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Detrimental effects of Magnesium (II) on hydroxyapatite precipitation from synthetic industrial brines

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Abstract

The influence of Mg(II) on phosphorous recovery as hydroxyapatite (Hap) from alkaline phosphate concentrates using desalinated industrial brines as the calcium source in a batch reactor was evaluated. Two synthetic brines with Mg/Ca molar ratios of 2.2 and 3.3 were continuously fed to reach a Ca/P molar ratio of \( \sim 1.67 \) to promote Hap formation under different constant pH values (8, 9.5, 10.5, 11.5 and 12). For both brines, inhibition of Hap precipitation and formation of the amorphous mineral phases of Ca-, Mg- and Ca/Mg-phosphates were observed at pH >9.5. Mg(II) severely inhibited phosphate precipitation, allowing the formation of amorphous calcium phosphate from meta-stable clusters due to Mg(II) incorporation into Ca-phosphate. For the Mg/Ca (3.3) brine, a more soluble Mg-phosphate mineral (cattiite) was formed at pH 11.5. Thermal treatment of the amorphous solids to increase crystallinity confirmed the presence of
Hap and chlorapatite as Ca-phosphate, stanfieldite as Ca-Mg-phosphate and farringtonite as Mg-phosphate. In the experiments at pH 8, the formation of stable nanometre-sized pre-nucleation clusters promoted nucleation inhibition, even in supersaturated solutions, and no solids were recovered after filtration. Although sulfate was involved in some of the precipitation reactions, its role in the inhibition of Hap formation is not clearly elucidated.

Keywords: phosphate recovery; brines; hydroxyapatite; stanfieldite; cattiite; farringtonite

1. Introduction

Phosphorus (P) is a non-renewable resource, non-substitutable for agriculture and food production and directly linked to global food security, as well as being important in other industrial and technical uses. At the same time, P losses are the principal contributor to eutrophication of surface waters, globally the P footprint of human diets continues to increase and the world mineral phosphate reserves decrease and there is a debate about their extent and extractability and about their geographical concentration. Improving the efficiency of P processing and use, in industry, agriculture, livestock production, food processing, and developing P reuse or recovery-recycling can reduce costs, contribute to reducing nutrient pollution, and create jobs in the frame of circular economy [1].

Phosphate is typically present at low concentrations in urban wastewaters (from 10 to 30 mg P-PO₄³⁻/L) and in industrial wastewaters, such as detergent manufacturing, food processing or metal-coating processes (50 to 0 mg P-PO₄³⁻/L) [2–4]. The removal of phosphate from water bodies is important because it causes eutrophication, which has a harmful effect on aquatic life, resulting in a reduction in biodiversity. On the other hand, the recovery of phosphate from phosphorus-containing wastewater is essential for developing an alternative phosphorus source to overcome the global challenge of its scarcity [5]. However, one of the disadvantages that complicates phosphate recovery is the low concentration of phosphate in the target effluents.
Many different processes have been proposed for pre-concentration of phosphate, such as adsorption, ion exchange and biological treatment [6–8]. The introduction of new P-selective sorbents (e.g., hydrated metal oxide based sorbents) would generate alkaline phosphate concentrates due to the requirements of the sorbent regeneration with NaOH solutions [6].

Chemical phosphorus recovery using Ca(II) and Mg/NH₄ salts to precipitate or crystallize phosphate as NH₄-Mg or Ca salts are the primary solutions postulated [9]. To address this objective, the use of industrial wastes as alternative Ca(II) sources for Ca-phosphate precipitation has been suggested. Ca-phosphates can be recovered by crystallization of Hap in appropriate reactors via pH and chemical dosing control, as reported previously [10-13]. Recently, the use of seawater reverse osmosis and nanofiltration brines for the recovery of economically valuable constituents [14] or specifically as an inexpensive Mg(II) and Ca(II) source, and for struvite recovery from anaerobic digesters in municipal wastewater treatment plants was suggested [15-17].

