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Consolidation of bone-like apatite bioceramics by spark plasma sintering of amorphous carbonated calcium phosphate at very low temperature

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ABSTRACT

Various carbonated calcium phosphate powders were synthesized by aqueous precipitation and ceramics consolidation by spark plasma sintering (SPS) at very low temperature was investigated. The objective was to preserve low crystallinity and avoid material decarbonation. SPS at low temperature only leads to little or no sintering when crystallized powders are used. Amorphous powders are required. In this case, consolidation occurs at temperatures below 150 °C. It is accompanied by crystallization of the amorphous phase into calcium-deficient carbonated apatite $Ca_{10-x,y}(PO_4)_{6-x}(HPO_4)_y(CO_3)_{2+x}(OH)_{2-y}$ (CO$_3$)$_2$. The resulting ceramics are microporous and highly cohesive with good mechanical properties (flexural strength = 18 MPa). The sintering mechanism, called "crystal fusion", is based on solid state diffusion of chemical species at the grain boundary and crystal growth within the amorphous particles. These bioceramics that mimic the composition of the bone mineral are expected to have a higher bioactivity than well crystallized carbonated hydroxyapatite ceramics obtained by conventional sintering.

1. Introduction

Synthetic hydroxyapatite (HA) of chemical formula $Ca_{10}(PO_4)_{6}(OH)_{2}$ is commonly used as bone substitute due to excellent biocompatibility and osteoconductive properties [1,2]. Calcium phosphate apatite can accept multiple ionic substitutions as well as vacancies in its crystal lattice [3,4]. Thus, the mineral part of bone is composed of a low crystalline and poly substituted nonstoichiometric apatite [5]. Carbonate (CO$_3^{2-}$) is the main substituted ionic group in biological apatites, with amounts ranging from 3 to 8 wt% [6,7]. A mean chemical formula of bone mineral apatite was proposed by Legros et al. as follows: $Ca_{10.8}(PO_4)_{6.3}(CO_3)_{3.7}(CO_3,HPO_4)_{3.3}$ [5]. As shown in this formula, carbonate ions can be incorporated in the apatite lattice in replacement of hydroxide ions (OH$^-$), named "A type substitution" (referred as $CA$ in the text), leading to the formula $Ca_{10}(PO_4)_{6}(OH)_{2}$ $2x(CO_3)_{x}$ with $0 \leq x \leq 1$. Synthetic A type carbonated HA ($CaHA$) can be produced by calcination of a stoichiometric HA powder under dry CO$_2$ flux at 900 °C for several hours according Trombe's method [8,10]. Carbonate can also substitute for phosphate ions (PO$_4^{3-}$), named "B type substitution" (referred as $CB$ in the text) leading to the formula $Ca_{10}(PO_4)_{6-x}(CO_3)_{x}(OH)_{2}$ with $0 \leq x \leq 2$. Synthetic B type carbonated HA ($CB$) can be obtained by aqueous precipitation [11,14]. Additionally, mixed AB type carbonated HA ($CA$), $Ca_{10}(PO_4)_{6-x}(CO_3)_{x}(OH)_{2}$ $2x(CO_3)_{x}$ can be produced by heating a $CaHA$ under controlled atmosphere containing CO$_2$ and water vapor partial pressures [14]. Due to these substitutions, such carbonated apatites are expected to exhibit better biological properties in terms of resorbability than pure HA [15-18]. We have shown recently that mixed $CaAB$ ceramics containing carbonates in similar amount to that found in the natural bone, exhibited enhanced biodegradability associated to an optimal balance for osteoblast and osteoclast activity (i.e. the main cells involved in bone regeneration) [19].

However, a serious drawback of carbonate apatites is their thermal instability. This material decomposes by decarbonation at "low" temperature (from about 450 °C under air atmosphere) which makes the consolidation of ceramic parts difficult by conventional natural sintering [20,23]. Consequently, these ceramics are sintered between 800 °C and 1000 °C under a flow of CO$_2$ gas (wet or dry according to the desired substitutions) to prevent the decarbonation reaction to occur, preserve carbonate groups located in B sites of the structure and avoid the formation of secondary phases such as lime or calcite [14,23]. Another approach to avoid the decomposition of the material could be sintering faster and/or at lower temperature using a non conventional sintering method such as Spark Plasma Sintering (SPS).

SPS has been investigated since the 1990s for the flash sintering of several ceramic materials. Its principle is based on a pulsed current...
passing through a conductive pressing die containing the sample, and consecutive heating by Joule effect, under compressive stress. This process is generally known to limit the grain growth and to lower the sintering temperature [24,25]. SPS of HA has been investigated by several research teams. The SPS temperature ranged from 900 °C to 1000 °C and led to high bulk density (> 99% of the maximum value) without the formation of any secondary phase. This increased the strength of the ceramic and limited grain growth [26–28].

Low crystallinity and nonstoichiometry, as in natural bone apatite, are other key factors to improve the biodegradability of apatitic synthetic materials. With this aim, synthetic nanocrystalline apatites were investigated. It was shown that these materials exhibit low crystallinity degrees were prepared by aqueous precipitation. Then, SPS of HA has been investigated to consolidate iodate substituted lead to thermal decomposition and eliminate the hydrated non apatitic surface layer of particles. In order to preserve such features previous studies explored SPS of non carbonated nanocrystalline apatites at low temperature (typically lower than 300 °C) [34–36]. Recently, this method has also been investigated to consolidate iodate substituted hydroxyapatite for nuclear application in the conditioning of radio active iodine 129 [37].

From this context, and in order to further increase the resorbability of apatitic compounds by incorporation of carbonate ions in the lattice, the objective of this work was to investigate the processing ability of low crystalline carbonated apatite ceramics. With this aim, powders having various chemical compositions (i.e. carbonate contents) and crystallinity degrees were prepared by aqueous precipitation. Then, SPS of these powders at very low temperature (150 °C) was performed. The structure, chemical composition and mechanical resistance of the processed ceramics were then investigated.

