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Nanoscale conversion of chlorapatite into hydroxyapatite using ultrasound irradiation

Habiba Bouyarmane,^a Asmae Gouza,^a Sylvie Masse,^b Sanaa Saoiabi,^a Ahmed Saoiabi,^a Thibaud Coradin,^{b,*} Abdelaziz Laghzizil^{a,*}

^aLaboratoire de Chimie Physique Générale, Université Mohammed V, Faculté des Sciences BP.1014 Rabat, Morocco.

^bSorbonne Universités, UPMC Univ Paris 06, CNRS, UMR 7574, Laboratoire de Chimie de la Matière Condensée de Paris, F-75005 Paris, France.

*Corresponding authors :

- T. Coradin ; e-mail : thibaud.coradin@upmc.fr; tel : 33-144271528 ; fax :33-144274769
- A. Laghzizil; e-mail: laghzizi@fsr.ac.ma; tel/fax: 212-537775440

Abstract: The ultrasound-assisted conversion of chlorapatite nanoparticles obtained from a natural phosphate rock into hydroxyapatite nanocrystals was achieved within one hour. The process had no significant impact on particle crystallinity but led to a significant decrease of the colloidal size. Similar variations were obtained for hydroxyapatite nanoparticles treated in the same conditions, suggesting that these variations are not due to the compositional modification. Analyses of the solubility properties of the two phases and of the ionic contents of the reaction medium suggest that the undersaturated conditions of the reaction and the cavitation phenomena favor surface exchange and erosion mechanisms over bulk amorphization.

Keywords : hydroxyapatite ; sonochemistry; topotactic conversion; nanomaterials

1. introduction

Natural phosphates are one of the major reservoirs of both phosphorus and calcium in the Earth's crust [1]. Fluoroapatite ($Ca_{10}(PO_4)_6F_2$), and its carbonated form francolite, is the main component of Moroccan natural phosphates [2-4]. Among other applications, the cation-exchange properties of phosphocalcic apatites as well as their rich surface chemistry make them valuable materials as sorbents for water remediation [5-8]. However, natural apatites exhibit low specific areas that limit their sorption capacity [9-11]. Therefore many physical and chemical routes have been developed to obtain phosphocalcic apatites as high surface area powders from inorganic salts [12]. Of particular interest here are sonochemistry routes that allow control of the particle size and crystallinity through the modulation of irradiation power and duration [13-21]. As a more sustainable alternative, a process allowing for the conversion of natural phosphate into porous hydroxyapatite ($Ca_{10}(PO_4)_6(OH)_2$) was recently described [22,23]. The procedure involves the dissolution of the natural ore in nitric acid, filtration of the remaining minerals (mainly quartz) and reprecipitation by neutralization.

Although this approach has the benefits of using raw materials available in large quantity and being performed in purely aqueous conditions, the use of nitric acid at a large scale is associated with explosive hazards related to its strong oxidizing power. In this study, we examined hydrochloric acid as an alternative dissolution medium. The precipitated material was a nanopowder of chlorapatite ($Ca_{10}(PO_4)_6Cl_2$), whose reactivity is lower than hydroxyapatite due to the key role of OH⁻ in its surface properties. Topotactic, *i.e.* shape preserving, conversion of chlorapatite and other phosphocalcic apatite into hydroxyapatite has already been described in hydrothermal conditions from well-crystallized powders [24,25]. These approaches required strong alkalinity (KOH, 6.25 M), high temperatures (> 300 °C) and relatively long reaction times (> 12 h) but allowed for conversion of large crystals (several mm). Alternatively, for nanoscale particles with high interfacial reactivity and easy dispersion, topotactic transformations of calcium phosphates using ultrasound irradiation appears as a suitable and fast method [26,27]. On this basis, the effect of sonication on the topotactic transformation of chloroapatite nanocrystals into hydroxyapatite was demonstrated and the mechanisms by which the conversion can be influenced by ultrasound irradiation are discussed. In particular, the key role of undersatured conditions on the ultrasound-induced transformation is evidenced. These results demonstrate that the structural conversion can be achieved without particle amorphization but with size reduction, offering a simple and original method to obtain hydroxyapatite nanoparticles with tailored crystallinity and dimensions.