The significant Ca(II) concentration present in seawater brines (up to 0.4 g Ca/L) may enhance the precipitation of Ca-phosphate minerals (e.g., Ca₃(PO₄)₂ and Ca₅(PO₄)₃(OH), among others). However, the influence of high concentrations of Mg(II) up to 1 g Mg(II)/L is unknown. Salami et al. [18] reported no detectable effect of Mg(II) ions on the growth of dicalcium phosphate dihydrate, but they did report that the Mg(II) ions appreciably decelerated the rate of octacalcium phosphate growth, most likely by adsorption at active growth. More recently, Cao and Harris [19] studied the interactive effects of CO₃²⁻ and Mg(II) ions on Ca-phosphate precipitation under conditions simulating dairy manure-amended soil leachate and phosphate recovery from manure wastewater. The inhibition effects of Mg(II) and the synergistic effect of both of the ions on Hap crystallinity and the precipitation rate promoted the formation of amorphous Ca-phosphate (ACP), presumably due to Mg(II) incorporation into the crystal structure. However, the presence of Mg(II) or SO₄²⁻ ions in the case of using industrial desalinated brines at concentrations higher than the
calcium ions has not been studied. Moreover, few studies in literature are devoted to study the potential precipitation of Ca-Mg-phosphate minerals and the mechanism involved. It should be mentioned the work done by Golubev et al. [20, 21] who postulated the formation of \((\text{Ca,Mg})_4\text{H}(\text{PO}_4)_3\cdot x\text{H}_2\text{O}\) in the precipitation of phosphate with sea water and more recently Muster et al. [22] who postulated theoretically the formation of potential Ca-Mg phases.

Therefore, the goal of this study is to evaluate the potential inhibition of Mg(II) on hydroxyapatite (Hap) precipitation during the valorisation of concentrated phosphate effluents when using synthetic industrial desalinated brines as the calcium source. Two brines with different Mg/Ca molar ratios of (2.2) and (3.3) were used. The precipitation/crystallization of Ca- and/or Mg-phosphate processes at different constant pH values were evaluated in a batch reactor and the precipitate properties were also studied. The variation of the Ca- and Mg-phosphate nucleation profiles was used to elucidate the formation mechanism of Hap or Mg-phosphates with high Mg(II) concentration brine.

2. Materials and Methods

2.1 Experimental set-up and procedures

The precipitation of phosphate (P(V)) was performed in a 2 L glass batch reactor at constant pH values (8, 9.5, 10.5, 11.5 and 12), following the conditions defined in a previously study [13]. These alkaline pH values were selected based on the thermodynamic prediction for the precipitation of Ca and Mg phosphates. Stirring at 250 rpm was achieved using a mechanical stirrer (IKA RW 20). The pH was monitored on-line using a pH potentiometer (Crison pH 28), when the pH was 0.1 units above or below the set point, 1 M HCl or 1 M NaOH was dosed using a peristaltic pump. Batch experiments were performed by mixing a 1.0 g P-PO\(_4^{3-}\)/L solution with Mg/Ca brine. NaH\(_2\)PO\(_4\) was used to prepare the phosphate solutions. Composition was fixed according to the expected conditions of the elution of ion exchange resins on the recovery of...
phosphate from treated waste water effluents. Two synthetic solutions with different Mg/Ca molar ratios (2.2 and 3.3) were prepared by mixing given amounts of NaCl, CaCl₂·2H₂O, Na₂SO₄ and MgCl₂·6H₂O. The compositions of both of the brines are summarized in Table 1. The presence of antiscalants typically present on desalination brines (e.g. 1-2 mg/L) and the temperature were not included in the experimental design.

Brine solution was added at a flow rate of 0.3 mL/min (using a Gilson Minipuls 3 peristaltic pump) to reach a Ca/Pa molar ratio of 1.67 suitable for Hap precipitation. Experiments were performed at room temperature (22±2 °C) in duplicate.

Table 1.

<table>
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<th>Batch reactor aqueous samples were obtained during the experiments and then filtered through a 0.22-µm filter. The total concentrations of ions were determined by ion chromatography using an Ionex Liquid Chromatograph (ICS-1000). The accuracy of the measurements was higher than 95%. At the end of the experiments, the solid phase was removed from the reactor by filtration, washed with deionised water several times and dried at 60°C for 24 h.</th>
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2.2 Particle analysis

The solid phase particle size distribution was analysed by LS with a Coulter diffraction particle size analyser (LS 13 320 Laser Diffraction Particle Size Analyser Instrument, Beckman Coulter). The crystal size distribution range (CSD) varied from 0.04 to 2000 µm. Particles were analysed as obtained directly from the batch reactor without any thermal treatment or particle size separation.

The phase purity and crystallinity of powder were analysed by X-ray diffraction with λ CuKα radiation (λ= 1.54056 Å) at a scanning rate of 19.2 and 57.6 s, a steep angle of 0.015° and 2θ over range of 4 to 60°. The solids in powder form were identified by the Joint Committee Powder Diffraction Standards (JCPDS) file and were compared with the Powder Diffraction File (PDF) no.
00-009-0432 for Hap (Ca_{10}(PO_4)_6(OH)_2), 00-011-0231 for stanfieldite (Ca_4Mg_5(PO_4)_6), 00-025-1373 for farringtonite (Mg_3(PO_4)_2) and 00-001-1011 for chlorapatite (Ca_{10}Cl_2(PO_4)_6) [23].