2. Experimental

2.1. Synthesis of carbonated phosphocalcic powders

Powders were prepared by precipitation in aqueous medium according to a protocol described previously [14]. Briefly, a solution of diammonium hydrogen phosphate ((NH4)2HPO4, 98.7%, Fisher Chemical) and ammonium hydrogen carbonate (NH4HCO3, > 99.0%, Sigma Aldrich) is added to a calcium nitrate solution (Ca(NO3)2·4H2O, 99%, Sigma Aldrich) at constant pH and temperature. The C/P ratio, designating the molar ratio in which the phosphate and carbonate ions were introduced, was set at 0.125. The amount of reagents was also fixed in order to have a Ca/P molar ratio equal to 1.667. The pH was maintained at a value of 9 by addition of an ammonium hydroxide solution (NH4OH 35%, Fisher Chemical). Three synthesis temperatures were investigated: 90 °C, which was close to the boiling point of the solution, 60 °C as an intermediate temperature and 37 °C the physiological temperature of the human body. The synthesis reactor was put under argon flow. When the last drop of phosphate and carbonate solution was added, the mixture was left to mature for 30 min. After this maturation, the precipitate was filtered on a Buchner funnel, then washed with 250 mL of distilled water per gram of powder and finally lyophilized.

2.2. Spark plasma sintering

Sintering of ceramic pellets was performed using a SPS Dr.Sinter 825 from Fuji Electronic Industrial Company (Japan). It involves application of a uniaxial compressive load on the sample and simultaneous heating by a pulsed continuous electrical current.

The sintering behavior of powders synthesized at different temperatures (37 °C, 60 °C and 90 °C) was studied. Weighed 300 mg of powder was introduced into a 10 mm diameter graphite die previously covered with Papyex®, a thin graphite foil, to facilitate the demolding of the sample after SPS. Fig. 1a gives the SPS temperature and pressure cycles used in this study. The sintering temperature was set at 150 °C with a heating rate of 20 °C min⁻¹. The pressure was set to 80 MPa, applied in 1 min. After application of the compressive stress a dwell was made at 50 °C during a few minutes to homogenize the temperature in the enclosure of the SPS device. Each test was carried out under argon atmosphere.

Influence of the applied pressure (60 MPa, 80 MPa, 100 MPa and 120 MPa) was also investigated. For this study, the SPS cycle was modified (Fig. 1b): whatever the final pressure, the load was applied within 1 min. The final temperature was set at 175 °C for a short dwell (1 min) with a 40 °C min⁻¹ heating rate and without dwell at 50 °C. Pressureless SPS was also performed. The powder was initially compacted under a 80 MPa compressive pressure applied for 1 min before heating. Then, the mechanical pressure was removed and the thermal cycle described in Fig. 1a was applied.
The in situ densification curves were registered from the measurement of the moving piston displacement and the temperature measured by a thermocouple placed close to the sample inside a hole drilled in the graphite die. The relative density (or densification ratio) of SPS samples corresponded to the ratio between the apparent density, determined by Archimedes’ method in water, and the real solid density, obtained by helium pycnometry measurements (Micromeritics, AccuPyc II 1340).

Complementary hot pressing (HP) experiment was performed in a Goliath LPA (2000 °C, 5 tons), ECM Technologies (Applied Physics). The same protocol of sample preparation as that described for SPS was used. The temperature was measured on the external wall of the graphite die containing the sample. The dwell temperature was set at 400 °C for 10 min with a heating ramp of 15 °C min⁻¹. A 80 MPa compressive stress was applied at the beginning of the heating cycle. At the end of dwell at 400 °C, the temperature and the load were decreased gradually. The displacement of the piston applying the mechanical load was measured during the test.

2.3. Samples characterization

X Ray diffraction patterns of precipitated powders and SPS ceramics were acquired on a BRUKER D8 Advance diffractometer using the CuKα radiation. The patterns were recorded in a 20 range 20° - 60° with a step size of 0.02° and a dwell time of 0.51 s. Phase identification was performed by comparing the experimental diagram with the database of diffraction patterns (PDF) of the International Center for Diffraction Data (ICDD). Lattice parameters were refined by profile fitting using the JANA2006 software (Czech Republic). Diffraction peaks were fitted according to a pseudo Voigt function, allowing the system to be anisotropic. The crystallite size \( L_{\text{diff}} \) was determined from two peaks (200) and (310) giving respectively the mean crystallite length and width) by the Scherer Eq. (1) [38] and using the Peakoc software (France):

\[
L_{\text{diff}} = \frac{\lambda}{\cos\beta} \sqrt{\frac{\beta_{\text{exp}}^2 - \beta_0^2}{\beta_{\text{exp}}^2}}
\]  

with λ, the radiation wavelength; \( \beta \), the full width at half maximum of the peak and \( \theta \), the diffraction angle of the associated (hkl) plane. The lattice distortion parameter \( g_{\text{diff}} \) in the crystallographic direction (00l) was also calculated based on the model of Hoseman and Vogel (Eq. (2)) [39]:

\[
\sqrt{\beta_{\text{exp}}^2 - \beta_0^2} = \frac{1}{L_{\text{diff}}} + \pi \frac{g_{\text{diff}}^2}{d_{\text{diff}}} m^2
\]  

with \( m \) the reflexion order and \( d_{\text{diff}} \) the reticular distance of (hkl) planes family, applied here to the family of planes (00l).

Temperature programmed X Ray diffraction was performed using the same diffractometer, in an ANTON PAAR, NTK 1200N oven. Powder was placed in an alumina crucible lined with platinum foil. The maximum temperature was set at 1000 °C with a heating rate of 10 °C min⁻¹ under nitrogen flow. The patterns were acquired in the 20 range 27° - 38° with a step size of 0.02° and a dwell time of 1.0 s, every 20 °C during a temperature plateau of 11 min.

