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Comparison of Physical-chemical and Mechanical Properties of Chlorapatite and Hydroxyapatite Plasma Sprayed Coatings

Imane Demnati¹, David Grossin¹, Olivier Marsan¹, Ghislaine Bertrand¹, Gérard Collonges², Christèle Combes¹, Maria Parco³, Inigo Braceras³, Joel Alexis⁴, Yannick Balcaen⁴ and Christian Rey¹,*

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Abstract: Chlorapatite can be considered a potential biomaterial for orthopaedic applications. Its use as plasma-sprayed coating could be of interest considering its thermal properties and particularly its ability to melt without decomposition unlike hydroxyapatite. Chlorapatite (ClA) was synthesized by a high-temperature ion exchange reaction starting from commercial stoichiometric hydroxyapatites (HA). The ClA powder showed similar characteristics as the original industrial HA powder, and was obtained in the monoclinic form. The HA and ClA powders were plasma-sprayed using a low-energy plasma spraying system with identical processing parameters. The coatings were characterized by physical-chemical methods, i.e. X-ray diffraction (XRD), Fourier transform infrared spectroscopy (FTIR) and Raman spectroscopy, including distribution mapping of the main phases detected such as amorphous calcium phosphate (ACP), oxyapatite (OA), and HA or ClA. The unexpected formation of oxyapatite in ClA coatings was assigned to a side reaction with contaminating oxygenated species (O₂, H₂O). ClA coatings exhibited characteristics different from HA, showing a lower content of oxyapatite and amorphous phase. Although their adhesion strength was found to be lower than that of HA coatings, their application could be an interesting alternative, offering, in particular, a larger range of spraying conditions without formation of massive impurities.

Keywords: Chlorapatite, FTIR, hydroxyapatite, low energy plasma spraying, mechanical testing, Raman, XRD.

INTRODUCTION

Orthopaedic implants, especially those intended to restore the function of load-bearing joints (hip, knee, shoulder, elbow, ankle, finger...) of aged or injured human beings, should possess several attributes to meet the high level of mechanical stresses, wear, and fatigue in the course of normal activity. Although they may differ depending on the type of joints, these attributes include mechanical properties (elasticity, yield stress, ductility, toughness, and wear resistance) as well as biological properties (biocorrosion resistance, biocompatibility, good and long-term osseointegration). Metallic materials are most commonly used for load bearing implants. Stainless steel, cobalt alloys and titanium alloys are accordingly commercially available for bone substitutes, artificial joints and dental implants [1, 2]. Durability and functionality of the implant are governed mainly by the properties of the bulk material, whereas the biological response depends, among others, on the surface characteristics of the implantable device. Indeed, a crucial step for the implant stability is the formation of the effective interface between the bone and the implanted material. Several surface characteristics favour this osseointegration: mechanical interlocking related to surface roughness and physical characteristics, and direct chemical interactions involving bonds between the implant surface and bone tissue constituents. To promote the osseointegration of metallic implants, morphological (3D texturing, surface roughness and porosity) and chemical (ion implantation, grafting, and coating) modifications of the implant surface have been proposed [3, 4]. Since the mid-1980s, plasma sprayed hydroxyapatite (HA) (Ca₁₀(Po₄)₆(OH)₂) coatings appear as the most developed and successful treatments of metallic, especially titanium-based, implants for a fast integration to bone tissues [5-13]. Indeed, atmospheric plasma spraying (APS) is the most widely used commercial technique due to its process simplicity, low cost, and bulk production [14]. During plasma spray, the material to be deposited (feedstock), typically as powder, is introduced into the core of a plasma plume emanating from a d.c. torch. The powder is fully or partially melted, and accelerated towards the work piece, upon which it impacts
and cools down to form a coating by overlapping of splats. Depending on the plasma-forming gases, electrical characteristics, feedstock attributes, and kinematics conditions, the thermal input transferred to the in-flight particles and coating, together with the cooling rate, can result in severe powder decomposition and in lack of crystallinity. Starting from HA feedstock it is noteworthy that plasma spray can induce the formation of amorphous calcium phosphate (ACP), tricalcium phosphate (α or β-TCP), tetracalcium phosphate (TTCP), calcium oxide (CaO) and dehydroxylated products (oxyhydroxyapatite or oxyapatite) [5, 7, 15-21]. The decomposition sequence occurs along the powder trajectory in the plasma jet that is determined mainly by the particles’ size, density and their dwell-time in conjunction with many parameters, for example the viscosity and density of the plasma. The degree of crystallinity of plasma sprayed HA coatings is usually between 65 and 85 vol. % (in some operating conditions it may reach 93 vol. %) [7, 15, 22-26]. Crystalline HA is seen as a crucial factor for long-term coating stability in vivo. Although the presence of relatively soluble non-apatitic calcium phosphate salts such as ACP, TCP, and TTCP has been found to accelerate the bone attachment, these phases are related to long-term non-uniform coating degradations that adversely affect the prosthetic stability of the implants [4, 27-34]. To limit the amount of easily soluble CaP phases, the steep temperature gradients during plasma spraying have to be reduced by increasing the substrate temperature and controlling the cooling rate and/or by limiting the thermal transfer. One method to control the degree of melting of the powder could be achieved by micro plasma spray (MPS, also called MIPS) or low-energy plasma spray (LEPS) techniques [35-42]. Due to the low heat input of these plasma torches, overheating of the powder particles as well as excessive local overheating of the coating can be avoided. LEPS also allows the production of highly porous (around 20 vol. %)-thick coatings (about 250 µm) without accumulation of residual stresses (tensile stresses are detrimental because they promote cracks and accelerate dissolution) and without negatively affecting the mechanical properties (fracture toughness of about 0.6 MPa.m\(^{1/2}\)). In a previous work, we have implemented such an innovative low energy plasma spray system that has resulted in producing thin dense HA coatings (about 250 µm) without accumulation of residual stresses and has been shown to lead to formation of stoichiometric hydroxyapatite, and has been shown to lead to formation of stoichiometric HA.