To elucidate the potential inhibition mechanism, a portion of the amorphous phases was heated at 1050°C for 4 h and cooled at room temperature to enhance the crystallinity of the precipitated phases. It cannot be discarded that the thermal treatment promotes additionally to an increase of crystallinity a change on the crystal structure or the chemical composition. However, the information provided after this treatment is valued as it is improving the mechanisms discussion [24].

2.3 Prediction of phosphate precipitation processes

Phosphate precipitation processes using Mg/Ca brines were studied using the HYDRA-MEDUSA [25] and the Visual Minteq codes [26]. The measured P(V), Mg(II), Ca(II), SO_4^{2-}, and Cl^- concentrations were compared when required to those estimated using both of the codes. Although conditions in the precipitation tests could be far away from the equilibrium, measured and predicted values were used to identify the potential reactions and mechanism involved, especially when the solids formed were not appropriately characterized.

The supersaturation index (SI) was calculated by Visual Minteq and using Equation 1, as follows:

\[
SI = \log \left( \frac{\text{IAP}}{K_{sp}} \right)
\]  

(1)

where IAP is the ion activity product, and K_{sp} is the solubility constant. Equilibrium solubility data for Ca-Mg-phosphates were critically reviewed from the HYDRA and PHREEQ C databases, and the selected values are shown in Table (S1) (Supporting Material) [11, 27, 28].

3. Results and Discussion
3.1 Influence of pH on the phosphate recovery with Ca/Mg brines

Precipitation of Hap with Mg/Ca (2.2) and Mg/Ca (3.3) brines was studied as a function of pH (8, 9.5, 10.5, 11.5 and 12). The change of total phosphate concentration and recovery (%) as a function of reaction time is shown in Figure 1 in which the dotted lines represent the expected total phosphate concentration if any precipitation reaction was involved. Phosphate recovery efficiency in the richest Mg brine (Mg/Ca 3.3) (Figure 1 c and d) is larger at pH up to 10.5. Higher recoveries were measured in only 18 hours, while for the Mg/Ca (2.2) brine were observed after 34 hours (Figure 1 a and b). A similar trend was reported by Su et al. [11] when precipitating phosphate with magnesium chloride solutions (Mg/P (2.1)) in alkaline media (pH 10, 11 and 12) in a fluidized bed reactor.

Figure 1.

The lowest phosphate recoveries (20%) were reported at pH 8 for Mg/Ca (2.2) brine and at pH 8 and 9.5 for Mg/Ca (3.3) brine. For both of the brines (at lower pH) at the end of the experiment after filtration, the solutions presented turbidity, and no precipitate was recovered on the 0.22-µm filter. This result was associated with the inhibition of the nucleation process and the formation of clusters of the nanometre size, as discussed in section 4.3.

The increase of phosphate recovery efficiency with increasing pH is explained by the change of P(V) speciation. At pH 8, 45% of P(V) is present in solution as HPO$_4^{2-}$, and less than 4% is present as PO$_4^{3-}$ for the initial additions of brine with 12-8 mmol/L P(V) concentrations. However, at pH 11.5, 41% of P(V) is present as PO$_4^{3-}$ and 20% as HPO$_4^{2-}$, and a higher SI is achieved. As demonstrated by Gunawan et al. [29], the degree of supersaturation and the type of precipitates formed depend on the pH. A higher pH leads to higher SI and accelerates the precipitation reaction as well as increasing its efficiency.
3.2 Influence of brine composition on Hap formation and precipitation inhibition

3.2.1 Recovery of phosphate by Mg/Ca (2.2) brines

The change of the Mg(II), Ca(II), SO$_4^{2-}$ and Cl$^{-}$ contents as a function of precipitation reaction time is plotted in Figure 2 in which the solid lines represent the total ion concentration added to the reactor throughout the experiment, which is the concentration expected to be measured for a species not involved in any precipitation or solid formation reaction.

![Figure 2.](image)

As can be seen in Figure 2a, Mg(II) concentration is reduced to less than 0.2 mmol/L, independent of the pH value, while the Ca(II) concentrations are maintained below 1 mmol/L, with the exception of pH 8, where values are equal to the total added Ca(II) concentration. In the case of SO$_4^{2-}$, the measured concentrations agree with the total added concentration, except for the experiment at pH 8 with values below 10%. For Cl$^{-}$, the measured concentrations showed a reduction from 700 to 400 mmol/L. This behaviour confirms that these ions (Mg(II), Ca(II) and Cl$^{-}$) are involved in the precipitation reactions.