2. Materials and methods

2.1 Materials

The phosphate rock sample used was from the Bengurir region (Morocco). The sample was washed and then sieved to give a $-400/+100\mu$ m size fraction using ASTM Standard sieves, this grain size being optimum for efficient dissolution. As reported earlier [4], it consists of a mixture of francolite and quartz (2.5 wt%) and its composition in major mineral elements is provided in Table 1.

2.2 Synthesis

The conversion of the natural ore into nanosized powders was performed according to a previously described procedure [23]. Dissolution of the phosphate rock was carried out in a 1 L open glass reactor with a spherical bottom at a rate of 200 rpm. For each experiment, the temperature was set to 25°C. The phosphate rock (20 g) was first dissolved in 80 mL of deionized water supplemented with 20 mL of concentrated hydrochloric acid (37%) or nitric

acid (65%) to obtain the solvated Ca²⁺ and hydrogenophosphate species. Insoluble matter, mainly quartz, was then separated by filtration. The remaining solution was neutralized using 100 mL of a concentrated NH₄OH solution (25%). The pH value of the precipitation reaction was maintained at pH 10 to avoid the formation of other non-apatitic calcium phosphates. The resulting powder was recovered by filtration and either washed several times with deionized water or re-dispersed in 100 mL deionized water under stirring for 30 min and then placed in an ultrasonic water bath (200 W, 35 kHz) for 1 hour. The resulting products were recovered by filtration and thoroughly washed with water. Both set of samples were dried overnight at 100°C. The precipitated samples are noted Cl-CP and N-CP according to the inorganic acid (HCl and HNO₃) used for dissolution; uCl-CP and uN-CP correspond to the powders obtained from sonication. Further heating of the samples was performed at 800°C. The overall flow diagram of the process is summarized in Figure 1.

2.3. Characterization. The crystalline phases were identified using a powder X-ray diffractometer (XRD) (Philips PW131 diffractometer) using the CuK α radiation. Infrared spectra were recorded from 400 cm⁻¹ to 4000 cm–1 on a Bruker IFS 66v Fourier transform spectrometer using KBr pellets. Particles were imaged using Transmission Electron Microscopy (TEM) on a TECNAÏ G2 instrument. Calcium and phosphorus content of the powders were determined by inductively coupled plasma atomic emission spectroscopy (ICP-AES) (ICPS-7500, Shimadzu, Japan). Chloride content was determined using the Mohr method. The N₂ adsorption-desorption isotherms for dried powders were obtained by multipoint N₂-gas adsorption-desorption experiments at 77 K using a Micromeritics ASAP 2010 instrument. The specific surface areas were calculated according to the Brunauer-Emmett-Teller (BET) method using adsorption data in the relative pressure range from 0.05 to 0.25 whereas the pore size and volume were estimated using the Barret-Joyner-Halenda (BJH) approximation.

3. Results

The X-Ray diffractograms of non-sonicated Cl-CP and N-CP dried at 100°C and heated at 800°C are shown in Figure 2a. The dried Cl-CP and N-CP samples exhibit a series of very similar low intensity peaks but differ from a multi-component signal in the $2\theta = 31-34^{\circ}$ range for the former and 32-35° domain for the latter. After heating at 800°C much narrower diffraction peaks are obtained allowing for complete resolution of these complex signals (Figure 2c,d). The replacement of OH^{-1} ions in hydroxyapatite $Ca_{10}(PO_4)_6(OH)_2$ (HAp) by CI^{-1} in chloroapatite $Ca_{10}(PO_4)_6(Cl)_2$ (ClAp) was reported to result in an increase of the a parameter (from 9.418 Å to 9.641 Å) and decrease in c parameter (from 6.884 Å to 6.771 Å) of the apatite structure [24]. These variations can easily be visualized by the shift of the (211) diffraction peak shift from 31.8° to 31.3° and the (300) from 32.8° to 32.3°. The (112) peak that is clearly evidenced for HAp at 32.2° is shifted to 32.4° in ClAp, becoming merged with the (300) (Figure 2b). Altogether, it is possible to assume that N-CP sample consists of HAp and Cl-CP of ClAp although the presence of minor amounts of HAp in the later cannot be fully excluded. As a comparison, sonicated samples obtained from natural phosphate dissolution in HCl (uCl-CP) and HNO₃ (uN-CP) showed only diffraction peaks corresponding to the HAp structure (Figure 2b). The only noticeable difference is the lower resolution of the 32-35° broad diffraction signal for uCl-CP at 100°C compared to uN-CP at the same temperature, indicative of its lower crystallinity. Importantly, no significant difference was observed between N-CP and uN-CP after drying.

