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Link to publication record in Ulster University Research Portal

Published in:
Ceramics International

Publication Status:
Published (in print/issue): 31/05/2018

DOI:
10.1016/j.ceramint.2018.01.206

Document Version
Author Accepted version

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Synthesis and characterisation of nanophase hydroxyapatite co-substituted with strontium and zinc

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ARTICLE INFO

Keywords:
Bioerodible
Nano-hydroxyapatite
Co-substitution
Strontium
Zinc

ABSTRACT

In order to develop new bioactive calcium phosphate (CaP) materials to repair bone defects, it is important to ensure these materials more closely mimic the non-stoichiometric nature of biological hydroxyapatite (HA). Typically, biological HA combines various CaP phases with different impurity ions, which substitute within the HA lattice, including strontium (Sr$^{2+}$), zinc (Zn$^{2+}$), magnesium (Mg$^{2+}$), carbonate (CO$_3^{2-}$) and fluoride (F$^-$), but to name a few. In addition to this biological HA have dimensions in the nanometre (nm) range, usually 60 nm in length by 5–20 nm wide. Both the effects of ion substitution and the nano-size crystals are seen as important factors for enhancing their potential biofunctionality. The driving hypothesis was to successfully synthesise nanoscale hydroxyapatite (nHA), co-substituted with strontium (Sr$^{2+}$) and zinc (Zn$^{2+}$) ions in varying concentrations using an aqueous precipitation method and to understand their chemical and physical properties. The materials were characterised using Fourier Transform Infrared Spectroscopy (FTIR), X-Ray Diffraction (XRD), X-Ray Photoelectron Spectroscopy (XPS) and Transmission Electron Microscopy (TEM) techniques. The FTIR results confirmed that all of the samples were carbonated, with a significant loss of hydroxylation as a consequence of the incorporation of Sr$^{2+}$ and Zn$^{2+}$ into the nHA lattice. The TEM results showed that each sample produced was nano-sized, with the Sr/Zn-10% nHA having the smallest sized crystals approximately 17.6 ± 3.3 nm long and 10.2 ± 1.4 nm wide. None of the materials synthesised here in this study contained any other impurity CaP phases. Therefore, this study has shown that co-substituted nHA can be prepared, and that the degree of substitution (and the substituting ion) can have a profound effect on the attendant materials’ properties.

1. Introduction

Synthetic hydroxyapatite (HA) [$\text{Ca}_{10}(\text{PO}_4)_6(\text{OH})_2$] has been used extensively as a bone augmentation biomaterial, producing scaffolds for bone tissue engineering and for coating implants to promote bone healing and more effectively enhancing osseointegration [1,2]. In biological HA, the non-stoichiometric nature of biological hydroxyapatite (HA) is thought to be due to impurities within the HA structure, such as magnesium (Mg$^{2+}$), carbonate (CO$_3^{2-}$) and fluoride (F$^-$), and the presence of Sr$^{2+}$ and Zn$^{2+}$ ions. The nano-size crystals are seen as important factors for enhancing their potential biofunctionality. The driving hypothesis was to successfully synthesise nanoscale hydroxyapatite (nHA), co-substituted with strontium (Sr$^{2+}$) and zinc (Zn$^{2+}$) ions in varying concentrations using an aqueous precipitation method and to understand their chemical and physical properties. The materials were characterised using Fourier Transform Infrared Spectroscopy (FTIR), X-Ray Diffraction (XRD), X-Ray Photoelectron Spectroscopy (XPS) and Transmission Electron Microscopy (TEM) techniques. The TEM results showed that each sample produced was nano-sized, with the Sr/Zn-10% nHA having the smallest sized crystals approximately 17.6 ± 3.3 nm long and 10.2 ± 1.4 nm wide. None of the materials synthesised here in this study contained any other impurity CaP phases. Therefore, this study has shown that co-substituted nHA can be prepared, and that the degree of substitution (and the substituting ion) can have a profound effect on the attendant materials’ properties.

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https://doi.org/10.1016/j.ceramint.2018.01.206
Received 20 November 2017; Received in revised form 19 January 2018; Accepted 24 January 2018

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Please cite this article as: Lowry, N., Ceramics International (2018), https://doi.org/10.1016/j.ceramint.2018.01.206
Table 1
Examples of various ions which can substitute within the HA lattice.