![Figure 3.](image)

The variation of the SI of the expected mineral phases along the reaction time, such as Hap, tricalcium phosphate (TCP), octacalcium phosphate (OCP), monotite and brushite for Ca(II) as well as Mg(OH)$_2$ and Mg$_3$(PO$_4$)$_2$(s) for Mg(II) is shown in Figure 3 for pH 11.5 and 8. At higher pH values (9.5 and 11.5), the precipitation of Ca-phosphates is favoured (up to 90% of P(V) recovery at pH 11.5, as shown in Figure 1). The SI of the Ca-phosphate mineral phases were close to zero for brushite and monotite, close to 4 for TCP and OCP, and close to 18 for Hap (Figure 3a). Therefore, nucleation of Hap, the most stable phase among the Ca-phosphates, is expected to occur instantaneously [30]. Typically, supersaturation of Hap is achieved by a simple increase in
pH [31], and then it follows a three-stage process in which the initially formed amorphous ACP may be redissolved and form HAP nuclei followed by formation of hydroxyapatite [32-34].

XRD analysis of the precipitates collected in both of the experiments at pH 9.5 and 11.5 reveals that the formed Ca-phosphates were amorphous, as shown by a broad peak between 23° and 35° (2θ) (Figure 4a). These patterns are typical of ACP [35], indicating that Mg(II) promoted the formation of the relatively unstable ACP, in the form of ACP-adsorbed Mg(II), and then hindered the expected Hap formation according to the saturation indexes [36, 37].

Figure 4.

Yang et al. [38] reported that Mg(II) ions reduce the nucleation rate of Hap in Ca-phosphate supersaturated solutions by stabilizing the gel-like ACP phase and increasing the induction and transformation time. Ding et al. [37] described that Mg(II) ion adsorption onto ACP is more effective than the phase incorporation at inhibiting phase transformation from ACP to Hap. Additionally, at these pH values, sulfate was not involved in the formation of solid phases (measured values agree with the total added concentration), and it is present in solution primarily as complexed species (e.g., MgSO₄(aq) and CaSO₄(aq)), avoiding the precipitation or re-dissolution of potential Ca-phosphate precipitates [7].

The SEM–EDX analysis of the amorphous solids confirmed the major presence of precipitates containing Ca–P–O and to a minor extent, Mg(II) and Cl⁻. Considering that the Mg(II) and Ca(II) removal ratios for both of the pH values were higher than 90%, the solubility data of different Ca/Mg-phosphate mineral phases (Mg₃(PO₄)₂(s), Ca₅OH(PO₄)₃(s)) were used to predict the expected P(V) concentration throughout the experiment at each given pH. For both of the pH values Figures (S1 b and c) (Supporting Material), the measured P(V) concentrations were better predicted when assuming the formation of Mg-phosphate minerals than when assuming the formation of Ca-phosphate minerals.
The XRD analysis of the amorphous precipitate at pH 9.5, after treatment at 1050°C to increase its crystallinity, identified the presence of a Ca-phosphate mineral (Hap (Ca$_5$OH(PO$_4$)$_3$)), a Ca-Mg-phosphate mineral (stanfieldite (Ca$_4$Mg$_5$(PO$_4$)$_6$)) and a Mg-phosphate mineral (farringtonite Mg$_3$(PO$_4$)$_2$) (Figure 4b). In the case of the amorphous precipitate at pH 11.5, in addition to the presence of Hap and stanfieldite, a Ca-phosphate-chloride mineral (chlorapatite (Ca$_5$Cl (PO$_4$)$_3$)) was detected (Figure 4b). Therefore, the consumption of chloride in the precipitation reactions was confirmed (Figure 2), and it was also identified by EDX analysis, as described in Table (S2) (Supporting Material).

In the experiment at pH 8 because Ca(II) was not consumed, and Mg(II) was completely consumed, the P(V) recovery (up to 20%) should be associated with the formation of Mg-phosphate or magnesium hydroxide. The SI indicates that the solution is not supersaturated in Mg(OH)$_2$(s) (Figure 3c); thus, the recovery of P(V) should be associated with the formation of Mg-phosphate and, potentially, with the formation of minerals containing sulfate because the measured values are lower than the total added concentration. A comparison of the measured and expected P(V) concentration provides a good prediction of the measured P(V) concentration profile considering the formation of Mg-phosphate, as shown in Figure (S1 a) (Supporting Material). Although solutions were supersaturated in Hap, OCP, TCP, brushite and monitite (Figure 3b), the observed Ca-phosphate mineral inhibition could be due to the formation of CaSO$_4$ as a precursor because sulfate is present at a substantially higher concentration than phosphate or due to the inhibition of Mg(II) ions. At the end of the experiment, no solid was recovered after solution filtration because the size of the precipitate is expected to be of nanometre size (ca. below 1 nm); thus, it was not possible to confirm its chemical or mineral composition or the potential inhibition effect of sulfate ions.

