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Accurate characterization of pure silicon-substituted hydroxyapatite powders
synthesized by a new precipitation route

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Abstract

This paper presents a new aqueous precipitation method to prepare silicon-substituted hydroxyapatites $\text{Ca}_{10}(\text{PO}_4)_{6-y}(\text{SiO}_4)_y(\text{OH})_{2-y}(\text{V}_{\text{OH}})_y$ (SiHAs) and details the characterization of powders with varying Si content up to $y=1.25$ mol mol$^{-1}$SiHA. X-ray diffraction (XRD), transmission electron microscopy (TEM), solid-state nuclear magnetic resonance (NMR) and Fourier transform infrared (FTIR) spectroscopy were used to accurately characterize samples calcined at 400°C for 2 h and 1000°C for 15 h. This method allows for synthesizing monophasic SiHAs with controlled stoichiometry. The theoretical maximum limit of incorporation of Si into the hexagonal apatitic structure is $y<1.5$. This limit depends on the OH content in the channel, which is a function of the Si content, temperature and atmosphere of calcination. These results, particularly those from infrared spectroscopy, express serious reservations about the phase purity of SiHA powders, pellets or scaffolds prepared and biologically evaluated in the literature.

Keywords: biomaterials; silicon-substituted hydroxyapatite; precipitation method, infrared spectroscopy, NMR spectroscopy.
1. Introduction

According to the literature, silicon-substituted hydroxyapatite (SiHA) is a highly promising material in the field of bioactive bone substitutes and bone tissue engineering. It is now well-established that silicon plays an important role in the early stage of cartilage and bone growth [1-4]. Soluble silicon species have been shown to stimulate spontaneous calcium phosphate precipitation (i.e. the mineral bone phase) [5] and to increase bone mineral density [6]. Moreover, silicon has been reported to have a positive influence on the synthesis of type I collagen by human osteoblast cells (MG-63 cell line) *in vitro* [7]. Thereby, it is expected that silicon could enhance the hydroxyapatite (HA) bioactivity [8, 9], and silicon-substituted hydroxyapatites (SiHAs) have become a subject of great interest in bone repair. The SiHA structure corresponds to the substitution of phosphate ions (PO$_4^{3-}$) by silicate ions (SiO$_4^{4-}$) into the HA crystal structure. Different mechanisms for charge compensation have been suggested [8, 10, 11]. The most cited one was proposed by Gibson et al. with the creation of anionic vacancies at OH$^-$ sites [8, 12]. This mechanism leads to silicon-substituted hydroxyapatites with the general formula Ca$_{10}$(PO$_4$)$_6$y(SiO$_4$)$_y$(OH)$_2$y(V$_{OH}$)$_y$, where $y$ represents the molar number of silicate groups introduced into the apatitic structure ($0 \leq y \leq 2$) and V$_{OH}$ stands for vacancies maintaining the charge balance. The incorporation of Si into the HA structure in solid solution, *i.e.* without the formation of other phases, seems to be limited. However, the value and the origin of this limitation are still not known, with for instance the following values: 5 wt% ($\approx 1.7$ mol$_{Si}$/mol$_{SiHA}^{-1}$) [13-15], 4 wt% ($\approx 1.4$ mol$_{Si}$/mol$_{SiHA}^{-1}$) [16, 17], 3.1 wt% ($\approx 1.1$ mol$_{Si}$/mol$_{SiHA}^{-1}$) [18], 2 wt% ($\approx 0.7$ mol$_{Si}$/mol$_{SiHA}^{-1}$) [11, 19], 1.0 wt% (0.36 mol$_{Si}$/mol$_{SiHA}^{-1}$) [20] or 0.28 wt% (0.1 mol$_{Si}$/mol$_{SiHA}^{-1}$) [21]. Additionally, it has been suggested that the concentration of 0.8 wt% of Si ($\approx 0.28$ mol$_{Si}$/mol$_{SiHA}^{-1}$) is optimal to induce the development of important bioactivity [22-24]. A value of 2.2 wt% of Si was also reported by Thian et al. [25]. Unfortunately, in spite of extensive studies in recent
years, these results remain heterogeneous, confusing and sometimes misleading. For instance, Hing et al. revealed faster bone apposition and improved adhesion and proliferation of osteoblast-like cells for SiHA compared to stoichiometric HA [23, 26], whereas Palard et al. found no significant difference in the behavior of MG-63 osteoblast-like cells between pure HA and SiHA pellets (three compositions: y=0.2, 0.4 and 0.6 mol\textsubscript{Si} mol\textsubscript{SiHA}\textsuperscript{-1}) [27]. Recent critical analyses of the published results regarding SiHAs have highlighted the lack of experimental evidence which could explain the real effects of Si substitution on biological activity in a biological environment [28, 29]. In particular, it has been criticized that the physico-chemical characterizations of SiHA bioceramics are not detailed (purity, solubility, rate of incorporation of Si inside the crystal lattice, etc.). Therefore, the available data do not provide sufficient information to establish the origin of the improved biological performance of SiHA: (i) a direct effect of SiHA by Si release, (ii) an indirect effect of SiHA by changes in the physico-chemical properties of HA due to Si substitution (microstructure, superficial chemistry, topography, etc.) or (iii) an effect of second phases (crystalline and/or amorphous). According to Boanini et al., the term “ion-substituted” is quite often used without any experimental proof regarding the incorporation of ions inside the crystal lattice of calcium orthophosphates [28]. The unclear bioactivity of SiHA ceramics could be explained by variations in the phase composition. The first evidence for this was provided by the few accurate analyses available in the literature which show that SiHA powders can contain crystallized [16, 30-35] and amorphous [16, 34, 36-38] impurities. The study by Kanaya et al. is representative of the characterization problems of SiHA samples [38]. Indeed, while the X-ray diffraction patterns show only the characteristic lines of the HA phase (PDF: 09-432), the $^{29}$Si MAS NMR spectrum revealed that only 10% of Si was incorporated into the HA lattice; the rest was on the particle surface in the form of polymeric silicate species [38]. An equivalent observation was made by Gasquères et al. [16]. Most studies do not evidently show
the purity of their SiHA samples (powders, pellets or scaffolds) [8, 9, 19, 21-24, 26, 27, 39-58]. They generally provide an imprecise physico-chemical characterization with assumptions based on X-ray diffraction patterns or incomplete infrared band assignment [8, 9, 19, 22, 32, 36, 39, 59-61]. Moreover, infrared vibrations at 692, 840, 890 and 945 cm$^{-1}$, detected on an SiHA sample containing 1.6 wt% Si (1200°C for 2 h), were attributed without evidence by Gibson et al. to the substitution of SiO$_4$ for PO$_4$ into the HA lattice [8, 12]. Unfortunately, this article set the standard for SiHA analysis by infrared spectroscopy and has been widely cited to prove the purity of SiHA powders synthesized using Gibson’s method [9, 23, 39, 40, 43, 44, 46, 47, 53, 57] or displaying the same new infrared bands [14, 32, 36, 48, 50, 54, 62].

