Viscoelasticity of Quartz and Kaolin Slurries in Seawater: Importance of Magnesium Precipitates

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Abstract: In this study, the viscoelastic properties of quartz and kaolin suspensions in seawater were analysed considering two distinct conditions: pH 8 and 10.7. Creep and oscillatory sweep tests provided the rheological parameters. An Anton Paar MCR 102 rheometer (ANAMIN Group, Santiago, Chile) was used with a vane-in-cup configuration, and the data were processed with RheoCompass™ Light software (ANAMIN Group, Santiago, Chile). The outcomes were associated with the formation of solid species principally composed of magnesium precipitates. The magnesium in solution reduced in the presence of quartz (68 wt %), from 1380 to 1280 mg/L. Since the difference was not large regarding the solid-free seawater, the disposition of solid complexes at pH 10.7 was expected to be similar. The jump in pH caused both yield stress and viscoelastic moduli to drop, suggesting that the solid precipitates diminished the strength of the particle networks that made up the suspension. For the kaolin slurries (37 wt %), the yield stress raised when the pH increased, but unlike quartz, there was significant adsorption of magnesium cations. In fact, the concentration of magnesium in solution fell from 1380 to 658 mg/L. Dynamic oscillatory assays revealed structural changes in both pulps; in particular, the phase angle was greater at pH 8 than at pH 10.7, which indicates that at more alkaline conditions, the suspension exhibits a more solid-like character.

Keywords: viscoelasticity; quartz; kaolin; seawater; magnesium precipitates

1. Introduction

Diverse mining companies have their deposits in arid areas, especially in northern Chile, southern Peru, Australia, Africa and Asia [1,2]. In these regions, water availability may be a serious challenge and is the primary purpose for plants to reduce their water demand while avoiding significant alterations in production. A current strategy implemented in several plants is the use of seawater (see Table 1), either through prior desalination treatment by reverse osmosis, where all salts are eliminated, or by simply applying it directly, without altering the salinity [3–5]. The direct use of seawater entails advantages associated with environmental impact and operational costs that involve the building of a desalination plant [6]; however, the complexity of such an undertaking is the subject of constant discussion and research for engineers and scientists [3]. For example, there is the issue of the location of
plants that are far from the coast and at a high altitude. This may require a great amount of investment in pumping, but it is the only option in some cases. In this scenario, the efficient use of hydric resources is inevitable, which means advancing effective closures of seawater circuits. Here, it is essential to acquire a deep rheological understanding of the mineral slurries, which restrict performance at the dewatering stages [7].

Table 1. Examples of seawater use in mining (adapted from Cisternas and Gálvez [3]).

| Plant                  | Country    | Metal                        | Technology       |
|------------------------|------------|------------------------------|------------------|
| El Boleo Proyect       | Mexico     | Copper, cobalt, zinc, manganese | Leaching         |
| Mount Keith (*)        | Australia  | Nickel                       | Flotation        |
| Sierra Gorda SCM       | Chile      | Copper, molybdenum           | Flotation, leaching |
| Black Angel            | Greenland | Lead–zinc                    | Flotation        |
| Batu Hijau             | Indonesia  | Copper-gold                   | Flotation        |
| Beverly Uranium Mine (*)| Australia  | Uranium                      | Leaching in situ |
| Minera Michilla        | Chile      | Copper                       | Leaching         |
| Antucoya Proyect       | Chile      | Copper                       | Leaching         |
| Minera Las Luces       | Chile      | Copper                       | Flotation        |
| Minera Algorta Norte S.A. | Chile    | Iodine                       | Leaching         |

* Saline water.

The high concentration of electrolytes in solution can hinder distinct stages of mineral processing, such as flotation and tailings management [8,9]. Ions alter the interactions between the particles that constitute the slurries, as well as the behaviour of several reagents which are commonly employed in solid–liquid separation. Consequently, the industry is being driven to implement new methods or technologies that can adapt to the challenges posed by a highly saline environment such as seawater [10,11].

Understanding the fundamentals that govern the rheology of mineral pulps requires an in-depth knowledge of the surface interactions that rule the stability of colloidal systems. It is generally considered that the total interaction energy corresponds to the addition of van der Waals forces with the electric double-layer forces. This is the origin of the classical Derjaguin-Landau-Vervey-Overbeek (DLVO) theory. However, additional interactions may arise in highly saline media that are induced by the solvation of ions and structural changes in the association of water molecules that interact with mineral surfaces such as metal oxides [12]. An essential issue for solvation phenomena is the type of ions dissolved in the solution, wherein maker ions such as Na\(^+\), Mg\(^{2+}\) and Ca\(^{2+}\) are small in size and generate a strong electric field that causes the surrounding water molecules to become highly structured. Otherwise, breaker ions such as K\(^+\) and Cl\(^-\) are larger than the maker ions and, therefore, produce a smaller electric field on their neighbouring molecules, disorganising the layers of associated water molecules around the ions [13–15].