Elemental analysis of the samples shows that the Cl-CP sample contains ca. 6 % of Cl (Table 2). The theoretical atomic % of Cl for the chloroapatite formula is 6.9 %, allowing to estimate that the powder is a mixture of 85 % ClAp and 15 % HAp. The Cl amount decreased to 0.05 % for uCl-CP, corresponding to less than 5 wt% remaining ClAp or formation of a

chloride substituted hydroxyapatite $Ca_{10}(PO_4)_6Cl_{2x}(OH)_{2(1-x)}$ with $x \approx 0.007$. As far as the Ca/P ratio is concerned, it is found superior than the stoichiometric ratio 1.67 of pure HAp and ClAp, supporting the formation of carbonated apatites. However, a decrease in the Ca/P ratio is observed after sonication for both HCl- and HNO₃-treated samples, suggesting that the degree of carbonatation is smaller after irradiation.

FTIR spectra of selected samples are shown in Figure 3. For dried powders, the main difference between the Cl-CP and uCl-CP samples is the vibration band at 3560 cm⁻¹ corresponding to OH groups of the HAp structure that is evidenced only in the uCl-CP spectra. This difference is even more clearly visible on the 800°C-heated samples, where the corresponding band is more easily visible for uCl-CP due to the decrease of the broad band attributed to the vibration of bound water. This is in agreement with the substitution of Cl⁻ by OH during the sonication process. Noticeably, at this temperature, uCl-CP and N-CP are almost undistinguishable, in agreement with XRD data. The vibration bands in the 1020-1100 cm^{-1} wavenumber range that correspond to PO₄ groups show a strong tendency to saturate for N-CP and uCl-CP at 800°C, in agreement with high degree of crystallinity of the HAp phase. At the same temperature, this band is broader for Cl-CP, reflecting the coexistence of HAp and ClAp phases in the powder. FTIR spectra of dried powders showed additional bands near 1450 cm⁻¹ and 1650 cm⁻¹ that almost disappear from the spectra of the heated samples. The former corresponds to carbonate ions incorporated in the apatite structure while the latter indicates adsorbed water molecules, both of which should be withdrawn from the apatite after dehydration and decarbonation.

TEM images of the dried powders are shown in Figure 4. Cl-CP and uCl-CP samples consist of well-defined nanoparticles of similar morphologies but different lengths, i.e. 42 ± 6 nm for Cl-CP compared to 19 ± 5 nm for uCl-CP. As a comparison, N-CP and uN-CP

powders consist of similar particles although with an apparent lower aspect ratio compared to HCl-precipitated samples. Particle length obtained from TEM image was 29 ± 3 nm for N-CP and 16 ± 3 nm for uN-CP.

After heating, all samples consisted of densely-packed aggregates or fused nanoparticles with various morphologies (Figure 5). Although a precise measurement of particle size is difficult from such TEM images, Cl-CP and uCl-CP give average diameter of 51 ± 11 nm and 44 ± 8 nm, respectively, whereas N-CP and uN-CP gives 70 ± 8 nm and 58 ± 8 nm respectively. This suggests that the initial size difference between sonicated and unsonicated samples is preserved upon heating, at least to some extent.

Nitrogen sorption isotherms of dried powders are gathered in Figure 6 and indicate that all powders are mesoporous. Specific surface area calculated using the BET model (S_{BET}) show that Cl-CP has the smallest value while uCl-CP and uN-CP have the highest, with N-CP being intermediate (Table 2). Although calculation of porous volume (Vp) and average pore size (Dp) of powders must be taken cautiously due to particle displacement during nitrogen filling, estimated values indicate close Vp values for all powders and significant variations in Dp. This means that the density of each powder is similar but that the difference in accessible surface is due to a variation in pore size. The obtained values are in good agreement with TEM imaging as the values of calculated S_{BET} follow the same trend as particle size evolution. This supports the assumption that the measured mesoporosity does not originate from the internal structure of the particles but from interparticle spacings in the powder form.