Several methods are used to prepare Si-substituted hydroxyapatites (SiHAs), such as the sol-gel route [48], resuspension processes [20, 33, 34, 63-66], solid state reactions [10, 67], hydrothermal techniques [17, 68, 69], mechanochemical methods [70], magnetron sputtering [14], pulsed laser deposition [51, 71], electrophoretic deposition [72] and precipitation from aqueous solutions. Aqueous precipitation methods are the most often described and set up in the literature. Two different procedures are used to produce Si-HA from aqueous solutions: (i) the acid-base neutralization [8, 9, 16, 19, 23, 30, 35, 37, 39, 46, 47, 49, 50, 53, 56, 57, 73-75], and (ii) the use of phosphorus and calcium salts [13, 21, 32, 36, 37, 60, 74, 76, 77]. Generally, far less attention has been devoted to the silicon reagent. Two organic compounds are mainly used as a source of silicate ions: tetraethylorthosilicate (TEOS, Si(OC$_2$H$_5$)$_4$) and tetraacetoxyxilane (TAS, Si(COOCH$_3$)$_4$) which are not miscible in water. Powders synthesized through these methods are poorly crystallized and often contaminated by second phases (e.g. $\alpha$ and $\beta$-Tricalcium phosphate, amorphous phase, silicocarnotite [16, 19, 30, 31, 36-38, 49, 59, 60, 63, 64, 67, 74, 76]) or undesired ions like Mg [56], S [77] or Na [78, 79].
In order to correctly describe the physical, chemical and biological properties of SiHAs and to compare them to routinely implanted HA and β-TCP, well-characterized pure SiHAs powders first need to be prepared. Therefore, this work was devoted to the development of a new route to synthesize monophasic SiHA powders with controlled stoichiometry. To this purpose, a solution of soluble silicate was first prepared from TEOS via a sol-gel route, and then accurate powder analysis was carried out by means of ICP/AES, X-ray powder diffraction, Rietveld refinement, high resolution electron transmission microscopy (HR-TEM) with energy dispersive spectroscopy (EDS) as well as infrared (FT-IR/ATR) and solid-state NMR spectroscopy. Two pH levels of precipitation were studied, as well as six Si/P molar ratios.

2. Materials and methods

2.1 Powder synthesis

HA and SiHA powders were prepared through an aqueous precipitation method using a fully automated apparatus. A diammonium hydrogen phosphate aqueous solution ((NH$_4$)$_2$HPO$_4$, 99%, Merck, Germany), and, if applicable, an alkaline silicate solution were added to a calcium nitrate solution (Ca(NO$_3$)$_2$, 4H$_2$O, 99%, Merck, Germany) using peristaltic pumps. The reaction was performed under an argon flow (4.8, AirLiquide) to prevent any excessive carbonation of precipitates. The pH of the suspension was adjusted by the addition of a 28% ammonia solution (Merck, Germany) by means of a pH stat (Hanna Instruments), and the temperature was controlled and regulated automatically with an external T-probe. The suspension was continuously stirred and refluxed. After complete introduction of the solutions, the suspension was matured for 24 h, and then filtered under vacuum. Finally, the precipitates were dried at 70°C overnight.

The solution of soluble silicates was prepared from tetraethylorthosilicate (TEOS ≥99%, Aldrich, Germany) via a sol-gel route. The original step consisted of the preparation of
a silica gel through the hydrolysis and condensation of this precursor. First, the alkoxide
groups (Si-OEt) of TEOS were hydrolyzed to silanol groups (Si-OH). The reaction was
catalyzed by a nitric acid aqueous solution (7.5 $10^{-3}$M). This generates a considerable amount
of monomers (Si-OH), which aggregate and form a colloidal suspension called the “sol”. This
reaction, based on Klein et al. [80], was carried out by maintaining the molar ratio of
TEOS/ethanol/acidified water at 1.0/8.5/4.0 and the solution at 25°C in an ambient
atmosphere. Second, the condensation of the silanol or alkoxide groups was initiated by
increasing the solution pH to a basic level, about pH 8, with a few drops of pure 28%
ammonia solution (Merck, Germany). This reaction creates siloxane bridges (Si-O-Si),
leading to the formation of a silica gel. The gelation reaction was continued for about 12 h at
25°C. Finally, a solution of soluble silicates was obtained by depolymerization of the silica
gel. This was achieved through the nucleophilic attack of siloxane linkages (Si-O) by
hydroxide ions in pure 28% ammonia solution (pH > 11). After filtration through a Millipore
filtration unit (Magma nylon, 0.8 µm), the soluble silicate ion concentration of these solutions
was measured by means of an ion-exchange chromatography system (DIONEX DX-500)
equipped with a SiO$_4^{4-}$ ion exchange column (AS4A) with a UV/Vis detector (410 nm). The
soluble silicate ion concentration in these solutions was, on average, 1076 ± 34 ppm (n=14
solutions). This value is a function of the preparation process (e.g. hydrolysis, condensation
and depolymerization reactions). Thereby, the volume of the solutions of soluble silicates was
adjusted for each synthesis.

A pure HA powder Ca$_{10}$(PO$_4$_6)(OH)$_2$ was first prepared to be used as a reference
material for further SiHA syntheses and analyses. This was obtained at a pH and temperature
of precipitation of 9.5 and 50°C, respectively, with a maturation time of 24 h, and a reagent
(Ca/P) molar ratio equal to 10/6.
The reagent ratios used to prepare the SiHA powders were calculated by assuming, first, the general formula \( \text{Ca}_{10}(\text{PO}_4)_{6-y}(\text{SiO}_4)_y(\text{OH})_{2+y}(\text{V}^-)_{y} \) and, second, a constant molar ratio \( \text{Ca}/(\text{P}+\text{Si}) \) equal to 10/6. The volume of the calcium and phosphate aqueous solutions was fixed at 500 mL and 250 mL, respectively. Table 1 summarizes the amount (mol) of Ca, P and Si used for each (Si)HA powder prepared with \( y \) equal to 0, 0.25, 0.50, 0.75, 1.00, 1.25; designated hereafter as HA, Si_{0.25}HA, Si_{0.50}HA, Si_{0.75}HA, Si_{1.00}HA and Si_{1.25}HA, respectively. Precipitations were achieved at 50°C for 24 h of maturation time, according to the preliminary tests performed on pure HA. Two parameters of precipitation were examined, the pH of precipitation and the stoichiometric number \( y \), \textit{i.e.}, the amount of silicate theoretically incorporated into the apatite structure. Precipitations were conducted either at pH 9.5, previously established for HA synthesis, or at pH 10.80 ± 0.05. The former was chosen according to phosphate and silicate speciation curves, in order to have HPO$_4^{2-}$ [81] and H$_2$SiO$_4$' ions [82] in solution as the main phosphate and silicate species, respectively. The studied parameters are listed in Table 1.

The as-synthesized powders were heated under air using an alumina crucible. Aliquots of each sample were calcined both at 400°C for 2 h (designated hereafter as “raw powders”), in order to remove synthesis residues, and at 1000°C for 15 h under air according to ISO norm 13779-3 in order to characterize powders with a well-defined structure. The heating and cooling rate was fixed at 4°C min$^{-1}$.