Copper concentration operations in freshwater are normally carried out under highly alkaline conditions (above pH 10.5) to depress pyrite and prevent contamination of the concentrate [16–18]; however, it is challenging to operate in seawater at this condition and, generally, the processes are restricted to the natural pH (approx. pH 8), requiring further reagents to deal with the quality of concentrates [5,19]. At a high pH, the seawater divalent cations (Mg\(^{2+}\) and Ca\(^{2+}\)) can hydrolyse to form soluble complexes (e.g., Mg(OH)\(^+\) and Ca(OH)\(^+\)) and varied solid substances (e.g., Mg(OH)\(_2\), Ca(OH)\(_2\), etc.). These species may adsorb onto the mineral’s surface, which includes the main component of copper tailings such as kaolin and silica. Some reports indicate that the presence of complexes reduces the magnitude of the anionic zeta potential of particles, while solid precipitates can even assign them cationic values [20]. This leads to a substantial impact on the interaction between particles, and consequently, the effect on the rheological properties can be remarkable [21,22].

The tailings produced in copper processing operations mainly contain a significant amount of clays and quartz, which are consistently the focus of study since these determine the rheological
behaviour of the tailings [23]. Teh et al. [24] analysed the yield stress and zeta potential at different pH levels in kaolin pulps. The results confirmed that, in most cases, the yield-stress–DLVO model is obeyed, which means that the highest yield stress occurs at low magnitudes of zeta potential. Kaolinite, the molecular formula of which is \[ \text{Al}_4(\text{Si}_4\text{O}_{10})(\text{OH})_8 \], has three crystallographically different surfaces: a silica face, an alumina face and the edges. Previous studies revealed that, at a low pH, the alumina face and the edges have cationic charges, while the silica face is anionic. This promotes the formation of strong bonds between particles and, consequently, high yield stress. However, at a high pH, the faces and edges have anionic charges, which causes the dispersion of the particles and thus leads to lower yield stress [25]. Avadiar et al. [26] studied the rheological behaviour of alumina, silica and kaolin slurries in the presence of calcium and magnesium salts. It was detected that the adsorption of Ca(OH)\(^+\) and the precipitation of Ca(OH)\(_2\) induced a stronger association of particles than that caused by adsorption of Mg(OH)\(^+\) and solids of Mg(OH)\(_2\). The authors ascribed this to the larger size of the calcium complexes and precipitates, and due to hydration/solvation phenomena, these possess a lower enthalpy of hydration than Mg(OH)\(^+\) and Mg(OH)\(_2\), respectively. Therefore, the former should adsorb more readily on the particles’ surface [27].

The yield stress is the most used rheological parameter to characterise the flow properties of mining tailings, indicating the stress that must be overcome by the pulps so that they begin to flow [28–30]. However, extending to a better rheological characterisation can provide more insight into the strength of the particle networks that make up the suspensions, giving further control opportunities in tailings handling, especially in the discharge of the underflow by rakes in thickeners. In this case, mineral pulps can exhibit viscoelastic behaviours, which are described by a viscous component, represented by the storage modulus (\(G'\)), and an elastic part, represented by the loss modulus (\(G''\)) [31]. The viscoelastic modulus can be obtained by means of dynamic methods of oscillatory rheology, which are carried out by subjecting the material to an oscillatory strain \(\gamma(t) = \gamma_0 \sin(\omega t)\), where the resulting stress in relation to time is \(\tau(t) = \gamma_0 (G'(\omega) \sin(\omega t) + G''(\omega) \cos(\omega t))\), where \(G'\) is a measure of the stored energy of the material and, therefore, is related to molecular events of an elastic nature, while \(G''\) is a measure of energy dissipated as heat, associated with molecular events of a viscous character [32]. Additionally, creep tests consist of applying permanent stress to a specific material, measuring the temporal evolution of its shear strain. The tests can be used as a descriptor of the strength of the particle networks since stronger structures deform less [33,34]. Jeldres et al. [15] studied the effect of the type of salt on the viscoelastic behaviour of silica suspensions prepared in monovalent brines. By increasing the size of the cations, higher values of yield stress, elastic modulus and complex viscosity were obtained, while the shear strain after application of fixed stress was lower. The authors explained that silica has a greater tendency to agglomerate in the presence of larger ions such as K\(^+\), forming stronger particle networks compared with smaller salts such as Na\(^+\) and Li\(^+\).