The objective of the current study is to compare chlorapatite and hydroxyapatite coatings obtained by a LEPS gun. Until today only a few authors have worked on the substitution of chloride ion in hydroxyapatite, and no commercial product is available. Hence, our first aim was to develop a simple and fast reaction synthesis transposable at an industrial scale. A method is first described to prepare easily chlorapatite from commercial HA preserving similar powders characteristics. The main physical-chemical and mechanical properties of the coatings obtained with a low energy plasma spraying process are compared using standardized procedures and specific techniques such as Raman microscopy mapping of the main mineral phases found in the coating and three points mechanical bending with stiffener.

**POWDER PREPARATION AND CHARACTERISTICS**

The hydroxyapatite used in this study was a commercial product (Teknimed, L’Union, France). The XRD analyses, according to ISO standard (ISO 13779-3: 2008), did not detect any crystalline impurity (Fig. 1) and the FTIR spectrum (Fig. 2) was the same as that of stoichiometric hydroxyapatite.

Regarding the synthesis of chlorapatite, several methods have been proposed [49, 50]. Precipitation methods in aqueous media do not lead generally to stoichiometric chlorapatite and the apatite obtained contains always hydroxide ions. Several dry synthesis methods giving stoichiometric CIA with a high purity have been proposed. The reaction of CaCl\(_2\) on Ca\(_{3}\)(PO\(_4\))\(_2\) (800-1100°C) [50] results in CIA with a high yield, especially when using an excess of CaCl\(_2\), which can be removed by washing. Other methods can be used, especially Cl-OH ion exchange reactions on HA (800-1100°C), which offers the advantage of using an existing commercial product. Several chlorinating agents can be used for this exchange reaction, such as CaCl\(_2\):

\[
\text{CaCl}_2 + \text{Ca}_{10} \text{(PO}_4\text{)}_6 \text{(OH)}_2 \rightarrow \text{Ca}_{10} \text{(PO}_4\text{)}_6 \text{Cl}_2 + \text{CaO} + \text{H}_2\text{O} \hspace{1cm} (1)
\]

This reaction leads to the formation of CaO, which is difficult to remove. Other chlorinating agents could be used and more interestingly, chlorinating gases, such as HCl, which would not produce any residues except water. This reaction has been studied by Hoekstra [49] at about 1050°C and has been shown to lead to formation of stoichiometric CIA:

\[
\text{Ca}_{10} \text{(PO}_4\text{)}_6 \text{(OH)}_2 + \text{HCl} \leftrightarrow \text{Ca}_{10} \text{(PO}_4\text{)}_6 \text{Cl}_2 + \text{H}_2\text{O} \hspace{1cm} (2)
\]

This reaction is supposed not to alter the powder characteristics (e.g. grain size, shape). One of the difficulties in such a reaction is to handle hydrogen chloride gas, and, to overcome this difficulty, we choose as a precursor of HCl, NH\(_4\)Cl which decomposes at about 340°C under atmospheric pressure, into NH\(_3\) and HCl. In such a synthesis we should keep in mind, that non-stoichiometry has been reported to occur in CIA heated at high temperature, which has been assigned to the release of CaCl\(_2\) [51]:

\[
\text{Ca}_{10} \text{(PO}_4\text{)}_6 \text{Cl}_2 \rightarrow \text{Ca}_{10-x} \text{(PO}_4\text{)}_6 \text{Cl}_2 \cdot 2x + x\text{CaCl}_2 \hspace{1cm} (3)
\]

Previous attempts to use the reaction of HA with NH\(_4\)Cl by A. Hoekstra [49] have probably failed to produce...
stoichiometric CIA because the treatment temperature (1180°C) was too high. The second point to consider is the fast hydrolysis of CIA into HA when traces of water molecules are present. Thus, we shall use a temperature as low as possible, resulting in a reasonable reaction rate and enabling to preserve a dry atmosphere.

The ion exchange reaction was performed in a tubular oven under a flux of dry nitrogen enriched with sublimated NH₄Cl in large excess placed in a zone of the tubular oven at about 400°C [52]. The unreacted HCl in the gas flow recombines with NH₃ and regenerates NH₄Cl, which crystallizes at the outlet of the oven and could possibly be recycled. The reaction was fast at 950°C and a few grams of HA were transformed into CIA in about 15 minutes. However, for production treatments of the order of several hundreds of grams on partly sintered HA grains of industrial HA the treatment time was two hours [52].

The XRD analyses (Fig. 1) of the samples obtained indicated the formation of chlorapatite in its monoclinic form, which is the stable low temperature isomorph. The monoclinic CIA has been shown to correspond to an order among the Cl⁻ ions of the chlorapatite [53]. The tiny peaks of the superstructure related to this ordering can be distinctly seen on the XRD diagram Fig. (1). The transition temperature to the hexagonal structure of CIA occurred as expected at about 200°C. The unit cell dimensions (Table 5) appeared consistent with that of a stoichiometric chlorapatite.

The morphology of the HA and CIA powders are shown in Fig. (3). Their main characteristics (Ca/P, particle size, specific surface area, flowability) are reported in Table 1.