2.2 Powder characterization

2.2.1 X-ray powder diffraction and Rietveld refinement

Crystalline phases were identified by means of a Siemens D5000 0/2θ X-ray diffractometer (XRD) using CuKα radiation and operating at 40 kV and 20 mA. XRD patterns were first collected over the 20 range of 10-60° at a step size of 0.03° and counting time of 4 s per step in order to determine the phase composition. Crystalline phases detected in the patterns were
identified by comparison to standard patterns from the ICDD-PDF (International Center for Diffraction Data-Powder Diffraction Files). In addition, high resolution XRD scans were obtained on calcined powders from 10° to 120° in 0.02° steps with a counting time of 10.5 s per step. These patterns were used to calculate the lattice parameters and the crystalline size by full pattern matching. In this mode, the positions of the diffraction peaks and the systematic extinctions are calculated from the cell parameters and the space group. The intensities of the lines are adjusted by the refinement program and not calculated from the atomic positions. The refinements were performed using the space group of the HA structure P63/m (PDF 09-432) by means of the Topas software (Bruker, Germany). The initial cell parameters were taken as \( a = 9.42 \text{Å} \) and \( c = 6.88 \text{Å} \).

The evolution of the crystallinity of the samples after calcination at 1000°C for 15 h was evaluated by means of the full width at half maximum (FWHM) of the (211) peak at \( 2\theta = 31.8° \), as it had the highest intensity and minimal overlap with neighboring peaks.

### 2.2.2 Infrared spectroscopy

Fourier transform infrared (FT-IR) measurements were carried out with the use of a MIR TF VERTEX 70 Spectrometer by means of the ATR system. The spectra were recorded over the range of 450-4000 cm\(^{-1}\) with a resolution of 2 cm\(^{-1}\). Spectra were obtained by signal averaging 32 successive scans. Every measurement was at least duplicated on two independent powder samples crushed by hand, with an agate pestle and mortar, and deposited on the ATR system. Spectra were normalized with respect to the \( \nu_4 \) band of the phosphate group at about 602 cm\(^{-1}\), according to a classical procedure [83, 84].

### 2.2.3 NMR spectroscopy

NMR spectra were recorded on a Bruker AVANCE 300 spectrometer: \( B_0 = 7.05 \text{T} \), \( \nu_0(^1\text{H}) = 300.13 \text{MHz}, \nu_0(^{31}\text{P}) = 121.49 \text{MHz}, \nu_0(^{29}\text{Si}) = 59.62 \text{MHz} \), using a CP-MAS Bruker probe with 4 mm rotors spinning at 14 kHz for \(^1\text{H} \) and \(^{31}\text{P} \) and with 7 mm rotors spinning at 5
kHz for $^{29}\text{Si}$. Chemical shifts are referenced to TMS for $^1\text{H}$ and $^{29}\text{Si}$ and 85% aqueous $\text{H}_3\text{PO}_4$ for $^{31}\text{P}$. Single pulse and CP (cross-polarization) MAS experiments have been used to study these materials. CP MAS relies on the heteronuclear dipolar interaction between an abundant spin X ($^1\text{H}$ here) and a low abundant nuclei Y ($^{31}\text{P}$ and $^{29}\text{Si}$ here). It allows the study of local molecular motion. The NMR parameters are summarized in Table 2.

2.2.4 Electron microscopy (HR-TEM, SAED and EDX)

High resolution transmission electron microscopy (HR-TEM) was conducted on calcined powders (1000°C/15 h) using a JEOL 2010 F Microscope at a voltage of 200 kV. Samples were prepared by dispersing the powders in ethanol. After sonication at 40 W for 5 min, a small drop of the suspension was placed onto copper mesh grids coated with a holey carbon film. Finally, a thin coating of gold (10 nm in thickness) was sputtered on half of the grid. Gold was then distributed as crystallized nano-domains which were used as a reference in the selected area electron diffraction (SAED) patterns to calculate as precisely as possible the lattice parameters. The SAED patterns obtained from regions with or without gold on the HA part were the same.

2.2.5 Elemental analysis

The silicon, phosphorous and calcium content in powders was determined by inductively coupled plasma atomic emission spectrometry (ICP/AES) (HORIBA Spectrometer, Jobin-Yvon, Activa model). Powder samples were dissolved in a nitric acid solution (0.5 M). Solutions were prepared (e.g. powder mass, dilution) in order to limit the measurement uncertainty as well as to determine the concentrations of Ca, P and Si within the highest sensitivity range of the ICP/AES device (around 30 ppm). Lastly, the carbon content in the powders was determined using an elemental analyzer with an infrared detector (LECO CS-444 carbon and sulfur analyzer).
The stoichiometric number \( y \) was determined by means of Eq. 4, assuming that powders containing \( 0 \leq \text{wt}\% \text{Si} \leq 2.81 \) (or \( 0 \leq y \leq 1.00 \)) are pure monophasic SiHAs.

\[
\text{Ca}_{10}(\text{PO}_4)_{6-y}(\text{SiO}_4)_y(\text{OH})_{2.8}(\text{V}_\text{OH})_y, \quad \text{without any second crystalline or amorphous phase.}
\]

\[
\%\text{wt Si} = (m_{\text{Si}} 100 / m_{\text{SiyHA}}) = (C_{\text{m Si}} V_{\text{sol}} 100) / (m_{\text{SiyHA}}) \quad (1)
\]

\[
n_{\text{Si}} = (\%\text{wtSi} m_{\text{SiyHA}}) / (M_{\text{Si}} 100) \quad (2)
\]

\[
M_{\text{SiyHA}} = M_{\text{HA}} - (19.895 y_{\text{Si}}) \quad (3)
\]

\[
y_{\text{Si}} = (n_{\text{Si}} M_{\text{SiyHA}}) / (m_{\text{SiyHA}}) = (\%\text{wtSi} M_{\text{HA}}) / (100 M_{\text{Si}} + 19.985 \%\text{wt Si}) \quad (4)
\]

where \( \%\text{wt Si} \) is the mass percentage of Si in the powder, \( C_{\text{m Si}} \) is the mass concentration of Si in the analyzed solution by ICP/AES (ppm or mg L\(^{-1}\)), \( V_{\text{sol}} \) is the volume of the analyzed solution (L), \( m_{\text{SiyHA}} \) is the mass of the powder dissolved in the former solution (mg), and \( M_{\text{SiyHA}}, M_{\text{HA}} \) and \( M_{\text{Si}} \) are the molar masses of SiyHA, HA and Si, respectively (g mol\(^{-1}\)).

### 3. Results and discussion

#### 3.1 Influence of pH on the SiHA phase composition

Si\(_{0.50}\)HA powders were synthesized at 50°C with pH values equal to 9.50 (Si\(_{0.50}\)HA-9.5) and 10.80 (Si\(_{0.50}\)HA-10.8). Fig. 1 shows the XRD diffractograms of powders calcined at 1000°C for 15 h. The pattern of the Si\(_{0.50}\)HA-9.5 sample presents two different crystalline phases matching the ICDD standard for hydroxyapatite (HA, PDF 9-432) and \( \alpha \)-tricalcium phosphate (\( \alpha \)-TCP, PDF 9-348). The Si\(_{0.50}\)HA-10.8 diffractogram displays only the characteristic diffraction lines of HA. This result indicates that the final composition of the powders is a function of silicate ion speciation. \( \text{H}_2\text{SiO}_4^- \) or more basic forms of silicate ions have to be maintained during the synthesis to obtain a thermally stable SiHA phase, i.e. a monophasic powder after heat treatment at 1000°C/15 h. Therefore, herein, the following results are presented for powders prepared at pH 10.8. Under these synthesis conditions, the general reaction of precipitation can be written as follows:

\[
10\text{Ca}^{2+}+(6-y)\text{HPO}_4^{2-}+y\text{H}_2\text{SiO}_4^-+(8+y)\text{OH}^- \leftrightarrow \text{Ca}_{10}(\text{PO}_4)_{6-y}(\text{SiO}_4)_y(\text{OH})_{2.8}(\text{V}_\text{OH})_y+(6+2y)\text{H}_2\text{O} \quad (5)
\]
To our knowledge, this is the first time that a true precipitation reaction has been given for SiHA.