3.2.2 Recovery of phosphate by Mg/Ca (3.3) brines
The change of the major ion concentration as a function of reaction time at different pH values is plotted in Figure 5. Mg(II) concentration was reduced from 15 mmol/L to less than 5 mmol/L at the end of the experiments at pH 9.5 and 11.5. The Ca(II) concentration was reduced to values of 1 mmol/L for the experiment at pH 9.5, while for the test at pH 8 and 11.5, the measured values approached the total added Ca(II) concentration (4 mmol/L), indicating that Ca(II) did not participate in any precipitation reaction. Sulfate concentration was reduced from 80 mmol/L to 60 mmol/L at the end of the experiments, indicating that sulfate was involved in the precipitation reactions. The measured chloride concentrations approached the total added concentration, indicating that it was not involved in any precipitation reaction, contrary to the observed behaviour for Mg/Ca (2.2) brine.

**Figure 5.**

XRD analysis revealed that the solid product collected at pH 9.5 was amorphous, while at pH 11.5, cattiite (Mg₃(PO₄)₂·22H₂O) was detected. These results confirm the profiles of Ca(II) and Mg(II) shown in Figure 5 in which the Ca(II) concentration was not reduced as the experiment progressed, as revealed when it was identified by EDX analysis, as summarized in Table (S2) (Supporting Material).

**Figure 6.**

At pH 11.5, the solution is supersaturated in Hap; however, the higher initial magnesium concentration inhibited its precipitation, and cattiite was found (logK_{s0} = -23.1) [39] as shown in Figure 6a. Note that cattiite was formed in the presence of a high Mg(II) concentration instead of the less-soluble solid Hap (logK_{s0} = 57.8) due to the effect of Mg(II) ions on the field stability of the solids, promoting the precipitation of more soluble solids [40, 41]. The measured P(V) concentrations were well predicted assuming the formation of cattiite, as shown in Figure (S2) (Supporting Material).
The amorphous precipitate at pH 9.5 shows a broad peak between 23° and 35° (2θ) (Figure 6a) of ACP. The SEM–EDS examination of the amorphous sample detected the presence of Ca-Mg–P–O solids and the minor presence of S. The XRD analysis of the treated sample at 1050°C identified the presence of stanfieldite (Ca₄Mg₅(PO₄)₆) (Figure 6 b and c), thus confirming the consumption of Mg(II) and Ca(II), as described in Figure 5. The higher Mg(II) concentration inhibits the Hap precipitation, favouring the formation of mixed Ca-Mg-phosphates, such as stanfieldite (Ca₄Mg₅(PO₄)₆), as detected by XRD. Mg(II) stabilizes ACP, which is the precursor phase during Hap formation from highly supersaturated solutions [42]. It was also described that Mg(II) could be included in the precipitated solid and could modify the solids by its smaller size and greater tendency to bond covalently [43]. Lahav et al. [17] postulated that the complexation of Ca(II) ions in the precipitation of P(V) using seawater desalination brines reduces their free concentrations, thus reducing their precipitation potential and reducing the purity of the Ca-phosphates. The analysis of solids by SEM-EDX detected the presence of S and Cl, indicating their presence in the precipitates obtained at basic pH; however, XRD analysis did not detect any crystalline form.

For the experiment at pH 8, Ca(II) and Mg(II) were partially removed (approximately 10%) with a phosphate recovery of up to 20%. The SI analysis indicated that the solution is not supersaturated in Mg(OH)₂(s), and the removal of P(V) should be associated with the formation of Ca-Mg-phosphates and, potentially, with minerals containing sulfate, because the measured values were lower than the total added concentration. A good prediction of the measured concentrations was obtained when considering the formation of Mg- and Ca-phosphates, as shown in Figure (S2 a) (Supporting Material). Although solutions were also supersaturated in OCP, TCP, brushite and monotite, the observed inhibition could be either due to Mg(II) ions or the formation of CaSO₄aq of MgSO₄aq species because sulfate is present at a substantially higher concentration than phosphate. As previously described for the Mg/Ca (2.2) brine, after solution
filtration, no solids were recovered due to the nanometre size of the formed clusters, and it was not possible to confirm its chemical or mineral composition or the potential inhibition effect of sulfate ions.

### 3.3 Evaluation of precipitation inhibition: Effects on nucleation growth kinetics

According to the XRD results, it was observed that nucleation of Hap begins with the formation of the ACP precursor during the early induction steps, and after a relatively long induction period, it proceeds to the appearance of nuclei [44, 45]. However, in the presence of Mg(II), the formation of Mg-phosphate ion-pairs reduces the P(V) species activity, thereby reducing the relative supersaturation and prolonging the induction period [42]. Figure 1 shows that the change of the P(V) concentration and recovery rate for pH values between 9.5 and 12 for Mg/Ca (2.2) brine and between 10.5 and 12 for Mg/Ca (3.3) brine is different than for the experiments at pH 8 for Mg/Ca (2.2) brine and at pH 8 and 9.5 for Mg/Ca (3.3) brine.

