Precipitation Process of Calcium Phosphate from Calcium Carbonate Suspension†

Nayane Macedo Portela da Silva, Fabienne Espitalier * and Ange Nzihou
I Université de Toulouse, Centre RAPSODEE UMR CNRS 2392, France

Abstract

The Ca-HA is synthesized by reacting calcium carbonate (CaCO₃) and ammonium dihydrogen orthophosphate (NH₄H₂PO₄) in stoichiometric proportions. In this precipitation process, the ratio of Ca/P by moles is 1.67. The experiments were performed in a batch reactor at 25 °C. From control parameters, the pH and the temperature were measured in line for all experiments. The tests were performed with two liquid-solid mass ratios (H₂O/CaCO₃) of 3 and 5. For each mass ratio, three stirring rates of 260, 400, and 600 rpm were tested. Samples of the synthesis were collected at different intervals and analysed by laser granulometry and by environmental scanning electron microscopy.

For the synthesis conducted at 260 rpm, synthesis monitoring was made using a contact probe in solution coupled to a Raman spectrometer in order to follow the formation of solid phase. This technique is valuable to follow the synthesis of Ca-HA in a concentrated solids suspension (around 20–30 wt%).

The results make it possible to propose a mechanism of the precipitation process of Ca-HA. It can be divided into four main stages: (i) dissolution of calcium carbonate (CaCO₃), (ii) precipitation of brushite (CaHPO₄·2H₂O), (iii) transformation of brushite into Ca-HA (Ca₁₀(PO₄)₆(OH₂)), and (iv) nucleation, growth and agglomeration of Ca-HA.

Keywords: precipitation, hydroxyapatite, brushite, Raman spectroscopy

1. Introduction

The exponential population growth has led to a similar increase in requirements for resources, energy, food, housing and land as well as an enormous increase in industrialization and urbanization. This rapid industrialization and urbanization have led to great demand for natural resources and created an increase in waste production. A continuing issue facing society today is the disposal of municipal and industrial wastes (Wey et al., 2006). Waste calcium carbonate (CaCO₃) from the paper, cosmetics, pharmaceuticals, catalyst, ceramics, rubber and paint industries often ends up in landfill (Wen et al., 2007). These CaCO₃ wastes can be used as the calcium compound in the synthesis of hydroxyapatite (Ca-HA). Ca-HA, Ca₁₀(PO₄)₆(OH₂), is an insoluble calcium phosphate mineral. It is a main constituent of bones and teeth. Due to its excellent biocompatibility, slow biodegradation, good mechanical stability, great sorption properties and heterogeneous photocatalytic degradation under UV irradiation (Wang et al., 2003), it has been used extensively for bone repairs, bone implants, bioactive materials and for purification and separation of biological molecules and organic contaminants. It can accept a series of cationic and anionic substitution within its structure (Elliott, 1994). Similarly, it has been reported to be an efficient adsorbent to treat heavy metals (Chen et al., 1997) and to have a very high capacity for removing divalent heavy metal ions (Ma et al., 1993). Consequently, hydroxyapatite is used for the removal of heavy metals from contaminated soils, wastewater and fly ashes (Elliott, 1994). Several methods have been developed to prepare Ca-HA, including a dry process, precipitation, hydrolyzation of calcium phosphate and hydrothermal synthesis. Spray pyrolysis, freeze drying, gel diffusion, a sol-gel technique and electrochemical deposition have all also been used. Much work has been done on the synthesis of Ca-HA using different methods such as solid-solid reaction (Nakamura et al., 2001; Silva et al., 2003), ultrasonic spray freeze-drying (Itatani et al., 2000) or spray-drying techniques generating a Ca-HA powder of controlled morphology (Luo and Nieh, 1996), sol-gel technique (Anee et al., 2003; Bezzi et al., 2003; Salimi et al., 2012), precipitation in aqueous solution (Jarcho, and Bolen, 1976; Rodriguez-Lorenzo et al., 2001; Bailliez and Nzihou, 2004; Swain et al., 2012), emulsion