4. DISCUSSION

The full dissolution of 20 g of the natural phosphate in 100 mL of a 2.4 M HCl solution leads to final concentrations of $[Ca^{2+}] = 2$ M, $["PO_4"] = 1.2$ M and $[Cl^-] = 2.4$ M, where $["PO_4"]$ represent the total phosphate content, *i.e.* $H_xPO_4^{3-x}$ species. Increasing the pH to 10

by adding 100 mL NH₄OH leads to $[OH^-] = 10^{-4} \text{ M}$, $[Ca^{2+}] = 1 \text{ M}$, $["PO_4"] = 6.10^{-1} \text{ M}$ and Cl^-

= 1.2 M. The reaction of chlorapatite formation is (eq. 1) [28]

$$10 \text{ Ca}^{2+} + 6 \text{ PO}_4^{3-} + 2 \text{ Cl}^- \Leftrightarrow \text{Ca}_{10}(\text{PO}_4)_6 \text{Cl}_2$$
(1)

with a solubility constant K_{sc} at 25 °C of 10⁻¹¹⁵

The reaction of hydroxyapatite formation is (eq. 2)

$$10 \text{ Ca}^{2+} + 6 \text{ PO}_4^{3-} + 2 \text{ OH}^- \Leftrightarrow \text{Ca}_{10}(\text{PO}_4)_6(\text{OH})_2$$
(2)

with a solubility constant K_{SH} at 25 °C of 10⁻¹¹⁰

The ionic products *IP* relevant for ClAp and HAp formation in the conditions of the experiments are

$$IP_{C} = [Ca^{2+}]^{10} [PO_{4}^{3-}]^{6} [Cl^{-}]^{2} = 6.7 \ 10^{-2}$$
(3)

$$IP_{H} = [Ca^{2+}]^{10} [PO_{4}^{3-}]^{6} [OH^{-}]^{2} = 4.6 \ 10^{-10}$$
(4)

Supersaturation *S*, calculated as the ratio between the ionic product and solubility constant *IP/K*, is therefore $S_C = 10^{+113}$ and $S_H = ^{+100}$ for ClAp and HAp, respectively. The free enthalpy of the reaction of precipitation can be written as (eq. 5)

$$\Delta G_r = -\operatorname{RTln}S\tag{5}$$

Since S_C and $S_H > 1$ and $S_C > S_H$, the free enthalpy values are negative for ClAp and HAp, indicating that the two reactions are spontaneous, with ClAp precipitation being thermodynamically favored compared to HAp formation

After resuspension in deionized water (pH 7), [Cl⁻] is negligible, shifting the equilibrium (1) towards dissolution, releasing Ca²⁺ and PO₄³⁻ with 10 [Ca²⁺] = 6 [PO₄³⁻]. In these conditions, $S_H > 1$, *i.e.* HAp can reprecipitate, as soon as [Ca²⁺] > 10⁻⁶ M. This is to be compared with the calcium concentration in the resuspension (2 M), suggesting that, despite initial undersaturated ionic conditions, as soon as chlorapatite particles start to dissolve, reprecipitation of HAp can occur (Figure 7a). It is worth pointing out that the conversion of ClAp micron-size single crystals into HAp in high temperature (>1000°C) conditions was

reported to occur via an ion-exchange mechanism [29]. The contribution of such an ionexchange reaction in our working conditions cannot be put aside but the observed accompanying erosion process (vide infra) strongly support that dissolution/reprecipitation mechanisms are prevalent.

Such a crystallographic conversion is associated with a preservation of the morphology of the particles but with a decrease in size from CI-CP to uCI-CP. However, a decrease of particle dimension is also observed with pure HAp particles prepared using HNO₃ after a similar ultrasound treatment. Several previous reports have demonstrated the ability of ultrasound to tune HAp particle size but these studies were devoted to particle growth from Ca^{2+} and phosphate salts [13-21] and often describes a continuous process occurring from an amorphous calcium phosphate phase to crystalline HAp. In these works, decrease of particle size upon irradiation was attributed to the formation of hot spots resulting from cavitation bubbles collapse that favor nucleation events. One recent report also describes the sonicationinduced conversion of brushite CaHPO₄.2H₂O into monocalcium phosphate monohydrate $Ca(H_2PO_4)_2$ ·H₂O via monetite CaHPO₄ formation [26]. These transformations are possible due to the increasing stability of the successive phases but can be favored by the cavitation phenomenon that enhances exchange reactions occurring at the particle surface, especially dehydration that leads to a local increase in supersaturation [30]. The other key effect of ultrasound irradiation is related to its ability to limit particle aggregation [16].