The grains in both samples show irregular shapes related to the preparation of HA granules by grinding of sintered particles and subsequent sieving. The surface irregularities can be seen on CIA grains, which were not apparent on HA ones. Granulometry (Malvern Matersizer 2000) revealed a similar grain distribution for both HA and CIA powders. The grain sizes extended from 0.7 to 140 µm with a main distribution peak between 50 and 80 µm and smaller ones between 0.7-3µm and 3-20 µm [52]. On chlorination, the median grain size d₀.₅ has decreased, presumably due to fragmentation of the initial HA grain during the chlorination process. The flowability was determined, according to the recommendations of the European Pharmacopoeia, by measuring the flow time of 100 g of powder through a funnel with specified dimensions [52]. The flowability of the

The other hand, chlorinating treatments have been shown to remove several impurities present in the apatite structure as volatile chlorides and this reaction has been proposed for the extraction of transition elements, rare earths and uranides from apatitic ores [54]. Although we did not determine the effect of the chlorination treatment on the level of impurities, this possibility could be an interesting additional advantage of the chlorination treatment.

FTIR data (Fig. 2) confirm the removal of the OH⁻ ions, the stretching (3570 cm⁻¹) and librational (633 cm⁻¹) lines of which cannot be seen on the spectra after chlorination. The lines characteristic of OH-Cl interactions in solid-solutions of HA and CIA, at 3495, 3510 cm⁻¹ [55] cannot be observed either, suggesting a complete replacement of OH⁻ ions by Cl⁻ ions in the apatite structure.

The theoretically stoichiometric HA should also exist as a monoclinic apatite at room temperature. However, most industrial HA crystallize in the hexagonal systems. This anomaly is often attributed to the existence of defects and impurities, which stabilizes the hexagonal form of HA. On the other hand, chlorinating treatments have been shown to remove several impurities present in the apatite structure as volatile chlorides and this reaction has been proposed for the extraction of transition elements, rare earths and uranides from apatitic ores [54]. Although we did not determine the effect of the chlorination treatment on the level of impurities, this possibility could be an interesting additional advantage of the chlorination treatment.

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<table>
<thead>
<tr>
<th>Powders</th>
<th>Ca/P</th>
<th>Specific surface area (m²/g)</th>
<th>Granulometry d₀.₅ (µm)</th>
<th>Flowability (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HA</td>
<td>1.66 ± 0.02</td>
<td>1.44 ± 0.07</td>
<td>69.5</td>
<td>27.26 ± 3.44</td>
</tr>
<tr>
<td>CIA</td>
<td>1.67 ± 0.02</td>
<td>1.48 ± 0.15</td>
<td>56.5</td>
<td>4.75 ± 0.32</td>
</tr>
</tbody>
</table>
powders appeared significantly different and this observation might be possibly related to a change in surface characteristics as observed by SEM, although this remains difficult to elucidate.

In addition, Rietveld analyses of XRD data reveal that the chlorination treatment induced, unexpectedly, a decrease in the average size of the ClA crystals compared to the industrial HA crystals in the starting powder (Tables 4 and 5).

Comparison of the phase diagrams, CaO-P₂O₅ for HA and CaCl₂-Ca₃(PO₄)₂ for ClA [50, 56] indicate a decomposition of HA on heating, whereby ClA melts without decomposition. This property is a major argument in favour of the use of ClA for producing plasma sprayed coatings: the formation of foreign phases is expected to be limited, except for the amorphous phase resulting of the quenching of a melted phase.

The comparison of the solubility of HA and ClA, which could determine possible differences in bioreorption of the coatings, is also interesting. However, published data on ClA solubility are scarce and provide incoherent values. Thus ClA has been reported to be less soluble than HA [57], although thermodynamic data strongly suggest that it should be more soluble than HA with a solubility product pKₘ₄(CIA) = 108 at 25°C (pKₘ₄(HA) = 116.8 for Ca₁₀(PO₄)₆(OH)₂) [58]. However, considering the concentration of Cl⁻ ions in body fluids of more than 10⁵ times higher than that of OH⁻ in serum at physiological pH, the effect of this difference would be minor. It should be noticed that the incorporation of chloride ions in biological apatites, especially enamel, has been confirmed [59].

**COATINGS DEPOSITION**

**Thermal Spraying**

The powders (HA and ClA) were sprayed in air on surgical grade (grade 4) Ti disks (12 mm in diameter and 3 mm in thickness), using a low energy plasma spray (LEPS) device [43]. This system was an experimental low energy plasma spray system designed and developed by Inasmet-Tecnalia (Spain) from commercial Sulzer Metco torch (Switzerland). Ti substrates were previously degreased and then sandblasted with corundum P150 using a pressure of 2 bars to produce a surface roughness (Ra) of about 1 µm. Coupons were separately sprayed with chlorapatite or hydroxyapatite under the same thermal spraying conditions (Table 2). For practical reasons, the gun operated with standard electrodes. The entire gun was cooled with water and a glass fibre-reinforced epoxy housing separated the electrical poles. A more detailed description of the LEPS system has been provided elsewhere [60]. Coatings with various thicknesses were obtained simply by changing the number of torch runs (1, 6, 12 and 20 runs) for both HA and ClA powders. We can see on Table 3 that the coating thickness depended not only on the number of runs, but also on the nature of the apatite: whatever the number of spray runs, the HA-coatings were always thicker than the ClA-coatings. This observation can be related to the decrease of the median grain size associated with the chlorination treatment, and possibly to a certain additional fragility of the grains, although other causes could be envisioned such as the viscosity of the melted phase for example, which is supposed to be totally liquid in the case of ClA whereas it may contain solid CaO in the case of HA.