† Received 13 May 2014; Accepted 10 December 2014
J-STAGE Advance published online 28 April 2015
1 Ecole des Mines d’Albi-Carmaux, Campus Jarlard F-81013 Albi Cedex 09, France
* Corresponding author: Fabienne Espitalier; E-mail: fabienne.espitalier@mines-albi.fr
TEL: +33-5-63493151 FAX: +33-5-63493025

©2016 Hosokawa Powder Technology Foundation
Original Research Paper
of CaCO₃ was introduced into a 0.5 L glass vessel (diameter 10 cm) with double jacket equipped with a helicoidal stirrer (diameter 4.5 cm). Agitation was powered by a Eurostar IKA labortechnick 2000 motor. The reactor was equipped with four baffles (Fig. 1). The NH₄H₂PO₄ powder was dissolved for a minimum time of 30 min at ambient temperature. Then, this solution was gradually pumped into the reactor with a peristaltic pump (Master flex pump) at a flow rate of 11.11 cm³/min in order to maintain the pH value higher than 7. The temperature was controlled by a water bath (Julabo VC F30). The pH value was monitored throughout the duration of the experiment using a pH meter (Meterlab PHM210). The reaction continued for 72 h at atmospheric pressure and temperature (25 ± 2 °C). The ratio of Ca/P by moles was 1.67. Two different liquid-solid mass ratios (water/CaCO₃ solid) of 3 and 5 and three stirring rates (260, 400 and 600 rpm) were tested. Addition times were equal to 18.6 min and 18.0 min, respectively, for a mass ratio of 3 and 5.

2.2 Analytical methods

The reactor suspension, composed of calcium carbonate and calcium phosphates, was sampled at different intervals in order to characterize the solid and the solution.

Particle-size distributions: The particle size distributions were measured by laser diffraction with a Malvern Master Sizer 2000 instrument covering a size range of 0.1–2000 μm. Suspension samples were dispersed in water by ultrasonication for 4 min before measurement to disaggregate the solid. Direct sampling of the reactor suspension was used for these measurements.

For the other analysis, the collected samples were filtered. Solids were washed and dried in a vacuum oven (Heraeus Vacutherm—VT-6025) at 60 °C for 24 hours to yield a fine white powder. The dry solid was not very cohesive and it was simply broken up by manual grinding.

Ion chromatography was used to analyse the liquid, and TG-DSC and ESEM for the solids.

Ion phosphate concentration: The phosphate ion concentration was analysed using ion chromatographic separation and it is performed using a Dionex ICS-300 ion chromatograph system consisting of: a single pump (Dionex ICS-3000 SP), an eluent generator (Dionex ICS-3000 EG), a conductivity detector (Dionex Conductivity Detector P/N 061830), a precolumn (Dionex IonPac AG 19, 4 × 50 mm) and a column (Dionex IonPac AS 19 (4 × 250 mm). The eluent was the ultra-pure water with the pH value kept constant by using the KOH eluent generator. The eluent flow was 1 ml/min corresponding to a stable pressure in the column of about 120 bar. The results were analysed by using the software Chromelon.

Habitus: A scanning electron microscope (Philips XL30 ESEM FEG) equipped with EDAX fluorescence analysis following platinum sputtering of the sample, was used to study the morphology of the gold-coated intermediate and final products.

Weight losses: Thermal gravimetric analysis was performed using a Thermogravimmetrical Analyser (TG-DSC 111 Setaram). The experiment was performed with a heating rate of 2 °C/min using nitrogen flow (25 ml/min). The percentage weight loss of the samples (approximately
Using PIXE was monitored from 20 to 800 °C. Raman spectra; For the synthesis conducted at 260 rpm, the synthesis monitoring was conducted using a contact probe in solution coupled to a Raman spectrometer RXN2C785, Mettler-Toledo/Kaiser analysers (Kaiser Optical Systems, Ann Arbor, Michigan, USA). A 400 mW external cavity stabilized by an Invictus NIR diode laser at 785 nm was used for sample illumination. All the spectra were recorded at a resolution of 4 cm⁻¹.