3.2 Influence of the Si content ($y_{Si}$)

3.2.1 X-ray diffraction analysis

Fig. 2 shows the diffraction patterns of HA and Si$_y$HA ($y_{Si} = 0.25, 0.50, 0.75, 1.00$ and $1.25$ mol) raw powders. Each pattern matches well with the standard pattern of HA (PDF 9-432). No other phase was detected. The samples exhibit broad diffraction lines which indicate very small crystallites and/or low crystallinity. More, the FWHM increased with silicon content. The diffractograms of heat treated powders (1000°C/15 h) are displayed in Fig. 3. They present no secondary phase besides hydroxyapatite for a silicon content up to $y_{Si} = 1.00$. On the other hand, HA (PDF 9-432) and α-TCP (PDF 9-348) were detected in the pattern of Si$_{1.25}$HA calcined at 1000°C for 15 h. Moreover, as observed for the raw powders, the crystallinity of Si$_y$HA calcined powders with $1 \geq y \geq 0$ decreased with an increase in the silicate content. Indeed, the higher the Si content, the higher the FWHM of the diffraction line. For instance, the FWHM of the diffraction line at 31.8° (2θ) increased as follow: 0.070 (HA) < 0.096 (Si$_{0.25}$HA) < 0.160 (Si$_{0.50}$HA) < 0.164 (Si$_{0.75}$HA) < 0.170 (Si$_{1.00}$HA). This is due to both direct (decrease in the crystallinity) and indirect (decrease in the crystallite size) effects of Si, but the individual contributions of these effects cannot be evaluated by X-ray diffraction.

The XRD patterns indicate that whatever the amount of silicon ranging between $0 \leq$ wt%Si $\leq 3.51$ (or $0 \leq y \leq 1.25$), the crystalline phase of the precipitates is a hydroxyapatite (PDF 09-432). Thereby, silicon can be incorporated in the apatitic structure, or in an amorphous phase or in both phases. However, only precipitates containing up to 2.81 wt%Si (or $y = 1.00$) are thermally stable at 1000°C for 15 h.

3.2.2 Lattice parameter refinement
Refinement was performed on monophased heat-treated powders \( (0 \leq y \leq 1.00) \). The results, displayed in Table 3, reveal changes in the apatitic structure parameters. The Si substitution causes \( c \)-axis and unit cell volume expansion as well as a reduction in the mean crystallite size. The same observation was reported in other studies \([10, 17, 32, 70, 85]\). On the other hand, and in contrast to the linear increase of the lattice parameter \( c \) with an increasing amount of Si, the \( a \) lattice parameter fluctuates at random between 9.416 Å and 9.428 Å. These results highlight that all or part of the silicon is incorporated into the apatitic structure.

### 3.2.3 Elemental analysis

The experimental Ca/(P+Si) molar ratios of the final calcined powders are presented in Table 1. Values are in accordance with the expected ones (1.67) for all compositions, considering experimental error. Moreover, the calcined powders were not carbonated since the carbon content by weight was equal to 0.03, 0.07, 0.05, 0.05 and 0.05 wt% (± 0.02 wt%) for HA, Si\(_{0.25}\)HA, Si\(_{0.50}\)HA, Si\(_{0.75}\)HA and Si\(_{1.00}\)HA, respectively.

The results, presented in Table 1, show that the chemical formulas of SiHA calculated first by the measured silicon contents, then by the theoretical formula \( \text{Ca}_{10}(\text{PO}_4)_6 \cdot y(\text{SiO}_4)_y(\text{OH})_{2-y}(\text{V}_{\text{OH}})_y \) and finally using Eq. 4 are in a good agreement with the nominal ones.

### 3.2.4 Electron microscopy

A low magnification bright-field image of heat-treated Si\(_{1.00}\)HA particles is shown in Fig. 4a. SiHA powders are composed of round particles, smaller than 100 nm in diameter for Si\(_{1.00}\)HA, whereas the HA powder shows bigger acicular crystals \([86]\). At higher magnification, grains do not present any dislocation or disorder within the central region. Similarly, high-resolution lattice images of the grain surface do not show any amorphous or disordered layer (see Fig. 4b). A selected area electron diffraction (SAED) pattern obtained on the same grain is displayed in Fig. 5a. SAED patterns show a diffraction pattern with well-defined spots regardless of the selected area. This indicates the high crystallinity of the heat-
treated SiHA powders and the homogeneity of the crystals. Second, a very good agreement appears between the SAED patterns and the PDF card 09-432 of HA as identified by X-ray diffraction. Moreover, no halo ring was observed, confirming the absence of the amorphous phase. Fig. 5b shows a SAED pattern for the [010] zone axis. Two other SAED patterns were obtained for the [1-10] zone axis. Indexation of these patterns allows for calculating the \( \frac{c}{a} \) ratio by means of the interval between two spots. Thus, for instance, according to the hexagonal crystallographic structure identified by X-ray diffraction and the distance \( D_1 \) and \( D_2 \) between two spots along the [h00] and [001] directions ([010] zone axis, Fig. 5b), respectively, the \( \frac{c}{a} \) ratio was calculated as follows:

\[
(c/a)^2 = \frac{3}{4} \left( \frac{D_1}{D_2} \right)^2
\]

The results are the average of about ten intervals per pattern. A \( c/a \) ratio of 0.730 ± 0.005 was determined. Moreover, other experimental patterns for different zone axes (not shown here) were obtained and compared to theoretical electron diffraction patterns calculated by means of the Java Electron Microscopy Simulation (JEMS) software [87]. The results indicate that the experimental and simulated patterns are perfectly superimposed for \( 0.734 \geq c/a \geq 0.729 \). Additionally, the \( c/a \) ratio value from PDF 09-432 (0.7309) is included in this range.