**Table 2. Torch characteristics and plasma spray conditions**

<table>
<thead>
<tr>
<th>Torch Type</th>
<th>Sulzer Metco; Switzerland</th>
</tr>
</thead>
<tbody>
<tr>
<td>Powder level (kW)</td>
<td>15 kW</td>
</tr>
<tr>
<td>Current (A)</td>
<td>400</td>
</tr>
<tr>
<td>Gas</td>
<td>Argon</td>
</tr>
<tr>
<td>Spraying velocity (mm.s⁻¹)</td>
<td>200</td>
</tr>
<tr>
<td>Plasma gas rate (Ar; L.mm⁻¹)</td>
<td>24</td>
</tr>
<tr>
<td>Stand-off distance (mm)</td>
<td>40</td>
</tr>
<tr>
<td>Carrier gas rate (Ar; L.mm⁻¹)</td>
<td>4</td>
</tr>
<tr>
<td>Powder particle size (µm)</td>
<td>50-80</td>
</tr>
<tr>
<td>Powder feeding rate (g.mm⁻¹)</td>
<td>8</td>
</tr>
<tr>
<td>Number of spray runs</td>
<td>1; 6; 12; 20</td>
</tr>
</tbody>
</table>
PHYSICAL-CHEMICAL CHARACTERISTICS OF THE COATINGS

Structural and Chemical Composition of the Coatings

Surface and bulk composition of HA and ClA coatings were determined. The coating surface was analysed directly on the as-sprayed coatings. For the bulk characterizations, the coatings were scrapped and ground as indicated in the ISO standard (13779-3: 2008). Investigations on coating microstructure were also performed on sample cross-sections prepared as follows: the coupons were cut with a diamond disc (Struers) and then the half-coupon was embedded into an epoxy resin under vacuum. The embedded cross-sections were polished using a silicon carbide paper (1000 and 1200 grade SiC paper). Further polishing was conducted using 2400 then 4000 grade SiC paper to obtain a flat surface suitable for analysis.

X-ray Diffraction Analysis of the Coatings

Standards have been published concerning the coating’s chemical composition (Ca/P ratio), phase composition, and crystallinity (ISO 13779-2 and 3:2008). These standards are based essentially on X-ray diffraction (XRD) analyses.

X-ray diffraction studies on both powders and coatings were carried out in order to identify the composition of the coating (obtained after 12 spray runs) and determine its crystallinity, according to the ISO 13779-3: 2008 standard. A Seifert XRD-3000 diffractometer (Germany) with a CuKα radiation over a 2θ range of 20-65° was used. Patterns were collected at a step increment of 0.02°, with a collection time of 24 seconds at each step. According to the standard, the HA-coatings crystallinity (Cr) was defined as the ratio between the sum of integrated intensities of ten diffraction lines (∑A_{hkl}) of the coating diagram, and of the same ten lines of a fully crystalline HA reference powder (∑A_{hkl}^0): Cr = (∑A_{hkl})/ (∑A_{hkl}^0). The same approach was used to determine the ClA coating crystallinity. However, the (102) line used in crystallinity determination of HA is weak in ClA and it has a minor contribution in the crystallinity index; thus it was replaced by (201) line. Both coatings diagrams can be indexed in the hexagonal system indicating that a change of the structure of sprayed ClA powder has occurred. As we will see later, this structural modification is related to the alteration of the ClA crystals in the coatings and the formation of oxyapatite (OA) as well.

For both coatings, we noticed an increase of the full width at half maximum (FWHM) value of the diffraction lines compared to that of the initial powders (Figs. 4 and 5), which is generally assigned to two main phenomena: the decrease of the crystallite size, and the increase of the microstrain. In order to evaluate these contributions, Rietveld refinement analyses were performed using MAUD software. The crystallographic parameters determined for the HA and ClA initial powders and for the 12-run HA- and ClA-coatings are reported in Table 4 and Table 5, respectively. For HA- and ClA-coatings, we can notice both a significant increase in micro-strains and a decrease in crystallite size. However, these observations and the related calculations neglect an important cause of line broadening associated with the formation of heterogeneous solid solutions of oxy-hydroxyapatite and oxy-chlorapatite, which would result in broadened diffraction lines, corresponding to the sum of unresolved diffraction peaks of individual crystalline domains considered as homogeneous. For the OCIA this effect would result in a higher broadening of diffraction lines than for OHA, due to larger differences in unit-cell dimensions between OA and CIA [60]. The average unit-cell dimensions for CIA coatings can be compared to that of the powder considering that b=2a in the monoclinic CIA powder. The formation of solid solutions (OHA or OCIA) can be related to a clear decrease of the a unit-cell dimensions noticed on the coatings. This effect appears

<table>
<thead>
<tr>
<th>Number of runs</th>
<th>HA-coating (µm)</th>
<th>CIA-coating (µm)</th>
<th>HA-coating (µm.run⁻¹)</th>
<th>CIA-coating (µm.run⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>28 ± 4</td>
<td>16 ± 4</td>
<td>28 ± 4</td>
<td>16 ± 4</td>
</tr>
<tr>
<td>6</td>
<td>85 ± 4</td>
<td>60 ± 4</td>
<td>14.2 ± 0.7</td>
<td>10.0 ± 0.7</td>
</tr>
<tr>
<td>12</td>
<td>161 ± 4</td>
<td>120 ± 4</td>
<td>13.4 ± 0.3</td>
<td>10.0 ± 0.3</td>
</tr>
<tr>
<td>20</td>
<td>274 ± 4</td>
<td>179 ± 4</td>
<td>13.7 ± 0.2</td>
<td>9.0 ± 0.2</td>
</tr>
</tbody>
</table>

Table 3. Coating thickness as a function of the number of runs and of the type of apatite.

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![Fig. (4)](image-url) X-ray diffraction patterns of HA powder and coating (12 spray runs).
stronger for OClA due to the strong difference in unit-cell dimensions of ClA and OA just mentioned. However, in the case of ClA coatings, another possibility for the variations of the unit cell dimensions, compared to those of the ClA powder, would be the formation of a non-stoichiometric ClA at high temperature due to the release of CaCl$_2$ [51].