XR diffraction: For the synthesis conducted at 260 rpm, the phase identification of some samples was carried out using XR diffractometry (Phillips Panalytical X'Pert Pro MPD diffractometer), which had a Cu Ka radiation (l = 1.54 A) at 45 kV and 40 mA. The reflections were collected in the 2θ ranges from 10 to 75° with a step size of 0.17° and a time of 13.02 s per step. The phase identification was carried out with the JCPDS database.

3. Results

During the process for the formation of hydroxyapatite, it was possible to form other calcium phosphates. Their solubilities decrease with the pH value (Ferreira et al., 2003). The comparisons of the solubility of the CaHPO₄·2H₂O named brushite and the solubility of hydroxyapatite Ca-HA for a pH value lower than 4, show that brushite is a stable compound and for a pH value higher than 4, the Ca-HA is prominent. Fig. 2 shows the evolution of the pH value during the synthesis for the two liquid-solid mass ratios (3 and 5) at 400 rpm. It can be observed that the pH value readily decreases from 10.2 to 6.9 during the first 15 minutes of reaction. Then it gradually increases to 8.5 and after 40 hours, the pH value stabilizes at about 9. The liquid-solid mass ratio does not influence the pH value.

At 25 °C in the pH range [8-9] (Fig. 2), it was possible to form three solids: the monetite (CaHPO₄), the brushite (CaHPO₄·2H₂O) and the hydroxyapatite (Ca₁₀(PO₄)₆(OH₂)).

Their formation follows these three reactions respectively (Chkir, 2011; Verwilghen, 2009):

1. Monetite (metastable phase):
   $$\text{CaCO}_3 + \text{NH}_4\text{H}_2\text{PO}_4 \rightarrow \text{CaHPO}_4 + \text{CO}_2 + \text{H}_2\text{O} + \text{NH}_3$$

2. Brushite:
   $$\text{CaCO}_3 + \text{NH}_4\text{H}_2\text{PO}_4 + \text{H}_2\text{O} \rightarrow \text{CaHPO}_4\cdot2\text{H}_2\text{O} + \text{CO}_2 + \text{NH}_4\text{OH}$$

3. Hydroxyapatite:
   $$10\text{CaCO}_3 + 6\text{NH}_4\text{H}_2\text{PO}_4 \rightarrow \text{Ca}_{10}(\text{PO}_4)\text{O}_6(\text{OH}_2) + 10\text{CO}_2 + 2\text{H}_2\text{O} + 6\text{NH}_4\text{OH}$$

4. Transformation of brushite into hydroxyapatite
   $$10\text{CaHPO}_4\cdot2\text{H}_2\text{O} + 4\text{NH}_4\text{OH} \rightarrow \text{Ca}_{10}(\text{PO}_4)_3(\text{OH}_2) + 4\text{NH}_4\text{H}_2\text{PO}_4 + 18\text{H}_2\text{O}$$

The analysis did not detect monetite. When any amount of ammonium dihydrogen phosphate is introduced into the reactor containing the calcium carbonate suspension, gas bubbles with an ammonium odor are observed. This confirms the formation of carbon dioxide and ammonium during the reaction.

As the pH showed low values at the beginning of the synthesis, this may have favored the formation of brushite. The end of synthesis, which sees the stabilization of the pH value at around 9, favors the formation of hydroxyapatite.

3.1 Influence of mass ratio

Two syntheses with two liquid-solid mass ratios of 3 and 5, respectively, carried out at a low stirring rate of 260 rpm in the presence of a Raman probe to study the influence of this ratio on the synthesis of Ca-HA.

Tsuda and Arends (1993) give the Raman bands observed for three pure minerals (brushite, octacalcium phosphate OCP and Ca-HA). The spectral features of solids can be characterized as Raman bands due to phosphate (PO₄³⁻) vibrational modes of ν₄, ν₂, ν₁ and ν₃. The intensity of the ν₁ band is the strongest, and can be used for the identification of minerals.