The presence of meta-stable pre-nucleation clusters (PNCs) decreases the energetic barrier, thereby facilitating nucleation (pH 9.5 and 11.5), while at pH = 8, stable clusters are considered to increase the barrier, thus inhibiting nucleation [46, 47].

**Figure 7.** Phosphate precipitation can be evaluated considering the variation of the SI at pH 8, 9.5 and 11.5 (Figure 7) and by using the LaMer model (Appendix) describing the crystallization processes as three well-defined stages. At pH 11.5, there was an initial stage in which no precipitation occurred; in the second stage in which the SI reached values from 17 to 18.5, homogenous nucleation occurred; and the third stage is completed with the aggregation of small particles of the homogeneously nucleated material and their heterogeneous deposition [48]. For the experiment at pH 9.5 from the initial additions, a supersaturation condition was observed (SI>18.5), and then the homogenous nucleation and final aggregation stages followed the trend.
defined at pH 11.5. The observed behaviour for the experiment at pH 8 (no solid was recovered) is associated with the formation of stable clusters, increasing the energetic barrier, hindering nucleation and achieving heterogeneous nucleation. The longer induction time for nucleation as the Mg(II) ions extend the induction and transformation time [37, 38] promotes the formation of nanometre-sized nuclei (Posner's clusters of 0.7 to 1.0 nm [49]). These nanometre-sized crystals could not be recovered by the 0.22-µm filter.

To evaluate the influence of the pH and the supersaturation on the Hap nucleation kinetics, the plot of ln(ts) versus 1/[ln(1+σ)]^2 was constructed for experiments with Mg/Ca (2.2) brine at pH 8 and 9.5, as shown in Figure 8.

Figure 8.

According to Equation A3 (in the appendix) the charge in the crystalline phase structures can be analysed for a given system (k_n and B are constant) in terms of the variation of the slope (k_nf(m)). As shown in Figure 8, the depicted functions can in a first approach, be fitted by two intersecting straight lines with two slopes, which divide the supersaturation space into two regimes.

For both of the pH values (8 and 9.5), the function has a positive slope (regime 2) reaching a transition point (indicated by a vertical dotted line) followed by a plateau (regime 1) with a decrease of the slope. When comparing both of the experiments, a case of nucleation inhibition was identified at pH 8, as was postulated by Jian et al. [50] who determined the inhibition effect by the increase in the slope and the decrease of the intercept. On the other hand, for the experiment at pH 9.5, a case of nucleation promotion was identified, with a factor of (k_nf(m2)=243), referring to regime 2, which was much lower than that reported at pH =8 (k_nf(m2)=760). This result indicates that at pH 8, it is possible to reduce the nucleation barrier by improving the interfacial structure correlation [47, 50]. Similar results (data not shown) were obtained for the Mg/Ca (3.3) brines.
The particle size distribution in terms of volume and the number of particles for both of the Mg/Ca brines at pH 11.5 is shown in Figure (S3) (Supporting Material). The number of particles with a mean size ($d_{50}$) increases with the Mg(II) concentration from 310 nm to 1400 nm for the Mg/Ca (2.2) and Mg/Ca (3.3) brines, respectively. The precipitate particle size analysis in terms of volume revealed the formation of aggregates with an average equivalent diameter of approximately 113 µm and 62 µm for the Mg/Ca (2.2) and Mg/Ca (3.3) brines, respectively. The initially formed particles are smaller, thus having a higher tendency to aggregate due to their amorphous state and small size [51]. Excess Mg(II) negatively affects the final powder crystal sizes because it causes a higher supersaturation, consequently increasing the nuclei population density, which suggests a higher nucleation rate. Therefore, crystals reach larger sizes, as described by [52] for Hap crystallization in the presence of excess of Mg(II) ions. The obtained precipitates consist of a population of nanometre-sized primary particles and a population of micrometre-sized aggregates. The aggregates most likely result from the aggregation of primary nanoparticles because small particles have a high surface area-to-volume ratio, resulting in a high surface tension, which tends to diminish by adhering to other particles. [53].

4. Conclusions

In this study, the influence of Mg(II) ions on phosphate recovery by Hap precipitation from basic solutions with desalinated industrial brines containing mixtures of Ca and Mg was confirmed. For both of the Mg/Ca (2.2 and 3.3) brines at higher pH values (9.5 and 11.5), the precipitation inhibition of Hap was observed, and although solutions were supersaturated, the process proceeded with the formation of typically amorphous mineral phases (e.g. Ca, Mg, and Ca-Mg-phosphates). The presence of meta-stable clusters decreases the energetic barrier, thereby facilitating nucleation (pH 9.5 and 11.5). In the case of experiments at pH = 8, formation of stable
clusters increased the barrier, thus promoting nucleation inhibition, and in this case, nanocrystals were formed, and solids were not recovered after filtration with a 0.22 μm filter.

