minerals with naturally high floatability [58].

3.2. Depressants

The faces of talc crystals consist of quite compensated oxygen atoms, and the edges of them have consisted of hydroxyl ions, silicon, oxygen, and magnesium ions. Since the faces of talc crystals have a very slight electrical charge and are nonpolar in water, the depressant adsorption is expected to have occurred on a greater area of the faces. On the other hand, the edges of talc, which are simply hydrolyzable, have a high electrical charge and are polar in water. Thus, the large particles had higher depressant adsorption than the finer ones, resulting in a lower

flotation recovery ^[32]. Therefore, particle size and the face-to-edge ratio was still considered the main factor for the depressant adsorption [44].

Some investigations documented that the adsorption of Dextrin WY and Polymer-H onto the silica models followed the Freundlich isotherm, which indicates a heterogeneous surface ^[59]. However, the later studies showed that the adsorption obeyed the Langmuir isotherm, indicating a homogeneous surface ^[33]. The adsorption isotherms of guar gum and dextrin onto talc followed the Langmuirian behavior and pH-independent. In general, it was also shown that the Langmuir adsorption isotherm fit the data better than the Freundlich model, except when the pH was under 2 (Table 6) ^[60]. These indicated that the adsorption mainly occurred in the dominant basal face of the talc ^{[<u>8][51</u>]. However, talc edge could also play a role in the adsorption of polysaccharides ^{[11][61]}. In general, there is} a significant relationship between adsorbed amount, the adsorbed morphology, layer thickness, depressant molecular weights, and the subsequent talc wettability. For some depressants, such as carboxymethyl cellulose (CMC), the enhancement of acidic pH and ionic strengths of the suspension medium increased the adsorption isotherms of the depressant on talc ^[62]. Adsorption isotherms of the polymers indicated that by increasing the concentration, the talc recovery decreased [23][63].

Table 6. Investigation of adsorption isotherms for different polymers on talc.

make talc faces hydrophilic ^[64]. The other depressants are CMC (a negatively charged polymer) [3][7][66][43][45][67], guar gum ^[68], and dextrin ^{[3][7]}, which also had various degrees of depression on other minerals ^[68]. Polymer structure and molecular weights are effective factors on the talc depressant adsorption. For example, the high molecular weight of starch leads to a greater adsorbed amount and apparent layer thickness than its low molecular weighted polymers, which decreased the talc wettability ^[33]. Because of the more favorable cis-configuration of hydroxyl groups in guar gum and its higher molecular weight, it had higher adsorption densities onto talc in comparison with dextrin (Figure 3) [16][20][41][44][69][70][71]. Between CMC and guar gum, the molecular weight of the latter has a more significant effect on talc depression [25].

Figure 3. Adsorption of different depressants on the talc surface.

As mentioned, selective talc depression is complicated. For instance, flotation experiments have shown that the P(Am-co-VPD) copolymer system depressed talc and pentlandite in mixed mineral flotation. However, the copolymer system depressed the talc better than guar gum. From an absorption point of view, the low selectivity of depressants could be attributed to the little variation of the Langmuir affinity constant of these two minerals ^[72]. The

effect of various types of depressants on the flotation of talc was summarized in **Table 7**. Other factors such as calcination also affected the adsorption density of guar gum onto talc. Calcination decreased the adsorption density for talc particles. This decrease was attributed to the desorption of nitrogen species found on the talc surface ^[20].

Table 7. Depressants and their effect on the talc flotation.

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frother adsorption capacity and is more efficient in terms of weight comparing pine oil and Aero 825 while obtaining the highest talc recovery. In collectorless flotation, the addition of MIBC alone could recover 90% of the talc in a micro-flotation test in a wide pH range (acidic to alkaline conditions) ^{[55][75]}. This increases relatively and depending on the molecular structure, which could be considered for different frother types (**Table 8**).

It is well understood that through flotation separation, the bubble size strongly depends on frother type and its concentration. Kho et al. (1989) investigated the effects of bubble size on the talc recovery in the column flotation system. The simulation results demonstrated that the air bubble diameter was the most significant parameter to increase talc recovery, where they had a negative correlation (**Figure 4**). Hence, the control of bubble size would be an essential factor in the talc flotation. Simulation through a column flotation indicated that bubble size significantly influences talc flotation recovery, although increasing the column height could also improve its recovery [76].

Figure 4. Effects of bubble size on the recovery of talc in the different column flotation (L/D: Length/Diameter) ^[76]).