Figs. 3, 4 and 5 show the Raman spectra during the synthesis made with a mass liquid-solid ratio of 5 and a stirring rate of 260 rpm. In Fig. 3, we observe the appearance of a peak at 983 cm⁻¹ after 20 min of synthesis. This peak is relatively wide. Its intensity increases with time. This peak noted on Fig. 4 corresponds to the brushite, a compound observed in previous studies (Chkir, 2011).

The presence of brushite is confirmed by XR diffraction analysis (Fig. 6).

After 19 h, a second peak close to the brushite peak appears at 962 cm⁻¹; this corresponds to the Ca-HA. While the peak intensity of brushite decreases, the peak inten-
The intensity of Ca-HA increases (Figs. 4 and 5). After 30 h, the intensities of all peaks are stable. This assumption has been verified qualitatively by XR diffraction analysis (Fig. 7). At 44 h, the brushite is no longer visible by this technique. The large peak observed at 30–33° corresponds to Ca-HA.

Fig. 3 Raman spectra measured every 19 min between 0 and 3 h, (liquid-solid mass ratio = 5 and N = 260 rpm). Times from the bottom to the top.

Fig. 4 Raman spectra measured every 4 h between 3 h 10 min and 24 h 19 min, (liquid-solid mass ratio = 5 and N = 260 rpm). Times from the bottom to the top.

Fig. 5 Raman spectra measured every 8 h between 19 h 20 min and 91 h 20 min, (liquid-solid mass ratio = 5 and N = 260 rpm). Times from the bottom to the top.

Fig. 6 XR diffraction spectra at 0.75, 2 and 4 hours (liquid-solid mass ratio = 5 and N = 260 rpm). Times from the bottom to the top.

Fig. 7 XR diffraction spectra at 44, 50 and 68 hours (liquid-solid mass ratio = 5 and N = 260 rpm). Times from the bottom to the top.

Fig. 8 Weight losses between 20 and 800 °C at 0, 0.75, 4, 20, 44, 50, 68 and 72 h (liquid-solid mass ratio = 5 and N = 260 rpm).

cium carbonate, two weight losses are observed: one between 350 and 400 °C and another after 600 °C. The first loss is due to the degradation of impurities contained in CaCO₃. The decomposition of calcium carbonate (CaCO₃) in calcium oxide (CaO) and carbon dioxide (CO₂) (decarbomation) induces the second loss.

For the other samples, three weight losses are observed.
The first is between 180 and 200 °C, the second between 350 °C and 400 °C and the third after 600 °C. This first is attributed to loss of the hydration water of brushite (detected by RAMAN spectra analysis), the second to impurities and the third to the decarbonation of unreacted CaCO₃. The weight loss observed between 50 and 120 °C is due to insufficient drying of these samples. From these mass losses, by assuming that only three solids are present in the reactor (CaCO₃, CaHPO₄·2H₂O and Ca₁₀(PO₄)₆(OH₂)), the mass percentage of these three components in the solid can be calculated (Fig. 9).

Four steps can be observed. The first step (I) corresponds to the dissolution of calcium carbonate and formation of apatite: a sharp decrease in calcium carbonate (around 40 %) during the first hour, and a sharp increase in brushite (around 40 %) and Ca-HA (around 20 %).

During the second step (II) between 4 and 20 h, the weight percentage of solids seems to be constant. After 20 h, we can observe an increase in the weight percentage of Ca-HA up to 80 wt% and a decrease in the CaCO₃ and brushite weight percentages. This step (III) corresponds to the transformation of brushite into Ca-HA by dissolution-crystallization or by solid-solid transformation. However, the reaction is not complete and 20 wt% of the solid remaining is composed of calcium carbonate. After 68 h, two solids are in suspension: CaCO₃ and Ca-HA. The mass balance on calcium leads to the total weight as a function of time from mass fractions and initial weight of CaCO₃:

\[
m(t) = \frac{m_{\text{CaCO}_3} \cdot \frac{1}{M_{\text{CaCO}_3} \cdot w_{\text{brushite}} + 10 \cdot w_{\text{HAP}} + w_{\text{CaCO}_3}}}{M_{\text{brushite}} + 10 \cdot M_{\text{HAP}} + M_{\text{CaCO}_3}} (1)
\]

where \( m_{\text{CaCO}_3} \) is the initial weight of CaCO₃, \( M_j \) the molar weight and \( w_j \) the mass fraction of the component \( j \) in the solids mixture.