Infrared spectra were collected on a NICOLET 5700 spectrometer. The spectra were recorded in transmission mode, using KBr pellets, over the 4000 400 cm⁻¹ range with a resolution of 2 cm⁻¹ from an accrulation of 64 scans.

For quantitative chemical analysis, calcium and phosphorus contents were determined by Inductively Coupled Plasma/Optical Emission Spectroscopy (ICP/OES) on a PERKIN ELMER, Optima 8300, Optical Emission Spectrometer. The analysis was performed in solution from about 30 mg of powder dissolved in perchloric acid prepared at 6 mol L⁻¹ (70%, AnalR NORMAPUR) and diluted in 100 mL of water. Carbonate content of powders was determined by elemental analysis (HORIBA, Carbon analyzer, EMIA 321V) consisting of burning 500 mg of powder sample placed in a crucible with a metallic blend (iron, copper, tin) in order to initiate the combustion. Carbonate content ofapatite powders and sintered ceramics was also evaluated using FTIR spectroscopy according to Grunenwald et al. method [40]. FTIR was also used to estimate the proportions of carbonates within the various chemical sites (i.e. A or B type or labile, the latter corresponding to carbonates in non-crystallized environment).

The morphology of powders and the microstructure of SPS ceramics were observed by scanning electron microscopy working in secondary electron mode (JEOL, FEG SEM, JSM 7400F) and transmission electron microscopy (JEOL HR TEM 2100). TEM also allowed to record selected area electron diffraction (SAED) patterns. For powders, some grains were dispersed in acetone and placed on a steel sample holder prior to SEM examination and on a mesh Cu grid with a carbon membrane for TEM. The pellets observed by SEM were fixed on the sample holder with adhesive carbon tape.

Biaxial flexural strength was measured on 10 cylindrical samples obtained after SPS at 150 °C in a 30 mm diameter die and the strength value was calculated according to Meganck et al. method [41].

Specific surface area (\( S_{\text{BET}} \)) and bulk density (\( d \)) of powders were measured by the BET method (MICROMERITICS, ASAP 2020, 8 points) and by He pycnometer (MICROMERITICS, AccuPyc II 1340) respectively. Differential Thermal Analysis (DTA) was conducted on 50 mg of powder using in a NETZSCH STA 449F3 equipment from 30 to 1200 °C with a heating rate of 10 °C min⁻¹ under inert argon flow. Thermomechanical analysis (i.e. dilatometry) was realized on 3 mm thick pellet using the same thermal cycle as for DTA in a NETZSCH DIL402C device. The force applied by the probe was 25 nN. The sample was shaped by compaction of 300 mg of powder in a 10 mm diameter steel matrix under a pressure of 62.5 MPa.

3. Results and discussion

3.1. Powder synthesis and characterization

The XRD patterns of synthesized powders are presented in Fig. 2. Diffraction peaks corresponding to an apatitic phase (PDF 9 432) appear for the powders produced at 60 °C and 90 °C but with the lowering of the synthesis temperature, they become broader and poorly defined. The crystallite size (Table 1) is smaller for the powder synthesized at 60 °C than at 90 °C. In both cases the crystallites exhibit a preferred growth along the c axis of the apatite structure, the value of \( L_{\text{c(0002)}} \) being about half the value of \( L_{\text{c(0032)}} \). This anisotropic grain growth is typical of apatite crystals. Crystallinity is also related to lattice deformations, which can be evaluated using the lattice distortion parameter \( g_{\text{diff}} \). Its value is smaller for the powder prepared at 90 °C than at 60 °C (Table 1). Consequently, the powder synthesized at 90 °C shows less lattice distortions than the powder synthesized at 60 °C. This means

![Fig. 2. XRD patterns of synthesized carbonated calcium phosphate powders.](image-url)
that, in agreement with literature data [11,21,22], the increase of the synthesis temperature increases the crystallinity of the powder (larger crystallite dimensions and lower distortion parameters). In comparison with the lattice parameters of stoichiometric HA (a = 9.418 Å and c = 6.884 Å according ICCD PDF file 90 009 0432) both a and c are greater for the two carbonated powders (Table 1). According to Legeros et al., the parameter a decreases when substituting carbonates in B site, these ions being smaller than phosphate ones. Landi et al. showed that the parameter c increases with this substitution [42,43]. In contrast, in the case of A type substitution, Elliot demonstrated that the value of a increases and c decreases [44]. In addition, contraction of the c axis and expansion in the perpendicular direction are generally observed for non stoichiometric apatites [45]. Consequently, additional substituted species (carbonates Ca and/or Ca, or HP04 2−) may have caused the enlargement of the crystallographic lattice in both directions a and c, which will be confirmed hereafter by the spectroscopic and chemical analyzes of these powders.

The XRD pattern of the powder synthesized at 37 °C, does not show any defined peak (Fig. 2), suggesting that the grains are amorphous. Only two broad and diffuse peaks with maxima at about 20 = 30° and 50° are registered which cannot be ascribed to an apatite feature. Moreover, this diffractiongram is very similar to that of amorphous calcium phosphate (ACP) found in the literature data [46,47].

This also agrees with SEM and TEM observations (Fig. 3). The powder synthesized at 37 °C is made of agglomerated spherical grains which size varies from about 35 to 250 nm. Thus, in this powder grain growth during the synthesis process was isotropic, which was not the case of the crystallized powders. The amorphous character of this sample is also confirmed by the diffraction pattern produced by SAED. Here also, this powder presents the same morphological characteristics as ACP grains composed of so called Posner’s clusters of composition close to (Ca0.6(P0.34Ca0.65)PO4)x0.6H2O0.4 described in the literature [48,49]. Observations of the powder synthesized at 60 °C (Fig. 3) highlights nanometric needle like grains having an average size of about 40 nm length and 20 nm width. Due to its low crystallinity and very small grain size, this powder exhibits properties close to the nanocrystalline apatites described by Eichert et al. [31]. The powder synthesized at 90 °C is also made of rod shaped grains (Fig. 3), but of much larger size: about 150 nm length and 60 nm width. SAED patterns confirm that the grains precipitated at 60 °C and 90 °C are crystallized according to the apatite structure.