Several studies have analysed rheological behaviour in saline media, with respect to the primary minerals that compose mining tailings, such as quartz and clays. The reported investigations covered interpretation in monovalent [15,35,36] and divalent salt solutions [26,37]. However, no systematic study has explained the viscoelastic behaviour of these minerals in seawater, especially at a high pH (pH > 10.5), where interference by complexes of calcium and magnesium may appear. These conditions are potentially attractive to carry out the concentration stages, particularly in the copper industry, when the ores have a high content of pyrite [38]. In this sense, the objective of this study was to examine the influence of the principal ionic species present in seawater on the viscoelastic behaviour of quartz and kaolin slurries. The varied pH levels were linked to the formation of soluble complexes and solid precipitates of divalent cations, and their remarkable impact on rheological properties were studied through creep and oscillatory rheology tests.
2. Methodology

2.1. Materials

Kaolin was acquired from Ward’s Science, and quantitative X-ray diffraction (XRD) analysis showed that it contained 84 wt % kaolinite \( \text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4 \) and 16 wt % halloysite \( \text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4\cdot2\text{H}_2\text{O} \) (Figure 1). A D5000 X-ray diffractometer (Siemens S.A., Lac Condes, Chile) was used and the data were processed with Total Pattern Analysis Software (TOPAS) (Siemens S.A., Lac Condes, Chile). The FTIR spectrum (Figure 2) showed a double peak at 3696 and 3654 cm\(^{-1} \), characteristic of the kaolin group. Three absorption bands at 3696, 3654 and 3621 cm\(^{-1} \) reflected the high structural order of the samples by the OH stretching of the inner-surface hydroxyl groups. There was Si–O stretching at 1115, 1032 and 1009 cm\(^{-1} \), Si–O–Si deformation appeared at 431 cm\(^{-1} \), and Si–O deformation appeared at 471 cm\(^{-1} \). The zeta potential at pH 8 in distilled water was −40 mV. This was measured using a Zetameter System 4.0 (Zeta-meter, Staunton, VA, USA) following the methodology of Jeldres et al. [15].

![Figure 1. X-ray diffraction (XRD) for kaolin powder.](image1)

![Figure 2. Fourier Transform Infrared Spectroscopy (FTIR) spectrum for kaolin powder.](image2)

Quartz was acquired from a local Chilean store, where the SiO\(_2\) content detected by quantitative XRD was over 99 wt % (see Figure 3) and the zeta potential in distilled water at pH 8 was −45 mV.
while the remaining pulp continued under mixing. Each aliquot was used for a single rheological test and then discarded.

The pH was raised with sodium hydroxide (NaOH) of analytical grade (over 98%), with the presence of sodium carbonate less than 0.5%. The pulps were mixed with a mechanical stirrer, and the reagent was added gradually. The pH was recorded with a PHS-3BW pH meter (Bante Instrument, Shanghai, China).

The volume weighted particle size distribution (PSD) was obtained using a Microtrac S3500 laser diffraction particle size analyser (Verder Scientific, Newtown, PA, USA). The PSD of the quartz and kaolin samples are shown in Figure 4, where it was found that 10% of the particles were smaller than \(d_{10} = 1.8\) and 3.8 \(\mu\)m in the samples of kaolin and quartz, respectively.

![X-ray diffraction (XRD) for quartz powder.](image-url)

**Figure 3.** X-ray diffraction (XRD) for quartz powder.

![Particle size distribution for quartz and kaolin in distilled water at natural pH.](image-url)

**Figure 4.** Particle size distribution for quartz and kaolin in distilled water at natural pH.

Seawater was obtained from the San Jorge Bay in Antofagasta, Chile. The water was filtered at 1 \(\mu\)m using a UV filter system to eliminate the bacterial activity. The cation concentrations were determined by atomic absorption spectrophotometry: \(Na^+\): 10.9 g/L; \(Mg^{2+}\): 1.38 g/L; \(Ca^{2+}\): 0.4 g/L; \(K^+\): 0.39 g/L. By the argentometric method, \(Cl^-\) was found to be 19.6 g/L. By acid-base volumetry, \(HCO_3^-\) was found to be 0.15 mg/L. The conductivity was 50.4 mS/cm at 25 °C and natural pH.