3.4. pH Influence

Although the talc floatability is pH-independent, a negative trend between coagulation of talc suspension and pH could be observed ^[77]. It should be observed that the surfaces of talc particles are net-negatively charged [69][78], and when the pH is enhanced, the zeta-potentials of the talc surface decreased ^[79]. At pH 9.4, the solubility of talc is at its lowest level; by reducing pH, the negative charge of talc was reduced, and its solubility increased [42][24][80] [81]. Thus, talc floatability could be altered by pH ^[68]. At neutral pH, talc has the highest natural floatability, and in both acidic and alkaline solutions, its floatability is reduced to various degrees ^[Z5]. The adsorption mechanisms of talc are different at various pH values ^[65]. The potential- specifying ions including H⁺ and OH⁻ of talc mineral are enhancing pH of the solution and, as a result, reducing the adsorption capacity ^[5]. Ionic strength and pH of the solution indicated some effects on the depressant effectivity [ZIBI[14][70], although it is mainly suggested that depressant adsorption is independent of pH as the surface charge has a negligible effect ^[44]. It was shown that at neutral or moderately alkaline pH values (pH 7–9.3), the lignosulfonates would be weak talc depressants; however, their depressing ability significantly enhance under highly alkaline conditions when lime is applied as the pH regulator ^[78]. Adsorption studies indicated that at pH 3 and pH 9, the adsorption of chitosan is through physical interactions, hydrogen bonding. At pH 9, aside from adsorption due to hydrogen bonding, chitosan changed the potential of talc gradually to zero. In the mixed mineral flotation of chalcopyrite and talc, chitosan was seen as a selective depressant of talc at pH 3 due to the presence of more hydroxyl bonds on talc than chalcopyrite but

unselective at pH 9 as chitosan gets deposited on both minerals $[14]$. At pH 9 with low ionic strength (0.001M KCI), the carboxyl groups of CMC dissociated; thus, electrostatic repulsion occurred between the negatively charged talc and CMC, which prevented the adsorption [11]. The pH also affected the adsorption, wherein CMC was protonated at acidic pH. At pH 3.5, the negative charge of talc was not significant; therefore, electrostatic repulsion is reduced, enhancing the talc depression with CMC [Z][8][48][82].

3.5. Ions Influence

By dissolution of talc, the neutral or charged magnesium hydroxo complexes are liberated and, through hydrogen bonding or the chemical interaction, could react with depressants at the hydroxylated talc surface and in the bulk solution [5][17][32]. Hence, the presence of ions influences adsorption on talc. It was indicated that the addition of cations alone (without any depressant) at pH 9 could effectively depress talc floatability [Zl66]. In other words, in the presence of metal cations, the importance of depressant molecular weight became less significant and the ionic strength became a more influential factor for the adsorption density ^[83]. A lower electrostatic repulsion between the talc surface and depressant in the presence of metal cations could be the main reason. The addition of monovalent potassium enhanced the range of depressant adsorption onto talc planes due to very few bonding sites' availabilities and depressed the talc floatability $^{[22]}$. However, divalent cations (e.g., Ca²⁺ and Mg²⁺) showed a higher talc depression level than monovalent cations (e.g., K^+ and Na⁺). At pH 9, the addition of these ions could lead to a significant degree of talc depression. At this pH, the hydrophobicity reduction occurred through the ions' adsorption via the addition of charges along the talc planes ^{[66][61][64][71][84]}. In the presence of Ca²⁺ ions, the CMC adsorption onto talc surfaces occurred from the hydrophobicity interactions between the hydrocarbon under the part of glucose rings and the hydrophobic face of talc \overline{C} 9. The combination of talc, CaCl₂, and sodium lignosulfonate, reduced the surface potential of talc due to the complex formation of lignosulfonate ions with Ca^{2+} ions on the talc surface [17].

In addition, trivalent metal ions had a deep influence on the talc hydrophobicity at different pH values. By strong adsorption of hydroxo complex species including Al³⁺, Cr³⁺, and Fe³⁺ onto the polar surfaces, the zeta potential of talc changed its sign from negative to positive, and its natural hydrophobicity markedly decreased ^[85]. The Si⁴⁺ in the tetrahedral silica layer of the talc structure could be replaced with Al^{3+} . Because of this substitution and the interaction between the water molecules and the enhanced polarity of the talc surface state, the water film on the surface is stable. Thus, the contact angle of the sessile drop was reduced by enhancing the aluminum content from about 80 to 0 degrees [61].

3.6. Pre-Treatment

Ultrasonic pretreatment was seen to reduce the flotation recovery of talc at about 15% with lower depressant dosage and 8% with high depressant dosage. This could not be fully explained, but the possible reason could be the release of Mg ions after talc dissolution that might form Mg hydroxo complexes, which by H bonding and chemical interaction reacted with the polymer ^[20]. Ultrasound could create defects on the talc face that act as active sites and improved depressant adsorption. Furthermore, ultrasound could clean the edge surface and reveal the Mg sites where depressants could be adsorbed ^[32]. In industrial processing, the inherent hydrophobicity of talc

is detrimental as it results in poor dispersion ^{[86][87]}. To address this, Yi et al. (2019) applied thermal treatment (calcination at 960 °C) instead of surfactants that break the siloxane structure to amorphous silica. By thermal treatment, the surface wettability changed from 68.87° to 55.34° attributed to the transformation from the siloxane structure (hydrophobic) to amorphous silica (hydrophilic) [88][89].