The bulk density d and the specific surface area S BET of the powders are given in Table 2. The values of density measured for the crystallized powders (i.e., 2.96 and 2.94 g cm−3 after synthesis at 60 °C and 90 °C respectively) are not significantly different and close to that obtained by Lafon et al. (2.83 g cm−3 for a CaHAP precipitated at 90 °C in similar conditions) [14]. The density obtained for the amorphous powder synthesized at 37 °C is significantly lower than those of the crystallized powders. Its value (2.52 g cm−3) approaches the bulk density calculated for ACP by Holt et al. at 2.31 g cm−3 [50].

The specific surface area of the raw powders is linked to the grain size observed by electronic microscopy. The amorphous calcium phosphate consisting of large spherical grains has the smallest specific surface area. The carbonated apatite powder synthesized at 90 °C, with the largest needle like grains has a greater surface area. It can be noted that the value obtained here is close to that of stoichiometric hydroxyapatite (68 m2 g−1) [51] but slightly higher than that found by Lafon et al. for carbonated apatite (36 m2 g−1) [14] synthesized in similar conditions. The powder synthesized at 60 °C has the largest surface area in accordance with its nanometric grain size.

FTIR spectra (Fig. 4a) were collected on the raw synthesized powders. For powders synthesized at 60 °C and 90 °C, typical vibration bands of phosphate groups in the apatite structure are observed at 472 cm−1 (ν2) 560, 575 and 600 cm−1 (ν4), 960 cm−1 (ν1), 1020 1120 cm−1 (ν3). The bands at 630 and 3560 cm−1 are assigned to apatitic OH groups [52,53].

Regarding the powder synthesized at 37 °C, the spectrum exhibits broad bands which are characteristic of a non crystalline material. These bands are similar to those found for ACP prepared by

![Fig. 3. Microscopic observations of powders synthesized at different temperatures: a) SEM, b) TEM and SAED patterns.](image)
precipitation in an alkaline medium [53,54].

The amount of water, which can be estimated by the broad band between 3700 and 2800 cm\(^{-1}\) and the band at 1650 cm\(^{-1}\), decreases with increasing the synthesis temperature. This water is mainly due to the presence of a hydrated layer at the surface of apatite nanocrystals as it was previously described for nanocrystalline apatites [31,45]. Water seems present in greater amount in the amorphous powder produced at 37 °C, which can be explained by the fact that constitution water can be found in the interstices between the amorphous clusters [48,50].

Carbonate ions can be observed in two regions of the spectra: the \(\nu_2\) vibrations of carbonate groups between 850 and 890 cm\(^{-1}\) and the \(\nu_3\) vibrations between 1400 and 1570 cm\(^{-1}\). The \(\nu_4\) CO\(_3\) band (in the range 760-665 cm\(^{-1}\)) [53] is too weak to be visible on the spectra and cannot be exploited. For each vibration mode, contributions can be attributed to the various environments of carbonate groups: A site, B site and labile carbonates. Table 3 gathers the characteristic wavenumbers of these contributions based on literature reports.

Zooms of \(\nu_2\) and \(\nu_3\) CO\(_3\) bands (Fig. 4c and d) focus on the main contributions of carbonate ions. For the powder synthesized at 90 °C, typical \(\nu_2\) bands of carbonates in B sites around 1412 cm\(^{-1}\) and 1462 cm\(^{-1}\) are observed. A shoulder is also observed around 1480 1500 cm\(^{-1}\) which could be attributed to labile carbonates. However, it could also be attributed to carbonates in A sites in mixed AB type apatites [55]. This hypothesis can be confirmed on the one hand by the presence of a very small peak at 1542 cm\(^{-1}\) attributed to A site carbonates and on the other hand by the examination of the asymmetrical \(\nu_2\) CO\(_3\) band centered at 875 cm\(^{-1}\), implying a major contribution of the B sites at 872 cm\(^{-1}\) with a minor A sites contribution at 878 cm\(^{-1}\). Additionally, a light contribution of labile carbonates can be responsible for the enlargement of the band down to 860 cm\(^{-1}\).

Table 3

<table>
<thead>
<tr>
<th>Band</th>
<th>CO(_3) in A-site</th>
<th>CO(_3) in B-site</th>
<th>Labile CO(_3)</th>
</tr>
</thead>
<tbody>
<tr>
<td>References</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(\nu_2) CO(_3)</td>
<td>878 cm(^{-1})</td>
<td>872 cm(^{-1})</td>
<td>866 cm(^{-1})</td>
</tr>
<tr>
<td>(\nu_3) CO(_3)</td>
<td>1465 cm(^{-1})</td>
<td>1412 cm(^{-1})</td>
<td>1417 cm(^{-1})</td>
</tr>
<tr>
<td>(\nu_4) CO(_3)</td>
<td>1542 cm(^{-1})</td>
<td>1462 cm(^{-1})</td>
<td>1480 cm(^{-1})</td>
</tr>
</tbody>
</table>

The carbonate content (Table 4) was determined by calculating the ratio \(r_{CO_3}/P\) between the integrated intensity of \(\nu_2\) CO\(_3\) (1550 1350 cm\(^{-1}\) domain) and that of \(\nu_4\) PO\(_4\) (1230 912 cm\(^{-1}\) domain) using a calibration curve according to Grunenwald et al. method [40].
carbonate content was found to be around 3 wt% with a significant decrease when the synthesis temperature is high (90 °C) in accordance with previous studies [21,22,59]. The results obtained from this spectroscopic method were in excellent agreement with carbon elemental analyses. Consequently carbonate quantification on SPS samples will be evaluated from FTIR data using the Grunenwald et al. method. Indeed, the measurement of carbonate from carbon elemental analysis could be biased by the presence of residual carbon at the sample surface resulting from Papex® used during SPS.