3.2.5. Solid-state NMR analysis

3.2.5.1. \(^{31}\text{P} \) MAS NMR

The \(^{31}\text{P} \) MAS NMR spectra of HA, \( \text{Si}_{0.50}\text{HA} \) and \( \text{Si}_{1.00}\text{HA} \) calcined at 1000°C for 15 h are shown in Fig. 6a. The spectra present a main narrow peak which corresponds to the single P site of hydroxyapatite [88], and whose position shifts to higher frequencies with increasing silicon content: 2.8 ppm for HA, 3.0 ppm for \( \text{Si}_{0.50}\text{HA} \), and 3.1 ppm for \( \text{Si}_{1.00}\text{HA} \). Likewise, the FWHM broadens: 166 Hz < 188 Hz < 233 Hz for raw HA, \( \text{Si}_{0.50}\text{HA} \) and \( \text{Si}_{1.00}\text{HA} \), respectively (figure not included), and 55 Hz < 132 Hz < 233 Hz for calcined HA, \( \text{Si}_{0.50}\text{HA} \)
and \( \text{Si}_{1.00} \text{HA} \), respectively (Fig. 6a). The chemical shift and peak broadening indicate that the SiHA structure is different from a perfect hydroxyapatite short-range structure. The increasing line width suggests an increase in local disorder around those phosphate groups where Si is incorporated in the HA structure. Besides the 2.8 ppm peak, two additional weak broad peaks at 4.6 and 5.8 ppm were detected in the \(^{31}\text{P}\) spectra. The peak at 4.6 ppm was only revealed in the \(^{1}\text{H} \rightarrow ^{31}\text{P}\) CP MAS spectrum (Fig. 7b). HA powder did not present the 4.6 ppm signal (Fig. 7a). According to Hartmann et al. [89], the \(^{31}\text{P}\) MAS peak at 5.8 ppm corresponds to \( \text{PO}_4 \) tetrahedra in a proton-free region of a hydroxyapatite channel structure containing OH vacancies (\( i.e. \) along the crystallographic \( c \)-axis). These proton-free regions, observed in oxyhydroxyapatite \( \text{Ca}_{10}(\text{PO}_4)_6(\text{OH})_{2-2x}(\text{O})(V\text{OH})_x \) (OHA), must exist in silicon-substituted hydroxyapatite \( \text{Ca}_{10}(\text{PO}_4)_{6-y}(\text{SiO}_4)_y(\text{OH})_{2-y}(V\text{OH})_y \) (SiHA), or silicon-substituted oxyhydroxyapatite \( \text{Ca}_{10}(\text{PO}_4)_{6-y}(\text{SiO}_4)_y(\text{OH})_{2-y-2x}(\text{O})(V\text{OH})_{x+y} \) (SiOHA). The assignment of the peak at 4.6 ppm, that seems related to the incorporation of Si in the apatite lattice, will be discussed later on (section 3.2.5.3).

### 3.2.5.2. \(^{1}\text{H}\) MAS NMR

The \(^{1}\text{H}\) MAS NMR spectra of pure HA, \( \text{Si}_{0.50} \text{HA} \) and \( \text{Si}_{1.00} \text{HA} \) powders calcined at 1000°C for 15 h are shown in Fig. 6b. Two signals were observed, one sharp at 0 ppm and one broader at about 5.2 ppm. The former (0 ppm) is characteristic of OH groups in crystalline HA [89]. Moreover, and similarly to the \(^{31}\text{P}\) signal at 2.8 ppm, the greater the Si content in the apatitic structure of the calcined powders, the broader the FWHM of the signal at 0 ppm (FWHM = 108 Hz for HA, 191 Hz for \( \text{Si}_{0.50} \text{HA} \) and 228 Hz for \( \text{Si}_{1.00} \text{HA} \)). This suggests a change in the environment around the hydroxyl groups, highlighting the modification of O-H bond distances due to Si. As postulated by Gomes et al., this concomitant broadening of the \(^{31}\text{P}\) and \(^{1}\text{H}\) resonances with Si content confirms the incorporation of Si atoms into the apatite lattice [30]. The signal at 5.2 ppm was attributed by Hartmann et al. to OH\(^{+}\) positions missing...
only one neighboring ion in OHA channels [89]. This type of OH\(^{-}\) group along the channels of
the apatite structure leads to the formation of “proton pairs” stabilized by a hydrogen bond.
The intensity of this proton position is higher in SiHA samples than in HA, but does not seem
to be an exclusive function of the Si content (see Fig. 6b). Therefore, \(^1\)H MAS NMR indicates
the presence of two types of OH\(^{-}\) groups along the channels of the apatite structure: an
unchanged proton line position with continuous OH\(^{-}\) groups (0 ppm) and OH\(^{-}\) pairs
surrounded by vacancies (5.2 ppm).

3.2.5.3. \(^1\)H-\(^{31}\)P HETCOR CP MAS

To confirm that the entire proton position is located in the apatite channel structure, two-
dimensional \(^1\)H-\(^{31}\)P HETCOR CP MAS experiments were performed on raw and calcined
powders. Fig. 8 presents \(^1\)H-\(^{31}\)P HETCOR CP MAS spectra recorded on raw Si\(_{1.00}\)HA (Fig.
8a) and calcined Si\(_{0.50}\)HA (Fig. 8b) powders. The spectra show two main 2D correlation peaks
on both raw and calcined powders. The same peaks were observed for Si\(_{0.25}\)HA, Si\(_{0.50}\)HA and
Si\(_{1.00}\)HA (HETCOR CP MAS results for calcined Si\(_{0.25}\)HA and Si\(_{1.00}\)HA not included). The
stronger one (peak A, Fig. 8b), characteristic of an ideal HA structure, is due to the dipolar
interaction between the P sites (\(\delta^{(31}\)P) = 2.8 ppm) and the OH sites (\(\delta^{(1}\)H) = 0 ppm). It
corresponds to an undisturbed hydroxyapatite short-range structure [89]. The second one
(peak B, Fig. 8b) appears between the \(^{31}\)P signal at 4.6 ppm and the \(^1\)H signal at 5.2 ppm. Its
intensity increases with heat treatment (Fig. 8) and the Si content (data not included). This
correlation is not observed for the HA sample (data not included). Moreover, a low extent of
peak B is observed with the main \(^{31}\)P peak at 2.8 ppm (peak C, Fig. 8b). The intensity of this
2D correlation peak becomes stronger with Si content as well. More generally, the higher the
amount of Si incorporated into the SiHA structure, the stronger the dispersion of the main
correlation between A and B along the \(^{31}\)P axis. Finally, as expected, the HETCOR spectra
show no correlation peak of the \(^{31}\)P line at 5.8 ppm, confirming that it corresponds to a
proton-free phosphate position. To conclude, first the broadening of peak A indicates an increase in the distribution of P-H bond distances when silicate is incorporated into the HA structure, corresponding to the loss of local ordering in the phosphate environment. Second, peaks B and C correspond to two distinct P sites in the vicinity of isolated pairs of protons (OH− pairs) present in the channel structure of silicon-substituted oxyhydroxyapatite 

\[ \text{Ca}_{10}(\text{PO}_4)_{6-y}(\text{SiO}_4)_y(\text{OH})_{2.5-2x}\text{O}_x(\text{V}_{\text{OH}})_{y+x} (\text{SiOHA}) \]: (i) peak B represents a very strongly distorted hydroxyapatite short-range structure and (ii) peak C represents a less distorted one. Peak B is clearly related to SiO$_4$ substitution and proves once again that charge compensation implies OH vacancies along the channels, leading to the formation of OH− pairs in the vicinity of the substituted PO$_4$ tetrahedra. Peak C can be due to SiO$_4$ substitution as well as dehydration of SiHA, both leading to the formation of OH vacancies in SiOHA.