4. Summary

Talc is a common gangue phase in sulfide ores and remains a challenge in the beneficiation of target commodities from concentration techniques such as froth flotation and pyrometallurgical processes due to the presence of MgO in the latter. Besides being a gangue phase, there is an increasing demand for talc due to its specific properties such as thermal and chemical stability, which find extensive application in cosmetics, papermaking, paint, etc. Based on its natural hydrophobicity, talc remains a challenge in froth flotation, besides its proven versatility in mineral separation.

Due to its chemical composition and structure, talc is a naturally hydrophobic mineral. However, talc is composed of two different surfaces, the basal surface, and the edge, with different surface properties, the former being the majority (approx. 90%) hydrophobic surface, and the latter being the minority (approx 10%) hydrophilic surface. In determining the level of talc hydrophobicity, the

ratio is often the basis. Zeta potential measurements of talc showed that it is charged negatively at pH 2–12 and its IEP is at pH 2–3. The talc flotation showed that the IEP had little to no change when reagents such as depressants were added to the pulp, although the talc's potential decreased at certain pH due to shifting the slips surface double layer towards the outside. Talc's contact angle also showed that its floatability did not vary and remained high even though pH was changed until depressants (e.g., dextrin, guar gum, CMC) were added to the process. Impurities in talc can also decrease the talc contact angle. Although both inorganic and organic depressants have been reported in the literature, the latter seems to be gaining popularity. Given the possible influence of cations and the relatively hazardous nature of inorganic surfactants, using organic depressants is expected to grow. In some cases, using mixtures (inorganic and organic) is reported to improve selectivity, thus offering another potential solution. Considering nonionic polymers (such as starch, dextrin, guar gums) will be commonly opposed to ionic ones (such as CMC) due to their sensitivity to ions present in processed water due to the increased recycling of water.

Due to its natural hydrophobicity, the rejection of talc in the flotation process is challenging. In the collectorless flotation of talc, MIBC has shown to be the most suitable frother that can be used since it had the least frother adsorption capacity and could obtain high recovery. Typical talc depressants are CMC, guar gum, dextrin, etc. The ionic strength and pH of the solution also have some effects on the depressant effectivity. Molecular weight, degree of substitution, and concentration of depressants are some of the factors that affect the depressant performance in the talc flotation. In addition, different ions, including Ca²⁺, Mg²⁺, K⁺, Al³⁺, CO₃²⁻, H⁺, OH⁻, Cr³⁺ and Fe³⁺ influence talc flotation. The ions' presence has a desirable influence on talc depression. Di and trivalent cations, in comparison with monovalent cations, have higher talc depression effects. It is undeniable however that all these

depressants, even with the presence of different ions, do not seem to depress talc in mixed mineral flotation selectively, and so investigating for new and selective talc depressant is still of great interest. Pre-treatment of talc using ultrasound has also increased the talc recovery (~15%) and could be a promising method on an industrial scale. The poor selectivity and efficiency of depressants point to the lack of full understanding of the flotation and depression behaviors of talc and associated sulfide minerals. As such, there is a need for fundamental studies on the froth stability, surface properties, and their alteration by using more advanced and sensitive techniques such as TOF-SIMS.

Due to the existing limitation in surface modification methods, such as using surfactants (depressants), which have poor selectivity and non-uniformity, other techniques were suggested in the literature. Thermal treatment is shown to decrease the wettability of talc from 68.87° to 55.34°, which should also decrease the degree of hydrophobicity. Considering the siloxane structure is mainly composed of talc (001), and pyrophyllite (001) surfaces (basal surface) are mainly hydrophobic. The edge surface due to the broken bonds is hydrophilic on a microscopic scale. Thus any process that can alter the siloxane structure should reduce the hydrophobicity of talc. Pre-treatment of talc using ultrasound has been reported and decreased the talc recovery by 15 %. The well-reported hydrophobicity of talc is also further enhanced by the associated, very poor dispersion. This poor dispersion is detrimental to flotation due to the formation of flocs which may entrain and/or entrap valuable mineral phases, ultimately leading to poor separation performance. For improved dispersion, the application of hydrodynamic cavitation (nanobubbles) can be explored since the depressants-particle contact is very poor due to the tendency of talc to form flocs. These pretreatment techniques may have great potential in surface modification for talc and even other clay minerals.

Although much of the literature on the flotation of talc reports on the treatment of talc as a gangue phase and is set to report to tailings. With the growing number of waste valorization projects, the recovery of talc should also be considered in initial flowsheet development, considering the growing need for talc and also the challenges associated with talc in waste streams, such as poor water recovery in high talc-containing effluent. The discussion should also focus on the recovery of talc instead of its rejection for better resource utilization and reduced environmental impacts.

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