The Ca/P molar ratio determined from ICP/EOS titrations of calcium and phosphorus was found to be lower than 1.667 (i.e. 10/6 the value of stoichiometric HA) for the three powders. It increased with the synthesis temperature (Table 4). These results are usually encountered during the synthesis of calcium deficient hydroxyapatite because of partial substitution of HPO₄²⁻ for phosphates in the apatite lattice [51]. In addition, in the present study, the samples were carbonated. A type carbonate has no effect on Ca/P ratio, but for B type carbonates the stoichiometry is defined by Ca/(P + CB) = 10/6. Thus, in the presence of B type carbonation, the Ca/P ratio of powders must be greater than 10/6. Since the Ca/P values were all smaller than 1.667, the synthesized powders are clearly nonstoichiometric and must include other ionic species such as HPO₄²⁻ as hypothesized above.

Finally, the powder precipitated at 37 °C is an amorphous carbonate calcium phosphate whose stoichiometry is close to amorphous tri calcium phosphate composed of Posner’s clusters. Its approximate formula would be Ca₉₋ₓ(PO₄)₆₋₂ₓ(HPO₄)₂ₓ(CO₃)²ₓ(OH)₂ₓ, neglecting the light carbonation in A site and the presence of a hydrated surface layer.

### 3.2. Spark plasma sintering

#### 3.2.1. Influence of powder properties

The powders were heat treated by spark plasma sintering. Four distinct domains were highlighted (Fig. 5). In the first one, the registered increase of relative density, associated with sample shrinkage, is due to the application of the mechanical load and the associated reorganization and compaction of the powder bed. The second domain corresponds to the temperature dwell at 50 °C (see Fig. 1) performed in order to homogenize the temperature in the enclosure of the SPS device. There is almost no change registered in this domain. In the third domain of heating from 50 °C up to 150 °C, there is no significant change for the powders synthesized at 90 °C and 60 °C. However, regarding the powder synthesized at 37 °C, a significant shrinkage of about 20% is registered at about 130 °C. Then, in the last domain, which corresponds to the dwell at 150 °C, the three compositions behave similarly with a light shrinkage of about 3.5%.

The apparent bulk density (Archimede method), solid density (pycnometry) and the final densification ratio of SPS samples are given in Table 5. After SPS, the ceramics sintered from powders initially synthesized at 60 °C and 90 °C exhibit a quite similar densification with a final relative density of 49% and 52% of the maximum value, respectively. This behavior is similar to that observed by Grossin et al. during the SPS of uncarbonated biomimetic nanocrystalline apatites matured 1 day [34].

XRD patterns of SPS samples are given in Fig. 6. The lattice parameters, average crystallite size and lattice distortion δ₀₀₂ were refined from the XRD patterns. Results are summarized in Table 6. The XRD pattern of the ceramic produced from the powder synthesized at 90 °C remains unchanged (Figs. 6 and 2). Microscopic (Fig. 7) and visual observations of the sample confirm that it did not consolidate during SPS. Grains are still rod shaped with an average size of crystallites of about 48 nm long and 28 nm width (Table 6) without change compared to the initial powder (Table 1). The crystallites just seem to have coalesced to form grains of about 100 nm long without significant cohesion between them and the final compact remains very friable. The ceramic obtained from the powder synthesized at 60 °C also shows a

### Table 4
Chemical composition of synthesized powders.

<table>
<thead>
<tr>
<th>Powder synthesis temperature</th>
<th>CO₃ wt.% (FTIR)</th>
<th>CO₃ wt.% (elemental analysis)</th>
<th>Molar ratio Ca/P (ICP/OES)</th>
</tr>
</thead>
<tbody>
<tr>
<td>37 °C</td>
<td>3.1 wt.%</td>
<td>3.1 wt.%</td>
<td>1.45 ± 0.04</td>
</tr>
<tr>
<td>60 °C</td>
<td>3.6 wt.%</td>
<td>3.5 wt.%</td>
<td>1.55 ± 0.08</td>
</tr>
<tr>
<td>90 °C</td>
<td>1.9 wt.%</td>
<td>1.9 wt.%</td>
<td>1.62 ± 0.10</td>
</tr>
</tbody>
</table>

### Table 5
Bulk density of ceramic pellets obtained after SPS.

<table>
<thead>
<tr>
<th>Powder synthesis temperature</th>
<th>Apparent density (g cm⁻³)</th>
<th>Solid density (g cm⁻³)</th>
<th>Densification ratio (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>37 °C</td>
<td>2.11 ± 0.02</td>
<td>2.973 ± 0.005</td>
<td>71.0</td>
</tr>
<tr>
<td>60 °C</td>
<td>1.49 ± 0.02</td>
<td>2.926 ± 0.005</td>
<td>49.2</td>
</tr>
<tr>
<td>90 °C</td>
<td>1.51 ± 0.02</td>
<td>2.911 ± 0.005</td>
<td>52.0</td>
</tr>
</tbody>
</table>

Fig. 5. Relative density changes of samples during SPS.

Fig. 6. XRD patterns of ceramics after SPS at 150 °C. Powder synthesized at: a) 37 °C, b) 60 °C and c) 90 °C.
tered from powders synthesized at 60 °C and 90 °C. According to the a during SPS was accompanied by crystallization of the amorphous powder leads to a crystallized apatitic phase. Thus, consolidation sintered ceramic reveals that the low temperature SPS of the amor
particles of the initial powder (Fig. 2), a ceramic made of small needle change of grain morphology is also noticed: from the large spherical

The powder synthesized at 37 °C shows a much higher densification, the value reaching about 71% of the maximum density after SPS (Table 5). SEM observation of the ceramic (Fig. 7) points out a densified and cohesive material with some microporosity (27.8% measured by the Archimedes’ method). Biaxial flexural strength of SPS samples synthesized at 37 °C (using 5 replicates) is 18.3 ± 5 MPa. This result points out that consolidation and sintering are effective during SPS. A change of grain morphology is also noticed: from the large spherical particles of the initial powder (Fig. 2), a ceramic made of small needle like grains (Fig. 7) with crystallite size of about 33 nm length and 16 nm width (Table 6) is obtained. The observations also show that the acicular grains have grown preferentially in a plane perpendicular to the direction of the applied load during SPS. XRD pattern (Fig. 6) of the sintered ceramic reveals that the low temperature SPS of the amorphous powder leads to a crystallized apatitic phase. Thus, consolidation during SPS was accompanied by crystallization of the amorphous powder. The c lattice parameter is greater than that of ceramics so tere from powders synthesized at 60 °C and 90 °C. According to the literature, this expansion could be caused by insertion of CO$_3^{2-}$ in A sites or by the presence of HPO$_4^{2-}$ in B sites of the apatite.