3.2.5.4. $^{29}$Si MAS NMR

The $^{29}$Si MAS NMR spectra of Si$_{1.00}$HA powders, raw and calcined, are shown in Fig. 9. The raw powders present two resonances: one sharp at about -72 ppm that is clearly attributed to SiO$_4^{4−}$ ($Q^0$) in the apatitic structure [90], and a broad peak at about -100/-110 ppm which corresponds to $Q^3/Q^4$ silicon species [16]. The results confirm that the main part of the silicate is incorporated into the HA lattice during precipitation, according to the main reaction Eq. 1. The rest of the silicates are adsorbed at the crystallite surface, as observed for phosphates during the precipitation of apatitic calcium phosphates [91, 92]. After calcination, the broad peak at -100/-110 ppm disappears without the detection of new signals, similar to those reported in the literature (Si-TCP or α-Ca$_3$Si$_3$O$_9$) [16, 30, 38]. The same results were obtained for the Si$_{0.25}$HA and Si$_{0.50}$HA samples (data not included).

3.2.6. FT-IR analysis

3.2.6.1 As-synthesized samples
The IR spectra of the raw HA and SiHA powders with a silicon content up to $y_{Si} = 1.25$ mol (figure not included) present strong similarities and exhibit intense bands characteristic of hydroxyapatite. They correspond to the four vibrational modes of phosphate groups ($v_1$, $v_2$, $v_3$, and $v_4$), and the stretching ($v_S$) and librational ($v_L$) modes of the hydroxide groups [81, 92-94]. The intensity of the latter, at 630 cm$^{-1}$ ($v_L$) and 3570 cm$^{-1}$ ($v_S$), clearly decreased with the amount of silicon. These results confirm that the as-synthesized precipitates exhibit the hydroxyapatite phase, regardless of the amount of silicon ranging between $0 \leq \text{wt}\%\text{Si} \leq 3.51$ (or $0 \leq y \leq 1.25$), as observed on the diffractograms (Fig. 2). A more accurate description of the infrared spectra of the as-synthesized SiHA powders is available in a complementary article [95].

### 3.2.6.2 Calcined samples – general observations

Fig. 10 compares the infrared spectra of heat-treated HA and SiHA powders (1000°C/15 h). The HA and Si$_y$HA with $0 \leq y \leq 1.0$ powders mainly present bands characteristic of hydroxyapatite with the $v_1$ (962 cm$^{-1}$), $v_2$ (473 cm$^{-1}$), $v_3$ (1021 and 1085 cm$^{-1}$) and $v_4$ (562 and 600 cm$^{-1}$) modes of PO$_4^{3-}$, as well as the stretching ($v_S$: 3572 cm$^{-1}$) and librational ($v_L$: 629 cm$^{-1}$) modes of hydroxide groups [81, 92, 94]. A shoulder at 947 cm$^{-1}$ was also observed for HA and SiHA powders. The relative intensity of this shoulder was slightly higher for SiHA than for HA, but did not seem to be a function of the amount of Si. Besides the low hydroxyapatite vibrations, the spectrum of the Si$_{1.25}$HA powder presents mainly the characteristic bands of $\alpha$-TCP and traces of $\beta$-tricalcium phosphates ($\beta$-TCP): the weak band at 495 cm$^{-1}$ ascribed to the O-P-O $v_2$ vibrational mode of $\beta$-TCP, the bands due to the splitting of the $v_4$ mode at 567 cm$^{-1}$ in $\alpha$-TCP (551, 560, 580, 595 and 611 cm$^{-1}$), the bands at 945 and 955cm$^{-1}$ assigned to the degenerated symmetric P-O stretching vibration of the phosphate ions ($v_1$) in $\beta$- and $\alpha$-TCP, respectively, and the bands corresponding to the strong asymmetric P-O stretching mode ($v_3$) for $\beta$-TCP (988, 1025 cm$^{-1}$) and $\alpha$-TCP (988, 1013, 1025, 1031 and 1055cm$^{-1}$) [52,
This result is complementary to the X-ray diffraction pattern (Fig. 3) and indicates that the Si_{1.25}HA apatitic precipitate decomposes during heat treatment with the formation of TCP and amorphous silica species.

### 3.2.6.3. Calcined samples – accurate IR band assignment of the pure SiHA phase

As opposed to HA, the FT-IR spectra of the calcined Si_yHA powders, with 0 ≤ y ≤ 0.75, exhibit nine new bands or shoulders (sh) at 491 (sh), 504, 528, 750, 840, 893, 930 (sh), 985 (sh) and 1002 (sh) cm\(^{-1}\). The intensity of these new vibrations increased with the amount of Si, except for the weak band at 840 cm\(^{-1}\) and the shoulder at 930 cm\(^{-1}\) which remained relatively constant. Conversely, the intensity of the absorption bands attributed to the four vibrational modes of PO_4^{3-} and both modes of OH\(^-\) in hydroxyapatite decreased as the amount of Si increased in the range 0 ≤ y ≤ 1.00 (e.g. \(v_s\) OH : 0.49 (HA) > 0.39 (Si_{0.25}HA) > 0.35 (Si_{0.50}HA) > 0.12 (Si_{0.75}HA) > 0.07 (Si_{1.00}HA)). More precisely, the \(v_L\) mode decreased to the detection limit for y = 1, while the stretching the OH\(^-\) band decreased and broadened. In fact, two new vibrations at 3565 and 3552 cm\(^{-1}\), close to the main one at 3572 cm\(^{-1}\), were detected (Fig. 10). This confirms the mechanism of charge compensation with the creation of OH\(^-\) vacancies, and highlights the new environment and distribution of OH\(^-\) ions along the channels of the SiHA structure [93, 97] and, more importantly, the formation of low hydrogen bonding between closed O^{2-} and OH\(^-\) ions (H-bonds cause a shift of 20 cm\(^{-1}\) to the low wavenumber of the main band at 3572 cm\(^{-1}\) [98]). The FT-IR spectrum of Si_{1.0}HA displays the same eight new bands as those detected on Si_{0.25}HA, Si_{0.50}HA and Si_{0.75}HA. However, their relative intensity decreased significantly, reaching the detection limit for the band at 750 cm\(^{-1}\) and with a complete disappearance of the bands at 491, 504, and 528 cm\(^{-1}\). Additionally, seven new vibrations appeared at 500, 515, 535, 683, 798 and 873 cm\(^{-1}\). They were accompanied by an intensification of the shoulder at 947 cm\(^{-1}\). The bands at 683, 798 and 870 cm\(^{-1}\) were also detected in spectrum of Si_{1.25}HA calcined powder. According to these results
and an accurate review of the bibliography, summarized in Table 4, it is clear that the nine vibrations at 493, 504, 528, 750, 840, 893, 930, 985 and 1002 cm\(^{-1}\) are related to the incorporation of Si into the hydroxyapatite structure. The changes detected in the sample Si\(_{1.0}\)HA were due to the start of decomposition of the SiHA phase, emphasized by a decrease in the nine characteristic bands and the formation of an amorphous silica phase which presents vibrations at 500, 515, 535, 683, 798 and 873 cm\(^{-1}\). Silica was not detected by NMR due to the low amount of \(^{29}\)Si in the samples (high detection limit). The shoulder at 947 cm\(^{-1}\) could be attributed to \(\beta\)-TCP, which exhibits a symmetric P-O stretching mode \(\nu_1\) at 945 cm\(^{-1}\) [96]. However, this assumption is not convincing since other bands assigned to \(\beta\)-TCP are absent. In fact, this shoulder is a contribution of the Si-O in plane stretching vibrations in Si-OH and Si-O\(^-\) on the surface of the grains, and the symmetric stretching \((\nu_1)\) vibration of orthophosphate groups in the neighborhood of OH vacancies along the channels. The former, reported at 950 cm\(^{-1}\) (see Table 4), shifted toward lower frequencies due to hydrogen bonding between surface groups, thus creating the component at 930 cm\(^{-1}\). The band at 950 cm\(^{-1}\) is commonly observed for oxy-hydroxyapatite \(\text{Ca}_{10}(\text{PO}_4)_6(\text{OH})_{2-2x}\text{O}_x(\text{VOH})_y\), [93, 99-102]. Therefore, in the context of this study, OH vacancies \((\text{VOH})\) are due first to the incorporation of SiO\(_4\) into the HA structure, and second to the partial dehydration of SiHA to SiOH as follows:

\[
\text{Ca}_{10}(\text{PO}_4)_{6-y}(\text{SiO}_4)_y(\text{OH})_{2-y}(\text{VOH})_y \leftrightarrow \text{Ca}_{10}(\text{PO}_4)_{6-y}(\text{SiO}_4)_y(\text{OH})_{2-y-2x}\text{O}_x(\text{VOH})_y+x\text{H}_2\text{O} \quad (7)
\]