Fig. 10 presents the evolution of weight of each solid during a synthesis for both liquid-solid mass ratios, 3 and 5. Furthermore, by coupling the results of the loss of mass and the RAMAN spectra, it is possible to calibrate the area of RAMAN peaks observed in 983 and 962 cm⁻¹, respectively, for the brushite and the Ca-HA according to the mass fractions obtained by TGA. This calibration leads to the continuous curves given in Fig. 10. Both weight ratios lead to final masses that are close to CaCO₃ initial masses.

The conversion does not seem to depend on the initial weight ratio. This conversion is not total, but is of the order of 73–74 wt%. We also note that the maximum mass of brushite formed for a liquid-solid mass ratio equal to 3 is 13 times weaker than that formed for a liquid-solid mass ratio of 5 (Fig. 10 (c)).

Beyond 20 h for a liquid-solid ratio of 3 and 40 h for a
liquid-solid ratio of 5, the brushite seems to have totally disappeared. In parallel, the formation of Ca-HA is much faster for an L/S ratio of 3 within the first hours (77 wt% of the final mass is reached (Fig. 10 (a)). After 10 h of synthesis, a slowing down is observed.

The particles of brushite are formed preferably in the solution (Fig. 11), while the particles of Ca-HA are created on the surface of the brushite particles (Fig. 12) or (Fig. 11). The particles of CaCO₃ are attacked by orthophosphate ions. Therefore, CaCO₃ is used to form monetite. This compound is transformed into brushite which evolves to the hydroxyapatite (calcium phosphate thermodynamically more stable). When the liquid-solid mass ratio decreases, the available surface of CaCO₃ is larger, allowing the reaction (3) to occur on the surface of CaCO₃. This behavior limits the brushite formation by reaction (2). The mass growth rate of Ca-HA at the initial time can be written as:

\[
\frac{dm_{CaHA}}{dt} = K S \Delta C^8
\]

(2)

where \(K\) is a kinetic constant, \(\Delta C\) is the absolute supersaturation of Ca-HA, and \(S\) is the CaCO₃ disposable surface at the time at which Ca-HA is detectable.

As the initial size distribution of calcium carbonate is identical, the surface \(S\) is assumed proportional to mass.

The ratio of these initial rates for liquid-solid mass ratios of 3 and 5 leads to a coefficient \(j\) of the order of 12.5, thereby showing a strong influence of the available surface of calcium carbonate on the Ca-HA formation. As the Ca-HA layer forms on the surface of CaCO₃, the diffusion of orthophosphate and calcium ions is limited thus reducing the mass growth formation of Ca-HA observed beyond 10 h for a liquid-solid mass ratio of 3. In the case of a higher ratio, it can be assumed that reactions (2) and (4), which represent the CaCO₃ and brushite formations, respectively, occur in parallel.

Fig. 14 shows the evolution of the mass mean diameter as a function of the time based on both mass ratios (liquid-solid mass ratios of 3 and 5) for a stirring speed of 260 rpm. For a liquid-solid mass ratio of 5, the increase in average diameter (7–22 microns) between 0 and 3 is due to the brushite formation (Fig. 14 (a)). After 3 h, the mean diameter decreases assumed by brushite dissolution. Beyond 20 h, the mean diameter increases following the same trend as Ca-HA mass (Fig. 14 (b)).