FTIR spectra are given in Fig. 8. For all the ceramics the typical bands of an apatitic material are found with the vibration modes of phosphate groups (ν$_1$, ν$_2$, ν$_3$ and ν$_4$) and OH groups (ν$_1$ and ν$_2$) appearing at the wavenumbers detailed in subsection 3.1. Moreover, the shoulder on the ν$_4$ PO$_4$ vibration band at 534 550 cm$^{-1}$ attests the presence of hydrogenophosphate in the sample synthesized at 37 °C (arrow on Fig. 8b). The presence of ν$_2$ CO$_3$ and ν$_3$ CO$_3$ bands attests that carbonates are not eliminated during SPS.

Fig. 8c and d details ν$_2$ CO$_3$ and ν$_3$ CO$_3$ vibration bands and Table 6 reports the overall carbonate content estimated from FTIR spectra using the method of Grunenwald et al. [40]. The FTIR spectra of pellets obtained after SPS of powders synthesized at 90 °C or 60 °C do not show any significant change from those of the initial powders (Figs. 8 and 4).

Table 6
Crystalline characteristics of SPS ceramics obtained by XRD peak fitting and carbonate content evaluated by FTIR quantification according to Grunenwald et al. [40].

<table>
<thead>
<tr>
<th>Powder synthesis temperature</th>
<th>a lattice parameter (Å)</th>
<th>c lattice parameter (Å)</th>
<th>L$_{220}$ (Å)</th>
<th>L$_{002}$ (Å)</th>
<th>g$_{bc}$ (Å rad)</th>
<th>CO$_3$ wt.% (FTIR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>37 °C</td>
<td>9.45(9)</td>
<td>6.88(7)</td>
<td>161</td>
<td>326</td>
<td>0.0065</td>
<td>1.5 wt.%</td>
</tr>
<tr>
<td>60 °C</td>
<td>9.42(6)</td>
<td>6.89(1)</td>
<td>74</td>
<td>165</td>
<td>0.0073</td>
<td>3.2 wt.%</td>
</tr>
<tr>
<td>90 °C</td>
<td>9.42(5)</td>
<td>6.90(1)</td>
<td>275</td>
<td>477</td>
<td>0.0057</td>
<td>1.5 wt.%</td>
</tr>
</tbody>
</table>

However, a slight decrease in carbonate content is noted (Tables 6 and 4), indicating that part of the carbonate ions have decomposed and evolved as CO$_2$ gas during SPS, via the possible following reactions:

$$2\text{HPO}_4^{2-} + \text{CO}_3^{2-} \rightarrow 2\text{PO}_4^{3-} + \text{H}_2\text{O} + \text{CO}_2$$  \hspace{1cm} (3)

or

$$\text{H}_2\text{O} + \text{CO}_3^{2-} \rightarrow 2\text{OH}^- + \text{CO}_2$$  \hspace{1cm} (4)

For the amorphous powder a change of carbonate location and overall content after SPS is observed. The estimation of carbonate content by FTIR reveals a loss of half the initial carbonates during the SPS treatment. The profile of vibration bands shows the presence of carbonates in both A and B sites of the apatite as well as labile carbonates.

Because the powder synthesized at 60 °C and 90 °C did not allow the production of cohesive ceramics by SPS at low temperature (150 °C), the following subsection 3.2.2. focuses on the behavior of the amorphous carbonated calcium phosphate powder synthesized at 37 °C.

3.2.2. Thermal behavior of amorphous calcium phosphate synthesized at 37 °C

The shrinkage (fast increase of relative density) observed for carbonated ACP at about 130 °C during SPS is likely due to a concomitant consolidation and crystallization. In order to determine more precisely the behavior of this powder, other sintering conditions, including pressureless SPS, hot pressing and natural sintering were investigated. The chemical changes occurring during SPS were also analyzed.

3.2.2.1. Other sintering methods. At first, the powder was heat treated to investigate its natural sintering ability without mechanical load and pulsed electric current. DTA performed under argon flow shows an exothermic reaction starting at 630 °C with a maximum at 640 °C (Fig. 9).

In situ temperature programmed XRD was performed in a furnace under N$_2$ flow. Fig. 10 gives the typical diffractograms obtained from this experiment. The diffractograms registered up to 580 °C show an amorphous material whereas the pattern obtained at 620 °C exhibits a crystallized compound. Thus, the exothermic phenomenon registered by DTA is assignable to a crystallization event. The crystallized phase corresponds to beta tricalcium phosphate β TCP (ICCD 55 898) with traces of apatitic phase. This result is in accordance with the literature indicating that ACP expresses an exothermic crystallization at about 600 700 °C into β TCP [61]. But, it differs from the apatite phase ob tained after SPS at 150 °C. The stabilization of apatitic phase at low temperature is linked to the structural configuration of ACP clusters. Indeed, as reported in previous works, some calcium orthophosphates including hydroxyapatite present locally the ionic arrangement of Posner’s cluster [62]. Thisprobably helps to stabilize the apatitic phase at very low temperature. Moreover, this phase transformation into apatite is similar to that observed when ACP evolves in aqueous medium leading to calcium deficient apatite [63 65].