### 3.2.6.4. Calcined samples – OH vacancies and the limit of incorporation of Si in HA

The amount of hydroxide (%OH) in the silicon-substituted hydroxyapatite structure is reported versus the amount of silicon (mol mol\(_{\text{SiHA}}^{-1}\)) in Fig. 11. The theoretical values of OH were plotted by considering the formula \(\text{Ca}_{10}(\text{PO}_4)_{6-y}(\text{SiO}_4)_y(\text{OH})_{2-y}(\text{VOH})_y\) (i.e. %OH = 100 - 50 \(y\)). The experimental values were determined at room temperature on the calcined samples (1000°C/15 h) by means of the integrated area of the \(\nu_L\) and \(\nu_S\) OH bands. The average of the
integrated areas obtained for HA (y=0) was assumed to be representative of the full occupancy of the OH position within the channels (100% OH). The amount of OH vacancies \( V_{\text{OH}} \) determined experimentally was always higher than the theoretical one with respect to the chemical formula \( \text{Ca}_{10}(\text{PO}_4)_{6-y}(\text{SiO}_4)_y(\text{OH})_{2-y}(V_{\text{OH}})_y \). In other words, the dehydration of the calcined powders was greater than expected, and a higher Si doping level led to a greater difference. Thus, the incorporation of Si into the hydroxyapatite structure generates OH vacancies first to maintain the charge balance \( \text{Ca}_{10}(\text{PO}_4)_{6-y}(\text{SiO}_4)_y(\text{OH})_{2-y}(V_{\text{OH}})_y \) and second from another phenomena, which is an increasing function of the amount of Si. A possible explanation is an increase of the mobility of OH ions along the channel axis (local disorder) and their subsequent rate of diffusion and debonding from the HA lattice [103]. HA decomposition is a dynamic process, mainly controlled by the degree of dehydration [104-106]. In fact, the thermal stability of hydroxyapatite was found to depend on the fraction of \( V_{\text{OH}} \) and \( \text{O}^2- \) ions in the channels. Several “critical values” of \( V_{\text{OH}} \) in OHA \( \text{Ca}_{10}(\text{PO}_4)_6(\text{OH})_2 \cdot 2x\text{O}_x(V_{\text{OH}})_x \) have been reported beyond which the apatite channel structure is destroyed: \( x \leq 0.80 \) [103], \( x \leq 0.75 \) [107], \( x \leq 0.50 \) [89]. However, the most probable limit comes from work of Heughebaert and Montel on the crystallization of calcium phosphates during precipitation [108, 109]. They determined that the apatitic structure appears only when at least 25% of the OH positions along the channels are really occupied. In other words, the apatitic structure appears when \( V_{\text{OH}} < 1.5 \). Therefore, the theoretical maximum limit of incorporation of Si into a hexagonal apatitic structure is \( y < 1.5 \) as regards the formula \( \text{Ca}_{10}(\text{PO}_4)_{6-y}(\text{SiO}_4)_y(\text{OH})_{2-y}(V_{\text{OH}})_y \). However, due to the dehydration reaction (Eq. 7), this limit is a function of the temperature and the atmosphere of heat treatment. This assumption can explain why the Si_{1.0}HA and Si_{1.25}HA powders, which showed an apatitic structure after calcination at 400°C for 2 h, were not thermically stable at 1000°C for 15 h in contrast to SiHA with a silicon content ranging between \( 0 \leq y \leq 0.75 \). Control of the degree of hydroxylation, \textit{i.e.} the partial
steam pressure $p_{\text{H}_2\text{O}}$, is necessary to control the thermal stability of the SiHA phase. This assumption is confirmed by the infrared spectra obtained from the Si$_{1.0}$HA sample heat treated at 1000°C for 15 h under different partial pressures of steam, $p_{\text{H}_2\text{O}}$, in mbar (Fig. 12); a higher $p_{\text{H}_2\text{O}}$ led to a more stable silicon-substituted apatite phase. Thus, the Si$_{1.0}$HA powder, after calcination under high steam pressure ($p_{\text{H}_2\text{O}} = 200$ mbar), presented only the characteristic bands of SiHA with traces of amorphous silica at the IR spectroscopy detection limit ($\approx 0.1$wt%), whereas the same sample heat-treated under dry argon gas ($p_{\text{H}_2\text{O}} = 0$ mbar) contained amorphous silica species as well as TCP ($\alpha$ and $\beta$). Moreover, the vibrations related to Si-O in SiHA (750 and 890 cm$^{-1}$) as well as the vibrational modes $\nu_3$ and $\nu_4$ of the phosphate groups in HA decreased with a decrease in $p_{\text{H}_2\text{O}}$. Conversely, the vibrations due to amorphous silica species, at 683, 798 and 873 cm$^{-1}$, and TCP (e.g. 731, 942, 997 and 1137 cm$^{-1}$) increases with a decrease in $p_{\text{H}_2\text{O}}$.

These results confirm that the formation of amorphous silica and the decomposition of the SiHA phase are correlated. More precisely, the detection of amorphous silica indicates the start of decomposition, which is followed by the formation of TCP ($\alpha$ and $\beta$).

Finally, Trombe and Montel reported the variation of $a$ from 9.421 Å to 9.402 Å, without a significant variation in the $c$ lattice parameter, when HA transforms into OHA, Ca$_{10}$(PO$_4$)$_6$(OH)$_0.5$O$_{0.75}$V$_{0.75}$O$_{0.75}$ [102]. Thus, SiHA dehydration can explain the random fluctuations in the $a$ lattice parameter observed in this work and in the literature [30].

4. Conclusion

This paper presents a new route to synthesize monophasic silicon-containing hydroxyapatites Ca$_{10}$(PO$_4$)$_6$Si$_{0.5}$O$_{12}$V$_y$ (SiHAs) with controlled stoichiometry. The combination of solid state nuclear magnetic resonance (NMR) and IR spectroscopy (IR) demonstrated that silicate substitution for phosphate creates OH vacancies along the channels. Moreover, Si substitution increases the local disorder and the mobility of OH ions along these channels and
their subsequent rate of diffusion and debonding from the HA lattice. The control of the degree of hydroxylation of the SiHA phases, i.e. their thermal stability, can be carried out by means of the partial steam pressure, $p_{H_2O}$. Additionally, an exhaustive description of the infrared bands related to the incorporation of silicate groups into the HA structure was established. It allowed us to determine that the literature has incorrectly attributed some infrared bands to silicate groups in the apatite structure. These bands, e.g. 683, 798 and 873 cm$^{-1}$, due in fact to amorphous silica phases, raise questions regarding the phase purity of the great majority of biologically evaluated SiHA bioceramics.