The presence of pre-nucleation clusters in under-and super-saturated solutions and their participation in the phase separation process were proposed. Mg(II) severely inhibited precipitate crystallinity and the precipitation rate, allowing formation of ACP. This result is presumably due to Mg(II) incorporation into the Ca(II)-phosphate structure to form a Mg(II)-substituted structure that crystallized to stanfieldite $\text{Ca}_4\text{Mg}_5(\text{PO}_4)_6$ upon thermal treatment to increase crystallinity.

According to literature, this mineral has not been postulated previously in the precipitation of phosphate solutions with Mg/Ca brines. The surface adsorption of Mg(II) (rather than the incorporated Mg(II)) played a critical role in regulating the transformation rate of ACP to Hap.

Mg(II) altered the stability of the mineral phases, and the more soluble solids were precipitated (e.g., $\text{Mg}_3(\text{PO}_4)_2.22\text{H}_2\text{O}$) at pH 11.5. Sulfate ions have a high capacity to form complexes with Ca(II) and Mg(II), and participated in the precipitation reactions. However, although sulfur was detected by EDX, no mineral containing sulfate was identified by XRD.

The use of industrial desalinated brines containing mixtures of Cd and Mg could be a suitable source for the recovery of phosphate in the form of mixed Ca-Mg phosphates suitable for the chemical industries producing fertilizers.

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5. References


Appendix A. Fundamental precipitation inhibition effects on nucleation growth kinetics

Lamer and Dinegar [1] described the formation of colloidal nanocrystals in a solution phase through a crystal nucleation process involving the following three steps: i) ions start to aggregate into nuclei via self-nucleation as the monomer concentration increases in the solution to supersaturation levels, ii) monomers continuously aggregate on the pre-existing nuclei or seed, which leads to a gradual decrease in the monomer concentration, and iii) nuclei grow into nanocrystals of increasingly larger sizes until reaching an equilibrium state.

The nucleation and growth steps are two relatively separated processes, and the formation of nuclei occurs only at a reactant concentration substantially higher than the saturation concentration (C_s); otherwise, growth of the existing nuclei dominates. The subsequent growth steps will strongly govern the final morphology of the nanocrystals [2], [3] and [4].

The free energy change required for the formation of nuclei (ΔG) is determined by the free energy change for the phase transformation (ΔG_v) and the free energy change for the formation of a solid surface (ΔG_s) [5]. Then, the driving force (Δµ) required for Hap (Ca_5(PO_4)_3OH(s)) crystallization is defined by Equation A1, as follows [6]:

\[
\Delta \mu = kT \ln \left(1 + \sigma \right) = kT \ln \left[ \frac{a \left(Ca^{2+}\right)^{5} \left[a \left(PO_4^{3-}\right)^{3} \left[a \left(OH^{-}\right)\right]^{1}}{K_{so}(Hap)} \right] \right)
\]

where K is the Boltzmann constant, T (K) is the absolute temperature, K_{so} is the solubility product, \( a \) is the activity of species i, and \( \sigma \) is the relative solution supersaturation index.

At a given Δµ, natural nucleation is a kinetically controlled process in which the Hap nuclei overcome a homogeneous nucleation barrier (ΔG_{homo}^*) [6] that could be estimated by Equation A2, as follows:

\[
\Delta G_{homo}^* = \frac{16\pi \gamma_f^{3} \Omega^{2}}{3kT \ln \left(1 + \sigma \right)}
\]

(A2)
where $\gamma_{cf}$ is the specific interfacial free energy between the crystals and the mother phase, and $\Omega$ is the volume of the growth units.

The nucleation induction time ($t_s$) [7], [8] and [9] at different supersaturation levels could be used to characterize the kinetics of nucleation and could be calculated by Equation A3, as follows:

$$ln t_s = \frac{k_n f(m)}{[ln(1+\sigma)]]^2} - ln(VR_s^2 N_0 f'(m)[f(m)]^{1/2}B) \quad (A3)$$

where $R_s$ is the crystal radius, $N_0$ is the mineral density, $B$ is the kinetic constant, $V$ is the solution volume, and $m$ is a factor that depends on the interaction and interfacial structural match between the crystalline phase and substrate; it is expressed as a function of the interfacial free energy difference among the different phases, as shown in Equation A4 as follows [10]:

$$m = \frac{\gamma_{sf} - \gamma_{sc}}{\gamma_{cf}} \quad (-1 < m < 1) \quad (A4)$$

where $\gamma_{sf}$, $\gamma_{sc}$, and $\gamma_{cf}$ correspond to the interfacial tension between substrate and fluid, crystal and substrate, and crystal and fluid, respectively.