Dilatometry (TMA) under argon flow was also performed on this material. The dilatometric curve (Fig. 11) indicates that the
Fig. 8. FTIR spectra of SPS pellets: a) general spectra from 4000 cm$^{-1}$ to 400 cm$^{-1}$, b) zoom on the $\nu_4$ PO$_4$ band, c) zoom on the $\nu_3$ CO$_3$ band, d) zoom on the $\nu_2$ CO$_3$ band. Contributions are denoted A for A-site, B for B-site and L for labile carbonate. (Ts = Temperature of powder synthesis).

Fig. 9. DTA under argon flow of amorphous carbonated calcium phosphate powder synthesized at 37 °C.

Fig. 10. Temperature programmed XRD patterns performed under N$_2$ flow at: a) 580 °C and b) 620 °C. c) XRD pattern at ambient temperature of sample after SPS at 150 °C. (The diffraction peaks are shifted toward low angles for diagram (b) due to the thermal expansion of the powder and sample holder).

Fig. 11. TMA under argon flow of amorphous carbonated calcium phosphate powder synthesized at 37 °C.
crystallization at high temperature (in the temperature range around 600 °C) is accompanied by a very light compaction (< 0.2%) in comparison with the shrinkage obtained by SPS (densification > 10% at 130 °C). Thus, crystallization that induces an increase of bulk density is not the main phenomenon responsible for the consolidation and shrinkage of carbonated apatite by SPS.

Finally, it appears that the apatitic material produced by SPS is impossible to produce by a conventional sintering method.

Complementary pressureless SPS and HP of the powder synthesized at 37 °C were carried out in order to evidence the individual contributions of an applied pressure or electric current on the sample densification. The densification curves are given in Fig. 12 in comparison with the curve of the sample densified by SPS at 150 °C under 80 MPa. During HP a shrinkage phenomenon (fast increase of relative density) similar to that observed during SPS appears but at a much higher temperature around 325 °C. Moreover, the final relative density of the HP ceramic is only 63% whereas it was 71% after SPS. Pressureless SPS only leads to a very light densification with a final relative density of 48%. The shrinkage curve is similar to that registered during SPS (under 80 MPa) of the powder synthesized at 60 °C (Fig. 5). These results highlight that the sole application of an electric current or of a supplementary mechanical energy provided by the load allows the activation of the consolidation. It must be emphasized that the difference observed between the densification curve performed at 80 MPa in this assay and the curve presented in previous subsection for the same pressure (3.2.1 Fig. 5) may be explained by the changes of the sintering cycles (see Fig. 1).

3.2.2.2. Chemical analysis of SPS ceramics. Composition changes of the material during SPS (150 °C; 80 MPa) were investigated by FTIR spectroscopy. The carbonate content in the pellet was estimated according to Grunenwald’s method [40]. The overall amount of \( \text{CO}_3^{2-} \) ions is 1.5 wt.%, i.e. twice less than in the initial powder (3.1 wt.%). A fraction of carbones should have evolved as carbon dioxide gas during the treatment. The hypotheses mentioned above can be proposed to explain this loss of \( \text{CO}_3^{2-} \) (Eqs. (3) and (4)). The presence of HPO_4^{2-} ions, observed in the FTIR spectrum (arrow on Fig. 8b) and involved as reactants in Eq. (3), can be explained by an internal hydrolysis of PO_4^{3-} ions during heating according to Eq. (5):

\[
P O_4^{3-} + H_2O \rightarrow HPO_4^{2-} + OH^- \tag{5}
\]

In order to investigate the composition of the material, a thorough study was conducted on \( \nu_2 \text{PO}_4 \) and \( \nu_2 \text{CO}_3 \) vibration bands. A spectral decomposition (curve fitting) was performed to assess the area and relative intensity of the bands constituting the spectra. The decomposition of \( \nu_2 \text{PO}_4 \) (Fig. 14) used the wavelength position of the various contributions in accordance with previous studies on monoclinic apatites [45,65]. This decomposition reveals the presence of apatitic hydroxide ions. Because ACPs do not theoretically contain apatitic OH\(^-\), such hydroxide ions should appear during the consolidation/crystallization step according to the hydrolysis of phosphates and/or carbonates (Eqs. (4) and (5), respectively). The spectral decomposition also allows highlighting the various contributions of phosphates including two kinds of HPO_4^{2-} ions: an apatitic contribution of HPO_4^{2-} located in B sites and a non apatitic contribution. A fraction of these ions, mostly in non apatitic environments, could be formed during the internal hydrolysis of phosphates (Eq. (3)). This hydrolysis reaction had already been assumed to occur in the hydrated layer during the low temperature sintering of monoclinic apatites by Drouet et al. [35]. These observations tend to confirm our first hypothesis: decarbonation observed during SPS would be mainly caused by an internal hydrolysis of the PO_4^{3-} groups (Eq. (5)) followed by the decomposition of the carbonates by reaction with the non apatitic HPO_4^{2-} (Eq. (3)) rather than a direct hydrolysis of carbonates (Eq. (4)).