These two effects can be conjointly involved in the here-described process. On the one hand, ultrasound irradiation allows for an efficient dispersion of the particles, not only in the initial solution but also during the phase transformation where the high reactivity of the colloidal surfaces should confer them a strong tendency to aggregate. On the other hand, the supersaturation at the particle vicinity due to efficient surface dissolution should enhance the kinetics of the exchange of Cl⁻ by OH⁻. However it is important to point out that this

exchange is thermodynamically-favorable reaction, a fact that can explain that the use of a lower ultrasound irradiation power (160 W) did not significantly modify the size and structure of the recovered uCl-CP nanoparticles (data not shown).

It is interesting to note that we found no clear evidence of the amorphization of the CP particles upon sonication, contrasting from previous reports [21], but rather a decrease in particle size. This can be explained considering that the previous observations were obtained for particles in their mother liquor, *i.e.* at the solubility limit for the bulk liquid. In this case, the chemical equilibrium between hydroxyapatite and its constituting ions was towards the precipitation process so that the amorphous phase remains in a solid form (Figure 7b). In contrast, in the present work, the sonication process is performed in a salt-free solution so that surface erosion via dissolution and ion release becomes possible (Figure 7c).

Finally, the question is raised about the impact of the ultrasound irradiation of the chemical properties of the HAp nanoparticles. It has been previously shown that the erosion process enhanced the surface reactivity of metal nanoparticles as it allowed for the removal of the external metal oxide layer [31]. In contrast, for hydroxyapatite, it is well-known that the bulk phase is relatively inert [32] whereas the presence of a disordered outer layer makes its surface more reactive [33]. However, the generation of OH⁻ radical species upon water irradiation may impact on the particle surface composition and structure [34]. Therefore a more detailed study of the surface reactivity of these nanoparticles is required to fully understand the influence of ultrasonication on their reactivity

5. Conclusions

This work demonstrates that sonication provides a simple and fast method to achieve the topotactic conversion of chlorapatite into hydroxyapatite at the nanoscale. Such a conversion is made possible by the redispersion of the chlorapatite suspension in deionized water,

allowing for the favorable exchange between Cl⁻ and OH⁻. These conditions also favor erosion over amorphization, therefore providing a method to tailor hydroxyapatite particle size without impacting on its crystallinity, an original route that must be explored further.

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% Ca	% P	% F	%Si	%S	%Na
37.84	15.03	2.84	1.78	0.78	0.79

Table 1. Chemical analysis of the phosphate rock used in this study [4].

Table 2. Elemental analyses, specific surface area (S_{BET}), porous volume (V_p) and average pore diameter (D_p) for dried powders obtained from conversion of natural phosphates before and after ultrasound treatment.

Sample	% Ca	% P	% Cl	Ca/P	S_{BET}	V_p	D_p
					$(m^2.g^{-1})$	$(cm^{3}.g^{-1})$	(nm)
uCl-CP	37.55	16.78	0.05	1.73	165	0.36	7.0
Cl-CP	37.01	15.85	5.99	1.80	120	0.37	10.8
uN-CP	38.05	15.52	< 0.01	1.89	160	0.38	9.0
N-CP	38.66	15.33	< 0.01	1.95	150	0.39	11.5

Figure captions

Figure 1. Flow diagram of the preparation of nanosized apatite samples

Figure 2. XRD patterns of the dried and 800°C-heated powders (a,b) before and (c,d) after ultrasound irradiation. (OH) and Cl correspond to hydroxyapatite and chloroapatite structures, respectively

Figure 3. FTIR spectra of selected powder samples.

Figure 4. TEM images of the dried powders before and after ultrasound irradiation: (a) Cl-CP, (b) uCl-CP, (c) N-CP and (d) uN-CP (scale bar = 20 nm).

Figure. 5. TEM images of the 800°C-heated powders before and after ultrasound irradiation: (a) Cl-CP, (b) uCl-CP, (c) N-CP and (d) uN-CP (scale bar = 20 nm).

Figure 6. N₂-Sorption isotherms of the dried powders before and after ultrasound irradiation **Figure 7**. Possible effect of ultrasound irradition (US) on (a) chlorapatite nanocrystals (green) in water, leading to dissolution, erosion and reprecipitation of hydroxyapatite (light blue), (b) hydroxyapatite nanocrystals in mother liquor, leading to dissolution and reprecipitation of hydroxyapatite (dark blue), (c) hydroxyapatite nanocrystals in water, leading to dissolution and erosion.

Figure 1





Figure 3



Figure 4



Figure 5







Figure 7