The pH was raised with sodium hydroxide (NaOH) of analytical grade (over 98%), with the presence of sodium carbonate less than 0.5%. The pulps were mixed with a mechanical stirrer, and the reagent was added gradually. The pH was recorded with a PHS-3BW pH meter (Bante Instrument, Shanghai, China).

2.2. Rheology

Stock slurries of silica and kaolin were prepared at 68 and 37 wt %, respectively, by mixing the solid with seawater overnight with magnetic stirring. The pH was adjusted with sodium hydroxide either to pH 8 or 10.7. Once prepared, a 60 mL aliquot was extracted to perform the rheological assays, while the remaining pulp continued under mixing. Each aliquot was used for a single rheological test and then discarded.
The experiments were carried out on an Anton Paar MCR 102 rheometer, and the data were processed with Rheocompass software. A vane-in-cup configuration was used to minimise the wall-slip effects. The diameter of the vane was 2.2 cm and that of the cup was 4.2 cm.

Creep tests were performed by applying constant stress for 1200 s, which generated an increase in the strain angle ($\theta$) as a function of time. Then, the shear strain ($\gamma$) was obtained by Equation (1), considering that the material has linear viscoelastic behaviour:

$$\gamma = \frac{2\theta}{1 - (d_v/d_c)^2}$$

where $\theta$ is the strain angle, $d_v$ is the vane diameter and $d_c$ is the cup diameter. The apparent compliance ($J$) is the apparent shear strain divided by the applied shear stress as follows:

$$J = \frac{\gamma}{\tau_{\text{creep}}}.$$

If $J$ remains constant concerning the applied stress, the slurry exhibits a linear viscoelastic behaviour; otherwise, it is a nonlinear viscoelastic material. Equation (1) is valid only for linear viscoelastic materials, but errors due to nonlinear viscoelasticity are expected to be small because the main interest of this study was to capture only the trend of transition behaviour. The yield stress was determined as the average between the stress that causes a maximum strain (critical strain) and the minimum stress necessary for the pulp to start flowing instantly.

The oscillatory tests were carried out for a frequency sweep between 0.1 and 100 rad/s, keeping the shear strain at 0.5%, previously selected by a sweep amplitude. This ensured that the test was performed in the linear viscoelastic regime, wherein the deformation is proportional to the applied stress, and the viscoelastic moduli have physical meaning.

3. Results

This section analyses the viscoelastic behaviour of silica and kaolin suspensions in seawater, using the results of the creep and oscillatory sweeps tests. Different rheological parameters were obtained, such as apparent critical strain, yield stress, viscoelastic modulus and phase angle. These parameters were associated with the variations of pH and the appearance of magnesium precipitates, which are made in seawater at alkaline conditions, and were distinguished by speciation graphs and the consumption of sodium hydroxide.

3.1. Formation of Magnesium Precipitates

The principal chemical reactions that make soluble magnesium complexes and solid precipitates in seawater are the following:

- $\text{CO}_2(\text{g}) + \text{H}_2\text{O}(\text{l}) \leftrightarrow \text{H}_2\text{CO}_3$
- $\text{H}_2\text{CO}_3 \leftrightarrow \text{HCO}_3^- + \text{H}^+$
- $\text{HCO}_3^- \leftrightarrow \text{CO}_3^{2-} + \text{H}^+$
- $\text{Mg}^{2+} + \text{HCO}_3^- \leftrightarrow \text{MgHCO}_3^+$
- $\text{Mg}^{2+} + \text{OH}^- \leftrightarrow \text{MgOH}^+$
- $\text{Mg}^{2+} + \text{CO}_3^{2-} \leftrightarrow \text{MgCO}_3$
- $\text{Mg}^{2+} + 2\text{OH}^- \leftrightarrow \text{Mg(OH)}_2$

The formation of magnesium precipitates and complexes generates a buffer effect, which can be identified in a range where a high amount of reagent is required to raise the pH slightly. Nevertheless,
the presence of particles (concentrated slurries) interferes with the concentration of ions dissolved in solution, modifying precipitation reactions and the buffer effect. Indeed, a chemical analysis by Atomic Absorption Spectroscopy (AAS) determined that the magnesium in seawater reduced from 1380 to 1280 mg/L in the quartz pulps, while it dropped from 1380 to 658 mg/L in the kaolin suspension.