In order to identify and quantify the proportions of the different kinds of carbonates in the SPS ceramic, the \( \nu_2 \text{CO}_3 \) spectrum in the region 840 900 cm\(^{-1}\) was also decomposed. HPO_4^{2-} and CO_3^{2-} ions...
exhibit vibrations at similar wavenumbers around 870 cm\(^{-1}\) which makes it difficult to isolate the respective contributions of these ionic groups. To overcome this difficulty, a preliminary study was carried out using uncarbonated nanocrystalline apatite powders containing various amounts of \(\text{HPO}_4^{2-}\) ions. This allowed to relate the area of \(\text{HPO}_4^{2-}\) bands in the domain of \(\nu_4 \text{PO}_4\) band (Fig. 15b) to the area of the associated band at 870 cm\(^{-1}\) (Fig. 15a). Powders were synthesized by aqueous precipitation following a rapid addition of a solution of calcium nitrate to a solution of ammonium hydrogen phosphate. The precipitates were matured for various times, resulting in varying amounts of hydrogenphosphate ions in the powder. Then, the decom position of the FTIR bands according to the method proposed by Van decandelaere et al. [45] was performed. The calibration curve (Fig. 15c) obtained after calculation of the areas of hydrogen phosphate vibration bands gives the coefficient that relates the sum of \(\text{HPO}_4^{2-}\) bands areas (A2) in the \(\nu_4 \text{PO}_4\) region to the area (A1) of the associated band at 870 cm\(^{-1}\). This calibration curve was used to calculate the area (A1) of the \(\text{HPO}_4^{2-}\) band at 870 cm\(^{-1}\) in the domain of \(\nu_2 \text{CO}_3\) bands (Fig. 15) from the area (A2) of the \(\text{HPO}_4^{2-}\) bands in the domain of \(\nu_4 \text{PO}_4\) band of the SPS sample (Fig. 14).

Carbonate contributions were positioned at the wavenumbers given in Table 3 to decompose the \(\nu_2 \text{CO}_3\) bands. The carbonate species contained in the amorphous powder before SPS are hypothesized in a labile position. After SPS (Fig. 16), 28.6% of the remaining carbonates are labile and three quarters of the total amount have migrated in the A and B sites of the formed apatite with 27.8% and 43.6% respectively.

This decomposition highlights also the presence of non apatitic ions in the sintered ceramic, as the spectrum decomposition reveals the presence of some labile carbonates but also of non apatitic \(\text{HPO}_4^{2-}\) ions. Thus, it can be stated that the material obtained after SPS is similar to biomimetic nanocrystalline apatites [31], composed of non stoichiometric crystalline carbonated apatitic grains (Ca\(_{10-x}\)(PO\(_4\))\(_x\)(HPO\(_4\))\(_y\)(CO\(_3\))\(_2\)(OH)\(_{2-x-y}\)(CO\(_3\))\(_y\)) surrounded by a non apatitic hydrated layer which constitutes the grain boundaries as denoted in the

\[ A1 = 0.10 \times A2 + 0.01 \]
schematic representation of the resulting ceramic (Fig. 17).

3.2.3. Discussion

It was pointed out by Grossin et al. in the case of non carbonated nanocrystalline apatite sintered at low temperature by SPS that the classical mechanisms of diffusion and grain growth occurring during conventional sintering at high temperature cannot happen below 300 °C [34]. Classical solid state diffusion in crystallized compounds cannot be activated at such low temperatures. The authors stated the importance of the presence of a hydrated layer surrounding the apatitic core of nanoparticles which was characterized by a high ionic mobility. This statement is confirmed in our study since the powders synthesized at 60 °C and 90 °C which are crystallized and have a very small or an absence of hydrated surface layer could not consolidate by low temperature SPS at 150 °C. Amorphous calcium phosphate powders can be assimilated to fully non apatitic hydrated phosphocalcic grains (the clusters being surrounded by 15–20% of residual water in the inter stices of the Posner structure). This implies high solid state ion mobility at very low temperature favoring chemical reactions and leading to the formation of grains (apatitic core) that crystallize into an apatitic structure and grow preferentially along their c axis, while the sur rounding hydrated layers of adjacent grains fuse to form the grain boundaries. This results in high strength porous ceramics. Such a sin tereing, recalling the mechanisms involved in cold sintering of ferro electric materials [66,67], processes through a phenomenon previously called “crystal fusion” enhanced by the additional activation energy provided by an applied load [34,35]. In our case, calcium deficient carbonated apatite of chemical composition Ca_{10-x}(PO_4)_x(\mathrm{CO}_3)_y(OH)_2 \times 2y \cdot (\mathrm{CO}_3)_y crystallizes from the amorphous clus ters that formed the initial spherical grains leading to nanometric acicular grains after SPS. Starting from an amorphous powder was found to be necessary in carbonated systems to enhance the densification and provide a chemical composition very close to that of the mineral part of bone. This contrasts with the conventional sintering of calcium defi cient apatites that leads to mixtures of well crystallized stoichiometric calcium phosphates (Ca_3P_2O_7 + TCP, TCP or HA + TCP) from about 700 °C [68].

Although the mechanism involved in this process remains to be analyzed in more detail, it can be stated that the methods described previously in the literature, starting from crystallized powders, are un likely to achieve in the production of highly cohesive bioceramics of chemical composition and structure very close to the mineral phase of bone. Such “bone like” ceramics are expected to provide superior biological properties, the evaluation of which is under consideration.

4. Conclusion

Carbonated calcium phosphate powders of various composition and crystallinity level were synthesized by aqueous precipitation at dif ferent temperatures. These powders were heat treated by Spark Plasma Sintering at very low temperature (150 °C). Only the amorphous car bonated calcium phosphate powder, precipitated at the physiologic temperature of 37 °C, could be sintered at such a low temperature. The consolidation is accompanied with crystallization of the amorphous phase into carbonated apatite. The high flexural strength of the re sulting microporous ceramic confirms the efficiency of Spark Plasma Sintering. This ceramic is made of a low crystalline nonstoichiometric carbonated apatite. The grains are composed of an apatitic core sur rounded by a hydrated layer containing non apatitic ions which con stitutes the grain boundaries. The high mobility of chemical species of the initially amorphous powder appears responsible for the consolida tion of the sample at low temperature by surface ionic diffusion process enhanced by the applied load and electrical current during SPS. This leads to the formation of non apatitic grain boundaries while the core of grains crystallizes. It can be emphasized that the SPS of amorphous calcium phosphates makes it possible to consolidate a calcium deficient carbonated apatitic phase of generic composition Ca_{10-x}(PO_4)_x(\mathrm{CO}_3)_y(OH)_2 \times 2y \cdot (\mathrm{CO}_3)_y which is impossible to preserve by conventional sintering.

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