Brushite dissolution allows Ca-HA formation. This compound is thermodynamically more stable than brushite (Ferreira et al. 2003) and it forms on the surface of CaCO₃ causing the increase in diameter. For a liquid-solid mass ratio of 3 (Figs. 15 (a) and (b)), the same behavior is observed, meaning that the mass mean diameter increases, then decreases and finally increases again. However, beyond 20 h, the mean diameter does not change (about 20 µm) because the Ca-HA mass stabilizes.
Fig. 16 presents the mean diameters of the particles during synthesis for different stirring speeds (400 and 600 rpm) for both mass ratios (L/S = 3 and 5). The results obtained at 400 and 600 rpm show the same behavior obtained at 260 rpm (increase, decrease, re-increase and stabilization of mean diameter). After 40 h of synthesis, the measured diameters are identical (23 ± 1 μm) for a stirring speed equal to 400 rpm (Fig. 16 (a)). Larger agglomerates are observed at a low stirring speed after 72 h of synthesis. The stabilization of the mean diameter is observed after 40 h for all the syntheses except for the synthesis performed at 600 rpm. In this case after 20 h of synthesis, the mass mean diameter is stable and equals 16 ± 1 μm (Fig. 16 (b)).

4. Conclusions

In this study, we show that Raman spectroscopy is a valuable technique to follow the synthesis of Ca-HA in a concentrated solids suspension (around 20–30 wt%). This technique makes it possible to follow the formation of Ca-HA and brushite and to deduce from complementary measurements a formation mechanism described as fol-
The dissolution of calcium carbonate is observed in the first minutes of synthesis.

- The precipitation of brushite and Ca-HA particles is observed in the first four hours of synthesis.
- The transformation of brushite into hydroxyapatite begins after one day. It seems to be the limiting step.

In terms of morphology, agglomerates of hollow ellipsoidal particles were obtained corresponding to the calcium carbonate, while platelets agglomerated like stars corresponded to the brushite and agglomerated spongy spheres or round stick-like compounds of spongy spheres corresponded to the hydroxyapatite.

Through the mass balance obtained from thermogravimetric analysis and from the phosphate ions, it was also verified that at the end of the synthesis, the calcium carbonate was not completely consumed.

Therefore, the reaction is not complete at the end of the synthesis (72 h).

From the analysis performed by laser granulometry during the synthesis with a mass equal to five and three at 260, 400 and 600 rpm, there was a strong influence of the stirring speed on the process. The two effects (positive and negative) that explain the agglomeration process have been presented. For the synthesis with a mass ratio equal to five, the first effect called the size-enlargement phenomenon predominated. In contrast, for the synthesis with a mass ratio equal to three, a predominance of the second effect called the breakage phenomenon was observed. These phenomena were confirmed by the observations made using the environmental electron microscopy of the synthesized solids.

5. Acknowledgements

The authors would like to thank M. H. Lucas for support in the data analysis of RAMAN spectra. They also thank Ms S. Del Confetto, Ms C. Rolland, Ms N. Lyczko and M.P. Accart, respectively, for TG measurements, microscopy analysis and X-Ray, and particle size analysis.

References


Author’s short biography

Nayane Macedo Portela da Silva

Dr Nayane Macedo Portela da Silva was born in 1986. She completed her graduate studies at UFRN university of Natal, Brazil. Before her diploma, she joined the EMAC school at Albi, France, to do an internship concentrating on hydroxyapatite synthesis by reaction crystallization (precipitation). She prepared a PhD thesis on microfluidic systems under high pressure in the Laboratory Rapsodee of EMAC school and received her PhD from Toulouse University in 2014.

Fabienne Espitalier

Professor Fabienne ESpitalier is the head of the research group “Powder and Processes” at the Research Centre in Albi for Particulate Solids, Energy and Environment (Rapsodee research centre)-CNRS UMR 5302 at Ecole des Mines Albi in France. Her major research interests concern crystallization assisted or not by ultrasound, non-standard solvents such as ionic liquids, population balance simulation, characterization and particle technology.

Ange Nzihou

Professor Ange Nzihou is the Director of the Research Centre in Albi for Particulate Solids, Energy and Environment (Rapsodee research centre)-CNRS UMR 5302 at Ecole des Mines Albi in France. He is also editor in chief of the international peer-reviewed journal Waste and Biomass Valorization (Springer), and chair of the WasteEng conference series. He has developed outstanding research in the field of the engineering of phosphate-based materials for sorbents and catalysts, and heavy metals contaminants remediation. He has published about 87 papers in peer-reviewed journals.