The amount of sodium hydroxide used (g/L) to raise the pH of seawater, quartz pulp and kaolin pulp, normalised by the amount required to obtain pH 12.2, is shown in Figure 5. A marked buffer effect was observed for pure seawater and quartz slurry between pH 10 and 11, which coincided with the formation of magnesium precipitates, where carbonate precipitation started at pH 9.3 and ended at pH 10, while hydroxide deposition started at pH 10.3 and completed at pH 11 [39–41]. This effect was not clear in the kaolin pulp, indicating a lower formation of magnesium precipitates, which is compatible with the loss of magnesium in solution (from 1380 to 659 mg/L). Considering the low cationic exchange capacity of kaolinite [42,43], it is expected that magnesium cations are attached to the anionic sites of the clay’s surface by electrostatic attraction. The adsorption of magnesium ions reduces its concentration in solution, reducing the formation of solid complexes at a high pH.

Figure 5. Sodium hydroxide (NaOH) required to modify the pH of (i) seawater, (ii) quartz slurry in seawater and (iii) kaolin slurry in seawater. NaOH is mass (g) consumed per 1 L of solution, normalised by the concentration at pH 12.2 (seawater: 6.1 g/L; quartz: 5.7 g/L; kaolin: 9.1 g/L).

3.2. Creep Tests

Figure 6a–d display the creep tests for quartz and kaolin slurries prepared with seawater at pH 8 and 10.7. The temporal evolution of the apparent strain was measured under fixed stress for 1200 s, and in all cases, a maximum strain was observed that remained constant at high times. When the applied stress was greater than a critical value, the pulp started to flow, and its deformation diverged. An apparent instantaneous strain was observed for the quartz pulps when the stress was lower than half of the yield stress (Figure 6a,b). This effect was not observed for kaolin pulps, which exhibited a gradual increase in their apparent strains (Figure 6c,d).

Figure 6. Cont.
10.7. Interestingly, the decrease of yield stress by increasing the pH (from pH 8 to 10.7) coincided with the formation of solid complexes in seawater. In the literature, it has been reported that these products can adsorb onto the surface of the particles, having different consequences on the rheological properties [22]. The solid complexes may form bonds between quartz particles, mainly due to the reduction of zeta potential [20], but the strength of these bonds would be weaker compared with the cationic bridges generated by the divalent Mg\(^{2+}\) ions. For this reason, it is expected that at pH 8, where the concentration of Mg\(^{2+}\) ions in solution is higher, the yield stress is higher. On the other hand, the kaolin suspension at pH 8 had a yield stress of 38 Pa, while at pH 10.7, it increased to 42 Pa. This behaviour is contrary to the quartz pulp and can be justified by the low presence of solid precipitates.

![Figure 6](image_url)  
**Figure 6.** Creep test for quartz and kaolin slurries at varied applied stresses: (a) quartz at pH 8, (b) quartz at pH 10.7, (c) kaolin at pH 8 and (d) kaolin at pH 10.7.

The yield stress for each pulp examined in this study is shown in Figure 7. The quartz slurry at pH 8 reached a yield stress of 56 Pa, which was significantly higher than the 43 Pa obtained at pH 10.7. Interestingly, the decrease of yield stress by increasing the pH (from pH 8 to 10.7) coincided with the formation of solid complexes in seawater. In the literature, it has been reported that these products can adsorb onto the surface of the particles, having different consequences on the rheological properties [22]. The solid complexes may form bonds between quartz particles, mainly due to the reduction of zeta potential [20], but the strength of these bonds would be weaker compared with the cationic bridges generated by the divalent Mg\(^{2+}\) ions. For this reason, it is expected that at pH 8, where the concentration of Mg\(^{2+}\) ions in solution is higher, the yield stress is higher. On the other hand, the kaolin suspension at pH 8 had a yield stress of 38 Pa, while at pH 10.7, it increased to 42 Pa. This behaviour is contrary to the quartz pulp and can be justified by the low presence of solid precipitates.

![Figure 7](image_url)  
**Figure 7.** Yield stress for quartz and kaolin slurries at different pH levels.