The calculated crystallinity ratio, determined according to the standard, was equal to 68% for HA-coating and to 98% for ClA-coating. The ClA-coating showed a much higher crystallinity ratio than HA-coating, although it was not determined in the same way. This observation should not be only related to the amount of amorphous phase formed in the coatings. Other causes can produce crystallinity variations such as fast recrystallization of small apatite crystals from the melt or the amorphous phase, or the formation of solid solutions with OA.

The Ca/P ratio of HA- and ClA-coatings obtained after 12 spray runs was determined according to the ISO 13779-3: 2008 standard on a scraped and ground coating, which had been heated at 1000°C during 15 h. The XRD patterns showed only apatite crystals for both HA- and ClA-coatings, indicating that plasma spraying using the LEPS system did not appear to alter the stoichiometry of the apatite.

**Vibrational Spectroscopy Analysis of the Coatings.**

Several additional characteristics of the apatite coatings are more difficult to determine using X-ray diffraction analyses: the main phase proportions and their distribution within the coating, and, more importantly, the amount of amorphous phase, which is not specifically considered in the ISO standard, although it could determine the coating behaviour *in vitro* and *in vivo*. A more accurate characterization of HA and ClA powders and the related plasma sprayed apatite coatings can be proposed, using vibrational spectroscopies [61]. Fourier transform infrared spectroscopy and Raman spectroscopy, with a special focus on Raman micro-spectroscopy imaging on HA- and ClA-coatings.
**FTIR Spectroscopy Characterization**

Both initial HA and CIA powders and scraped HA- and CIA-coatings were analysed by transmission Fourier transform infrared (FTIR) spectroscopy (Nicolet 5700 spectrometer, ThermoElectron) using KBr pellets. We examined especially the 1100-950 cm\(^{-1}\) (\(\nu_1\)PO\(_4\) and \(\nu_3\)PO\(_4\)) and 600-450 cm\(^{-1}\) (\(\nu_2\)PO\(_4\)) domains, corresponding to the different phosphate vibration bands. In these domains, FTIR spectra of both HA- and CIA-coatings Figs. (6 and 7) showed broader and less resolved bands than that of the initial powders. This phenomenon can be related to a decrease in crystallinity. The enhanced relative intensity of bands observed in the \(\nu_2\)PO\(_4\) domain (434 and 471 cm\(^{-1}\)), in the spectrum of the HA coating, can be correlated to the splitting of the \(\nu_1\)PO\(_4\) band and the decrease in the relative intensity of the OH bands at 3572 and 630 cm\(^{-1}\) (Fig. 6) which have been shown to correspond to the formation of oxyapatite [61]. Although less visible than in HA, the formation of oxyapatite is also apparent on FTIR spectra of CIA coatings (Fig. 7) which exhibit lines with a higher intensity in the \(\nu_2\)PO\(_4\) domain than the starting powder.

**Fig. (6).** FTIR spectra of HA powder and coating.

**Raman Spectroscopy Characterization**

All phases detected in a plasma-sprayed coating of HA can theoretically be distinguished by Raman spectroscopy, especially the most important ones, in some instances with better sensitivity than according to XRD analyses [52]. In addition, Raman scattering imaging allows local investigation of the coating’s heterogeneity [52, 62]. The \(\nu_1\)PO\(_4\) domain, with strong thin lines, seems well adapted for imaging [6, 63]. The amorphous calcium phosphate (ACP) especially is characterized by a broad Raman line around 950 cm\(^{-1}\) and all crystalline phases show lines distinct from that of apatite at 961 cm\(^{-1}\). The xyapatite (OA), rather difficult to distinguish by other methods, exhibits very specific Raman scattering lines at about 951 cm\(^{-1}\) and 966 cm\(^{-1}\) [64]. The only phase that cannot be detected in this domain is CaO.

**Fig. (7).** FTIR spectra of CIA powder and coating.

Both initial HA and CIA powders and the as-prepared HA- and CIA-coatings were analysed by Raman spectroscopy in the range of 100-1200 cm\(^{-1}\) and 3500-3600 cm\(^{-1}\) (Horiba Jobin-Yvon HR 800 spectrometer equipped with a helium-neon laser: \(\lambda = 632.8\) nm). The quantitative evaluation of the components of the \(\nu_1\)PO\(_4\) band was performed using LabSpec 5 software (Horiba). The \(\nu_1\)PO\(_4\) band Raman spectra were decomposed according to the same protocol for all the samples in order to facilitate their comparison [52]: a baseline correction was performed on each spectrum. Peak positions and some curve-fitting parameters were recorded and used as initial input in the curve-fitting program. Iterations continued until the best fit was obtained.

**Fig. (8).** Raman spectra of HA powder and coating.
The Raman spectra of HA and CIA scrapped coatings are presented in Figs. (8) and (9) together with those of the initial powders. The most intense Raman line, $\nu_1PO_4$, appeared the most altered after spraying. The integrated intensity of the different contributing sub-bands has been determined to estimate quantitatively the changes induced by spraying. The resulting curve-fittings are presented in Figs. (10) and (11) for HA- and CIA-coatings respectively. The detection of oxyapatite phase in both coatings confirmed the results obtained by FTIR spectroscopy; however, the proportion of OA appears always significantly less important in CIA than in HA coatings. The same observation is true for the amorphous phase.

Fig. (10). A-Gaussian-Lorentzian curve fitting of the $\nu_1PO_4$ Raman stretching domain of HA coatings (20 spray runs), with the LEPS plasma gun. (1) ACP: amorphous calcium phosphate; (2) and (4) OA: oxyapatite; (3) HA: hydroxyapatite.

Fig. (11). A-Gaussian-Lorentzian curve fitting of the $\nu_1PO_4$ Raman stretching domain of CIA coatings (20 spray runs), with the LEPS plasma gun. (1) ACP: amorphous calcium phosphate; (2) and (4) OA: oxyapatite; (3) CIA: Chlorapatite.