Furthermore, $f(m)$ is the interfacial correlation factor describing the reduction of the nucleation barrier $\Delta G^*_{homo}$ due to the occurrence of the substrate and is defined by Equation A5; $f''(m)$ is the pre-exponential term describing the ratio between the average effective collision in the presence and absence of substrate and is defined by Equation A6. Finally, $k_o$ is the nucleation constant, which remains constant under a given condition $m$, and is defined by Equation A7.

$$f(m) = \frac{1}{4} (2 - 3m + m^3) \quad (A5)$$

$$f''(m) = \frac{1}{2} (1 - m) \quad (A6)$$

$$k_n = \frac{16\pi Y_2^2 \Omega^2}{3(KT)^3} \quad (A7)$$
For a crystalline phase \( m \), \( f(m) \) takes only those values corresponding to some crystallographically preferred orientations; then, it is possible according to Equation A3, to obtain a set of intercepting straight lines by plotting \( \ln t_s \) versus \( 1/\ln (1+\sigma) \). These lines with different slopes \( k_n f(m) \) in the different regimes indicate that nucleation is governed by a sequence of progressive heterogeneous processes, as described by Lamer and Dinegar (1950) [1].

**Appendix references**


Table 1. Composition of industrial desalinated brines used in this study.

<table>
<thead>
<tr>
<th></th>
<th>Ca (g Ca(II)/L)</th>
<th>Na (g Na(I)/L)</th>
<th>Cl (g Cl⁻/L)</th>
<th>SO₄²⁻ (g SO₄²⁻/L)</th>
<th>Mg⁺ (g Mg(II)/L)</th>
<th>pH&lt;sub&gt;initial&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mg/Ca (2.2) brine</td>
<td>0.23</td>
<td>23.5</td>
<td>34.2</td>
<td>3.4</td>
<td>0.30</td>
<td>8.8</td>
</tr>
<tr>
<td>Mg/Ca (3.3) brine</td>
<td>0.41</td>
<td>18.9</td>
<td>16.4</td>
<td>20.1</td>
<td>0.85</td>
<td>8.9</td>
</tr>
</tbody>
</table>

*The Mg content is low because Mg(II) was recovered as Mg(OH)₂*
Figure captions:

Figure 1. Effect of pH on a) the P(V) concentration variation and b) the P(V) recovery by precipitation using the Mg/Ca (2.2) brine, c) the P(V) concentration variation and d) the P(V) recovery by precipitation using the Mg/Ca (3.3) brine (dotted line represents the expected P(V) concentration if any precipitation.

Figure 2. Variation concentration of major components (Mg(II), Ca(II), SO$_4^{2-}$ and Cl$^-$) in experiments under different pH conditions using the Mg/Ca (2.2) brine (solid lines are the total ion concentration added throughout the precipitation experiment).

Figure 3. Saturation index (SI) for several minerals in the reactor for brine (Mg/Ca=2.2) at a) pH 11.5 for Ca-phosphate minerals, b) pH 8 for Ca-phosphate minerals and c) pH 8 and 11.5 for Mg-phosphate minerals.

Figure 4. XRD spectra of the particles obtained in the stirred batch reactor with Mg/Ca (2.2) brine a) ACP at pH 9.5 and 11.5 and b) Crystal solid at different pH values after thermal treatment.

Figure 5. Variation of major components (Mg(II), Ca(II), SO$_4^{2-}$ and Cl$^-$) in the batch experiments under different pH conditions using the Mg/Ca (3.3) brine (solid lines are the total ion concentration added along the precipitation experiment).

Figure 6. XRD spectra of the particles produced in the stirred batch reactor with Mg/Ca (3.3) brine at a) pH 11.5 and 9.5 and b) pH 9.5 amorphous solid and c) pH 9.5 after thermal treatment of amorphous precipitates.

Figure 7. Supersaturation index (SI) for Ca/Mg (2.2) brine with respect to Hap at different pH values (8, 9.5 and 11.5) as a function of precipitation time in the batch reactor.

Figure 8. Evaluation of the nucleation kinetics using the dependence of ln(t$_n$) versus $1/[\ln (1+\sigma)]^2$ [47] for Hap nucleation with Mg/Ca (2.2) brine at different pH values (8 and 9.5).
Figure 1.

Figure 1.
Figure 2.
Figure 3.
Figure 4.
Figure 5.
Figure 6.

a

b

2 Theta (Deg.)

2 Theta (Deg.)

Amorphous solid

1/ pH 9.5

1/ pH 9.5

2 Theta (Deg.)

2 Theta (Deg.)

C

Thermal treatment of Amorphous solid

1/ pH 9.5

Stanfieldite
Figure 7.
Figure 8.