Critical shear strain is an attractive property of materials, which is directly associated with the strength of the bonds that form the particle networks just before they begin to flow [44]. Figure 8 shows the critical strain, where for both minerals, the shear strain decreased as there was an increase in pH.
which was employed. This characteristic is shown in Figure 10, considering the compliance at the end of the experiment (1200 s). The kaolin pulp at pH 8 suggested linear viscoelastic behaviour when the stress was lower than 20 Pa, while at pH 10.7, the yield stress was displayed in a range of stress (40–46 Pa) instead of a single point. For that matter, the pulp began to flow depending on both the applied stress and the time of stress application (Figure 9). Otherwise, for pulps in which there were no solid precipitates, the yield stress was a precise value: quartz pH 8, 55 Pa; kaolin pH 8, 38 Pa; and kaolin pH 10.7, 42 Pa.

Mineral pulps may exhibit linear or nonlinear viscoelastic behaviour, or both, depending on the applied stress. A linear response implies that compliance remains constant, independent of the stress which was employed. This characteristic is shown in Figure 10, considering the compliance at the end of the experiment (1200 s). The kaolin pulp at pH 8 suggested linear viscoelastic behaviour when the stress was lower than 20 Pa, while at pH 10.7, the linear response limit was up to the stress of 35 Pa. On the other hand, the quartz slurry at pH 8 had a linear behaviour up to 30 Pa, while at pH 10.7, the system of which was affected by the presence of complexes and solid precipitates, the linearity decreased considerably to a stress lower than 10 Pa.

Figure 8. Critical shear strain for quartz and kaolin slurries at different pH levels.

Figure 9. Relationship between yield stress and applied time for quartz slurries at pH 10.7.
3.3. Oscillatory Rheology

Oscillatory rheology tests are a good descriptor of pulp strength. In this case, a frequency sweep was performed considering a constant amplitude of 0.5%, previously defined by an amplitude sweep test. This ensured that the pulp was in a linear viscoelastic regime. Figure 11 shows the response of both moduli $G'$ and $G''$ concerning the angular frequency for the silica suspension. Viscoelastic moduli had a higher value when silica was at pH 8, meaning a greater resistance of the particle networks, while at pH 10.7, both moduli were diminished. Interestingly, for both pH levels, the moduli $G'$ and $G''$ crossed within the range of frequencies examined. The crossover point was an indicator of the relaxation time, using the reciprocal of frequency under conditions where both moduli intersected. While at pH 8 the relaxation time was 8.6 s, at pH 10.7, the value increased to 10.8 s. For the kaolin pulps, the behaviour was different (Figure 12), and for both pH values (8 and 10.7), the storage modulus ($G'$) was higher than the loss modulus ($G''$). It can be argued that both moduli had a weak dependence on frequency, which is a typical gel behaviour [32]. The phase angle, as shown in Figure 13, was higher at pH 8 than at pH 10.7, for both minerals. This suggests that the pH changed the internal structure of the pulps, adopting a more solid-like character when the pH was more alkaline. In the silica suspension, it was observed that at frequencies greater than 30 rad/s, the phase angle was independent of the applied frequency, while for kaolin pulp, this happened at over 10 rad/s.

![Figure 10](image1.png)

**Figure 10.** Final apparent compliance as a function of stress for quartz and kaolin slurries.

![Figure 11](image2.png)

**Figure 11.** Frequency sweep for quartz slurries at different pH levels.
4. Conclusions

The viscoelastic properties of quartz and kaolin suspensions in seawater were analysed, examining two different conditions: (i) pH 8, which resembles that used by copper concentrating plants, and (ii) pH 10.7, which is attractive since it may enhance the quality of concentrates. Nevertheless, the latter is characterised by speciation reactions of divalent seawater cations, which lead to the formation of solid precipitates, mainly magnesium complexes. This feature changes for slurries according to the mineral species because the particles’ surfaces can adsorb a portion of the cations, reducing their content in solution. Here, quartz pulp reduced magnesium from 1380 to 1280 mg/L. Since it was not a large difference from the solid-free seawater, the disposition of solid complexes at pH 10.7 was similar. As rheological consequences, the increase in pH caused both yield stress and viscoelastic moduli to drop, suggesting that the solid precipitates diminished the strength of the particle networks that made up the suspension. For the kaolin slurries, the yield stress raised when the pH increased, but unlike quartz, there was significant adsorption of magnesium cations. In fact, the concentration of magnesium in solution fell from 1380 to 658 mg/L. Since it was not a large difference from the solid-free seawater, the disposition of solid complexes at pH 10.7 was similar.

The dynamic oscillatory assays revealed structural changes for both pulps; in particular, the phase angle was higher at pH 8 than at pH 10.7, showing that the more alkaline suspension exhibited a more solid-like character.

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