An accurate mapping of the different phases in apatite coatings can be obtained by micro-Raman imaging involving curve fitting in the $\nu_1PO_4$ domain, extraction of the characteristic line(s) associated with specific phases, and of their relative intensities [52]. It should be noticed that at this stage, without accurate standardization, and due to uncertainties in decomposition results. The data do not represent the real content of phases present in the coating but offer a base for visualizing the phase distribution in a first semi-quantitative approach. Figs. (12) and (13) show the main constituents distributions, i.e. HA or CIA, OA and ACP, in the cross section of HA and CIA coatings. The heterogeneity in the distribution of the constituents is immediately apparent. Domains with a high content in HA appear to be embedded in an amorphous phase acting as a binder. These observations are consistent with other descriptions of phenomenon occurring during plasma spraying [4, 5, 63, 65]. The nodules of HA, which appear as well delimited zones, are believed to correspond to the unmelt cores of the splats. In some cases smaller HA domains grouped in a zone may appear, corresponding possibly to the crash and dispersion into smaller particles of the solid core of splats impacting the substrate or to recrystallization of HA from the melted or amorphous phase. The OA rich domains are distributed more uniformly than the HA nodules and do not form well-defined zones with dominant concentrations. This distribution could result of a recrystallization process in the melted part of the splats, possibly associated with a direct formation of OA in the unmelt cores.

For CIA coatings the layered structure of the coating related to the splat accumulation is more apparent than in HA coatings, especially in the ACP mapping. The CIA domains appear elongated and rather diffuse, indicating a better spreading of the splats; they are surrounded by the amorphous phase. The OA seems more homogeneously distributed at the surface of CIA domains or within amorphous domains; in these domains, zones very poor in CIA and rich in OA can be clearly observed, suggesting that AO forms mainly in a recrystallization processes of the molten phase or the amorphous phase.

Therefore, Raman micro-spectroscopy can be used as a routine non-destructive tool to obtain rapid analysis of the composition of apatite coatings and even of different domains of a splat. This technique could be adapted for the chemical characterization of the coating directly on
biomedical implants. Image analyses could then be used to describe the coating characteristics with accuracy.

**Surface Roughness**

The average roughness of the CIA and HA coatings was determined by optical interferometry (Zygo) as the average of four measurements. The surface arithmetic roughness of HA and CIA coatings (Ra) was determined as a function of the number of spray runs Fig. (14). Ra increases strongly after the first run and reaches rapidly a constant value depending on the nature of the apatite: it is always higher in HA coatings than in CIA coatings. The increase of Ra is commonly observed and has been related to the structure of splats generally smaller at the fringe and larger in the centre [66]. These data indicate that HA splats are globally thicker than CIA ones, and seem consistent with the variations of the coating thickness according to the number of runs. The difference between HA and CIA coatings could be due, in part, to the alteration, already mentioned, of the particle size related to the chlorinating treatment.

The coating surfaces and cross-sections were studied by scanning electron microscopy (SEM). The coating cross-sections were carbon coated, then SEM micrographs were recorded in the secondary electron mode using a LEO 435 VP microscope. The typical surface morphologies of the as-sprayed HA- and CIA-coatings (12 spray runs) are presented as supportive materials.

**MECHANICAL PROPERTIES**

Several methods can be used to determine the adherence of a coating to its substrate. Concerning calcium phosphate coatings, an adherence measurement test was standardized (ASTM C633-01, ISO 13779-4) and the results obtained according to this procedure will be first presented. Among other methods used, the three point bending with a stiffener has also been applied and will be discussed.

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**Fig. (12).** Raman spectroscopy mappings of HA, OA and ACP in hydroxypatite coatings. Light blue oval: domain rich in HA, poor in ACP and with a little OA, tentatively associated with an unmelt core of a splat. White circle: domain rich in OA, poor in HA, with some ACP, attributed to the recrystallization of a molten domain or recrystallization of solid ACP phase.

**Fig. (13).** Raman spectroscopy mappings of CIA, OA and ACP in chlorapatite coatings. Light blue oval: domain rich in CIA, poor in ACP and OA, tentatively associated with an unmelt core of a splat. White circle: domain rich in OA, poor in HA, within two ACP layers, attributed to the recrystallization of a molten domain or of solid ACP phase. The layered structure of the splats is clearly visible on ACP maps (the top surface is corrupted by the spectra of the embedding resin).
Tensile Strength (ASTM C633-01, ISO 13779-4)

Test samples were made on 25 mm disks coated with HA or CIA (12 spray runs). These coated disks were glued to a dummy cylinder with the same diameter with FM1000 glue, (Cytec, NJ, USA) applied at 200°C for 90 to 100 minutes. The ultimate tensile strength of the assembly was determined on an Instron instrument, without pre-loading at a traction rate of 1mm/minute. The data obtained (mean of six measurements) are presented in Table 6. There was no significant difference between CIA and HA coatings. The failure appeared mainly cohesive for HA coupons (it occurred at the interface between the coating and the substrate), and it was mixed cohesive (at 80%) and adhesive for CIA coupons.

Table 6. Tensile strength of the coatings obtained at 12 spray runs

<table>
<thead>
<tr>
<th>Coating</th>
<th>Tensile bond strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HA</td>
<td>8.9 ± 1.5</td>
</tr>
<tr>
<td>CIA</td>
<td>8.2 ± 2.2</td>
</tr>
</tbody>
</table>

The ultimate loads observed are in the lower range of those reported for HA coatings obtained by plasma spraying [65]. The main drawback of this basic standardized test is the diffusion of the glue in the porous coating and its limitation to rather thick coatings (above 0.38 mm). Other measurements were performed especially the three point bending with stiffener.