Acknowledgments

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References


[96] Jillavenkatesa A, Condrate RA. The infrared and Raman spectra of b and a tricalcium phosphate (Ca$_3$(PO$_4$)$_2$). Spectroscopy letters 1998;31:1619-34.


Fig. 1. XRD patterns of heat treated Si$_{0.50}$HA powders prepared at 50°C and maintained at pH 9.5 and 10.8.

Fig. 2. X-ray diffraction patterns of raw SiHA powders synthesized at pH = 10.8 with different silicon concentration.
Fig. 3. X-ray diffraction patterns of heat treated (1000°C/15h) SiHA powders synthesized at pH = 10.8 with different silicon concentration.

Fig. 4: (a) TEM micrograph of heat treated Si$_{1.0}$HA crystallites (1000°C/15h), (b) High-resolution lattice image of a heat treated Si$_{1.0}$HA particle (1000°C/15h) along the zone axis [01-1]; plane spacing of 8.2Å.
Fig. 5: (a) selected area electron diffraction (SAED) pattern of heat treated $\text{Si}_{1.0}\text{HA}$ powder ($1000^\circ\text{C}/15\text{h}$), (b) SAED of heat treated $\text{Si}_{1.0}\text{HA}$ powder ($1000^\circ\text{C}/15\text{h}$) for the $[0 1 0]$ zone axis, $c/a = 0.730$
Fig. 6. (a) $^{31}$P and (b) $^1$H MAS-NMR spectra of the HA, Si$_{0.50}$HA and Si$_{1.00}$HA powders calcined at 1000°C for 15h.

Fig. 7. Comparison between $^{31}$P MAS-NMR and CP MAS NMR spectra of (a) HA and (b) Si$_{0.50}$HA powders calcined at 1000°C for 15h.
Fig. 8. $^1$H-$^{31}$P HETCOR CP MAS experiment recorded on (a) Si$_{1.00}$HA powder calcined at 400°C for 2h, and (b) Si$_{0.50}$HA powder calcined at 1000°C for 15h.

Fig. 9. $^{29}$Si MAS-NMR spectra of the Si$_{1.00}$HA powder calcined at 400°C for 2h and 1000°C for 15h.
Fig. 10. FTIR spectra of calcined HA and SiHA powders (1000°C / 15h) collected at room temperature, symbol ● shows additional vibrations for Si1.0HA at 500, 515, 535 cm⁻¹.

Fig. 11. OH content into the apatitic channels versus the Si content, for theoretical SiHA structure (y V_{OH}) and samples calcined at 1000°C for 15h (determined from integrated areas.
of $\nu_L$ and $\nu_S$ bands).

Fig. 12. FTIR spectra of calcined Si$_{1.0}$HA powder (1000°C / 15h) under different partial pressure of steam ($p_{\text{H}_2\text{O}} = 0, 80, 200$ mbar and air).
Table 1
Chemical conditions of preparation of HA and Si<sub>y</sub>HA samples and final composition of the calcined powders (1000°C/15h).

<table>
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<th>Sample</th>
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<th>Composition of the final calcined powders (1000°C/15h)</th>
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Table 2

NMR parameters.

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<td>60</td>
<td>SPINAL-64 (45kHz)</td>
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</table>
Table 3.

Lattice parameters, unit cell volume and mean size of the crystallites of heat-treated powders at 1000°C for 15h evaluated by Rietveld refinement.

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<tr>
<th>Sample</th>
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<th>Volume of the unit cell / Å³</th>
<th>Crystallites mean size / nm</th>
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</tr>
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<td>6.884 ± 3 10⁻⁵</td>
<td>1058 ± 1</td>
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<tr>
<td>Si₀.25HA</td>
<td>9.423 ± 4 10⁻⁵</td>
<td>6.892 ± 4 10⁻⁵</td>
<td>1060 ± 1</td>
</tr>
<tr>
<td>Si₀.50HA</td>
<td>9.428 ± 2 10⁻⁵</td>
<td>6.899 ± 2 10⁻⁵</td>
<td>1062 ± 1</td>
</tr>
<tr>
<td>Si₀.75HA</td>
<td>9.423 ± 2 10⁻⁴</td>
<td>6.908 ± 2 10⁻⁴</td>
<td>1062 ± 1</td>
</tr>
<tr>
<td>Si₁.₀₀HA</td>
<td>9.416 ± 7 10⁻⁵</td>
<td>6.920 ± 5 10⁻⁵</td>
<td>1063 ± 1</td>
</tr>
</tbody>
</table>
Table 4

IR bands and shoulders assignment proposed by our work

<table>
<thead>
<tr>
<th>$\lambda$ / cm$^{-1}$</th>
<th>Mode assignments proposed by this work</th>
<th>Literature support</th>
</tr>
</thead>
<tbody>
<tr>
<td>1002*</td>
<td>Degenerated PO$_4^{3-}$ asymmetric stretching ($\nu_3$)</td>
<td>[94, 110]</td>
</tr>
<tr>
<td>985</td>
<td>Si-OH at SiHA surface / and/or / Si in SiHA</td>
<td>[110-112]</td>
</tr>
<tr>
<td>947</td>
<td>Oxyapatite Si-OH at SiHA surface</td>
<td>[93, 99, 100, 102, 113]</td>
</tr>
<tr>
<td>930**</td>
<td>Si-OH at SiHA surface / and/or / Si in SiHA</td>
<td>[114-116]</td>
</tr>
<tr>
<td>893</td>
<td>Si in SiHA</td>
<td>[8, 113, 118]</td>
</tr>
<tr>
<td>873</td>
<td>Si in silica</td>
<td>[112, 113, 122, 123]</td>
</tr>
<tr>
<td>840</td>
<td>Si-R</td>
<td>[118, 124]</td>
</tr>
<tr>
<td>798</td>
<td>Si in silica</td>
<td>[34, 110, 112-115, 117, 120, 122, 125-127]</td>
</tr>
<tr>
<td>750</td>
<td>Si in SiHA</td>
<td>[32, 54, 74]</td>
</tr>
<tr>
<td>683</td>
<td>Si in silica</td>
<td>[113, 114, 116, 126, 127]</td>
</tr>
<tr>
<td>535</td>
<td>Si in silica</td>
<td>[128]</td>
</tr>
<tr>
<td>528</td>
<td>Si in SiHA</td>
<td>[111, 118, 121, 128]</td>
</tr>
<tr>
<td>515</td>
<td>Si in silica</td>
<td>[127-130]</td>
</tr>
<tr>
<td>504</td>
<td>Si in SiHA</td>
<td>[32, 56, 118]</td>
</tr>
</tbody>
</table>

* degeneration of the $\nu_3$ PO$_4^{3-}$ domain at 1020 cm$^{-1}$ due to the SiO$_4^{4-}$ in the environment of the phosphate ions

** degeneration of the band at 947 cm$^{-1}$