Three Point Bending Test with Stiffener

This test was developed particularly by A. Roche et al. [67] and lead to the ISO 14679-1997 standard. It is designed to determine adhesive or cohesive properties of paints, varnish and others coatings. A stiffener epoxy resin (Araldite® AW134B with HV997 hardener) is cured onto the coated substrate in accordance with the standard to generate a stress concentration at the interface substrate-deposit. Such loading is known to produce mainly shear stress at the substrate/coating interface and appears more consistent with the surgical use of coated prosthesis. The determination of the critical shear stress of the interface requires, without considerations to the actual geometry of the sample, to know the mechanical properties of all the materials implied in the elaboration of the samples: substrate, coating, and bulk stiffener Young’s modulus, as well as the thickness and elastic properties of a possible interphase between the coating and the stiffener. Once these parameters are determined, critical shear stress and critical strain energy release rate of the substrate coating interface can be determined from experimentally measured critical load, thanks to analytical and/or numerical models [68]. This complexity led us, as many authors [69, 70] to consider this practical adhesion test as a comparative test, considering critical load $F_{\text{max}}$ an indicator of adhesion.

The tests were performed on corundum-blasted titanium coupons plasma-sprayed at 12 spray runs. The stiffener was casted in a silicon mould, directly on coated samples. The mechanical test apparatus used in this experiment consists in an electromechanical Instron test machine fitted with a load cell capacity of 500 N. The crosshead speed is set to 0.5 mm. min$^{-1}$. A schematic illustration of the testing conditions is given as supportive material. Ten samples were successfully tested for CIA coatings, only eight for HA and two of them had the rupture of the stiffener itself. Load versus displacement curves are displayed on Fig. (15). To check the quality of the stiffener, we measured the stiffness of the sample. It was found to be $319 \pm 16.9$ N.mm$^{-1}$ for HA coatings and $305 \pm 9.83$ N.mm$^{-1}$ for CIA coatings. These preliminary results confirmed the good quality of the polymeric stiffener.

An abrupt decrease in load-displacement behaviour was observed beyond $F_{\text{max}}$. The maximum load $F_{\text{max}}$ achieved before rupture of the substrate-coating interface, thanks to the geometrical effect generated by the stiffener, is the indicator of the practical adhesion. For HA, $F_{\text{max}} = 165 \pm 30.7$ N (18.6 %), and for CIA, $F_{\text{max}} = 128 \pm 7.92$ N (6.16 %). Coatings exhibited mixed modes of failure (cohesive and adhesive). By image analysis on Energy-dispersive X-ray spectroscopy (EDX) maps, adhesive failure was estimated to be closed to 65% for both HA and CIA deposits. (EDX maps are given as supportive material)

DISCUSSION

The data reveal some differences between HA and CIA coatings obtained in similar deposition conditions with powders showing slightly different characteristics:

![Fig. (15), three point bending tests loading curves for HA (left) and CIA (right) coatings.](image-url)
Formation of oxyapatite occurs in both coatings but CIA coatings exhibit much lower OA contents than HA.

ACP content is also lower in CIA coatings compared to HA ones.

Thickness of the coatings for the same number of spray runs is higher for HA than for CIA.

Surface roughness is higher in HA than in CIA.

Mechanical properties appear also slightly different, showing lower strength in CIA than in HA coatings in a three point bend test with stiffener, although adherence tests according to the ISO standard 13779-4 did not reveal significant differences.

HA and CIA Powder Characteristics

The different physical characteristics of the HA and CIA coatings, for identical spray conditions, could be related to slight changes in the powder characteristics and specific differences in the apatites involved. The chlorination treatment was expected to preserve the crystal size and grain characteristics. Actually, the reverse treatment (the hydroxylation of CIA) on monocrystals allowed the preparation of HA monocrystals and its crystal structure determination [71], this ability of high temperature ion exchange reactions to preserve the crystalline structure having been recently confirmed [72]. This is not observed in our chlorination experiments and a significant decrease of the crystal size of the initial HA was observed during its transformation into CIA. The role of the change in unit-cell volume between HA and CIA could be involved, although it represents only 3.6 % of the total volume. It shall be noticed, however, that the change in unit cell dimensions between HA and CIA involves a decrease of the c dimension and an increase of the a dimension. These opposite variations in partly sintered grains could lead to a rupture of the initial crystals and eventually of the grains. The reduction of roughness in CIA coatings compared to HA ones, even for one spray run, indicates a better spreading of the splats, possibly due to a lesser amount of matter and/or higher content of liquid phase and/or a lower viscosity. The slightly lower melting point of CIA (1530°C) compared to that of HA (1570°C), according to phase diagrams, together with a smaller grain and crystal size, could facilitate its melting within sprayed grains, although other considerations on heat transfers in the plasma flame have also to be taken into account [73].

Oxyapatite Content

The strong differences between Raman spectra of HA and OA have been assigned to variations of the local environments of phosphate ions, i.e. OH ions in HA and lattice vacancies or O\(^2\) ions in OA. Involving a, correlative alterations of the vibrational energy level of the phosphate ion in the apatite lattice. The formation of OA in HA coatings is generally attributed to the dehydration of HA during heating. Actually, OA is not considered to exist as a separate phase but as a solid solution, OHA (oxyhydroxyapatite). Raman spectra do not allow to discriminate easily between solid solution of OHA and mixtures of HA and OA phases. However, Raman mapping reveals micro-domains containing both HA and OA environments, thus supporting the existence of solid solutions. In addition, very strong spatial variations are evidenced indicating a strong heterogeneity. The formation of OA environments appears as an intermediate stage before the decomposition of HA into TTCP and TCP takes place. A second process of formation of OA environments is due to the recrystallization of apatite in a melt practically deprived of OH ions. The existence of domains very rich in OA and very poor in HA are consistent with this recrystallization process.

The formation of OA in CIA coatings is rather puzzling as there is no precursor of oxygennated species. Thus in CIA coatings the formation of OA involves necessarily a reaction with a contaminant. A reaction may be proposed with residual water molecules remaining in the CIA powder, which was stored and manipulated in air. In fact, this is the reverse reaction of the chlorination synthesis, which has been used to synthesize hydroxyapatite monocrystals:

\[
2\text{H}_2\text{O} + \text{Ca}_{10}((\text{PO}_4)_6\text{Cl})_2 \leftrightarrow \text{Ca}_{10}((\text{PO}_4)_6\text{OH})_2 + \text{HCl} \leftrightarrow \text{Ca}_{10}((\text{PO}_4)_6\text{O}) + \text{H}_2\text{O} + \text{HCl}
\]

Such a reaction involves the formation of HA as an intermediate phase. Although there is no detection of OH ions in the CIA coatings by Raman or FTIR spectroscopy, this reaction could occur in the plasma or on the surface of the coupon during cooling in air.

A second possible reaction would be the direct oxidation of CIA with O\(_2\) from air:

\[
\text{O}_2 + \text{Ca}_{10}((\text{PO}_4)_6\text{Cl})_2 \leftrightarrow \text{Ca}_{10}((\text{PO}_4)_6\text{O}) + \text{Cl}_2
\]

although this analogue of Deacon’s reaction [74] has never been reported in CIA. This reaction would occur preferably at the surface of splats during the cooling stage, when in contact with the air.

Other reactions might be possible between TiO\(_2\) at the surface of the substrate and CIA. However, OA is distributed in the coating thickness and does not localize especially in close proximity to the titanium surface. OA rich domains are always observed in domains isolated from CIA rich ones, with no clear gradation, suggesting a preferential formation of OA in a recrystallization process from the molten phase, which would facilitate oxygen diffusion and reaction. The mode of formation of OA in CIA coatings, which implies reactions with contaminants, explains why these contain significantly less OA than the HA ones, where OA formation is a part of the thermal degradation.

Amorphous Phase Content

Concerning the formation of ACP, the Raman spectroscopic data appear more reliable than the crystallinity measurements done by XRD following standards. As already mentioned before, several causes may be behind the decrease in XRD crystallinity observed in the coatings, of which ACP formation is only one. On the contrary, the significant shift in the Raman v\(_1\) PO\(_4\) of ACP, compared to that of apatite HA or CIA (-12 cm\(^{-1}\)), appears as a distinctive characteristic of the amorphous phase. However the broadness of the amorphous PO\(_4\) lines in Raman spectra and the superimposition with an OA line, do not allow a very...
accurate quantitative evaluation. The formation of ACP is assigned to the rapid cooling of the molten grain reaching the surface. However, a recrystallization process is supposed to either happen during the cooling of the splat or during an additional upper layer spray run, involving a local re-heating of an underlying amorphous phase. Thus, a difference in ACP content could be related to different ratios of molten phases and/or different rates of recrystallization. All these phenomena are related to heat transfer in the plasma or in the coating, which are difficult to evaluate and control, although simulations have been proposed [73]. Several spray parameters such as the angle of spraying; the spray distance and the plasma gas flow rate have been shown to determine the amorphous content of coatings [25]. However, these parameters were maintained constant in this study. A crucial parameter seems to be the particle size, which is related to strong amorphous phase variations [25]. The characteristics of CIA powders compared to HA ones (lower melting point, smaller grain size and crystal size) should favour the formation of coatings with a higher content in amorphous phase, which is not observed. Therefore, the recrystallization rate could be a determining factor in the ACP content in these experiments.

Different recrystallization processes may occur in coatings: crystal formation directly from the melt or in solid state, from the ACP phase. The recrystallization in the melt is expected to be a faster process, although it can be interrupted by the rapid cooling and solidification of the splat. Recrystallization from the melt could be easier in the case of CIA as it does not involve intermediate phases like those theoretically observed in HA recrystallization according to phase diagrams. The recrystallization from the amorphous solidified phase would then follow and may occur at a rather low temperature. In HA coatings for example, the recrystallization of ACP has been observed at about 700°C [75]. The low recrystallization temperature could also occur when a new hot layer is deposited on a cold one, due to the low thermal conductivity of apatites.

Mechanical Properties

The difference in mechanical properties between CIA and HA coatings is especially apparent in the three point bending test with stiffener, which seems more reliable than the standardized test. The adhesive strength of coatings has been correlated to the porosity of the coatings, the presence of cracks related to stresses due to differences in thermal coefficient expansion, and its composition. In wet media, rehydration of the coating, inducing cracks, has been identified as a crucial parameter in the degradation of mechanical properties. Chemical interactions between the coating and the oxide layer of the substrate have been suggested, but their role in the adhesion has not been clearly identified. Concerning the phase content, it appears that a high amount of ACP, as well as a high amount of crystalline phase, is not favourable [76]. The weaker strength of CIA coatings could thus be related to too high a crystallinity.

CONCLUSION

The main characteristics of CIA compared to HA coatings is a higher content in apatite and a lower content in amorphous calcium phosphate and oxyapatite. It is probable that slight modifications in the coating process would permit to limit even further the formation of OA in CIA coatings, especially a careful elimination of water from the CIA powder, although the role of OA in coatings is not known and this phase should not behave differently from other apatites. The expected advantage of CIA coatings would be a lesser degradation in vivo. A persisting problem remains the poor adhesion of the CIA coating on titanium substrates. In vivo experiments have been realized and the explants are currently examined. The first results show a direct attachment of bone to the coating surface, as it is usually found for this kind of coating. These results will be published in a forthcoming report.

SUPPLEMENTARY MATERIALS

Supplementary material is available on the publisher’s web site along with the published article.

CONFLICT OF INTEREST

The authors confirm that this article content has no conflict of interest.

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