<table>
<thead>
<tr>
<th>Substituted ion</th>
<th>Position of substitution</th>
<th>Ionic radius (Å) Ca (0.99Å)</th>
<th>Biological function and effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strontium (Sr²⁺)</td>
<td>Ca²⁺</td>
<td>1.12</td>
<td>High concentration where new bone is forming. Boosts osteoblast (OB) cell proliferation. Reduces osteoclast (OC) activity and therefore rate of bone resorption.</td>
</tr>
<tr>
<td>Ca(I) High Concentrations</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Zinc (Zn²⁺)</td>
<td>Ca²⁺</td>
<td>0.74</td>
<td>Inhibits OC resorption of bone. Encourages OB activity stimulating new bone formation.</td>
</tr>
<tr>
<td>Ca(II) High Concentrations</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Magnesium (Mg²⁺)</td>
<td>Ca²⁺</td>
<td>0.72</td>
<td>Assists calcium ions to promote bone strength and health.</td>
</tr>
<tr>
<td>Ca(II) High Concentrations</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Silver (Ag⁺)</td>
<td>Ca²⁺</td>
<td>1.26</td>
<td>Exhibits an antibacterial effect on <em>S</em>aphylococcus <em>aureus</em>, <em>E</em> coli, <em>Pseudomonas aeruginosa</em>, <em>Candida albicans</em> and gram negative bacilli, with relatively low cytotoxicity. A minimum inhibitory concentration (MIC) of silver must be reached and maintained to inhibit later stage bacterial colonisation</td>
</tr>
<tr>
<td>Lithium (Li⁺)</td>
<td>Ca²⁺</td>
<td>0.76</td>
<td>Causes a reduction in bone remodelling, reducing bone turnover rates with lower amounts of bone loss. Stimulates the canonical Wnt signalling pathway, activating a major osteogenic signalling cascade, allowing MSC's to differentiate to OBs.</td>
</tr>
<tr>
<td>Copper (Cu²⁺)</td>
<td>Ca²⁺</td>
<td>0.73</td>
<td>Antimicrobial effect against <em>E</em> coli and <em>Candida albicans</em>.</td>
</tr>
<tr>
<td>Silicon (SiO₄²⁻)</td>
<td>PO₄³⁻</td>
<td>5.86</td>
<td>Increases solubility and dissolution.</td>
</tr>
<tr>
<td>Carbonate (CO₃²⁻)</td>
<td>CO₃²⁻</td>
<td>4.35</td>
<td>A type substitutes for hydroxyl ions B type substitutes for phosphate ions (A/B ratio = 0.7-0.9) Crystal morphology evolves as level of carbonate substitution increases from plate-like to needle-like to spherical. Reduced grain size and hardness value 200% larger than HA values, but not suitable for major load bearing applications. Fast dissolution times, in vivo and in vitro due to carbonate weakening the bonds within the HA lattice. Bioactive and osseointegrative, forming a bone-like surface apatite layer in 7 days compared to 24-28 days for stoichiometric HA</td>
</tr>
<tr>
<td>Fluoride (F⁻)</td>
<td>OH⁻</td>
<td>1.33</td>
<td>Substitutes for hydroxyl groups</td>
</tr>
</tbody>
</table>

References:

[5,6,8,9,32-34].

[6-8,30,31,35-39].

[8,40-42].

[8,43].

[34,44-47].

[48,49]

[8,30,50-52].

[8,16,53].

[8,54].
characterised and compared to the stoichiometric nHA using X-ray Diffraction (XRD), Fourier Transform Infrared Spectroscopy (FTIR), X-ray Photoelectron Spectroscopy (XPS) and Transmission Electron Microscopy (TEM). To the best of our knowledge, this paper represents the first attempts to produce, systematically characterise andphysicochemical properties of nanohydroxyapatite, co-substituted to these various Sr$^{2+}$/Zn$^{2+}$ combined weight percentages, produced under identical conditions of pH and temperature.

2. Materials and methods

The nomenclature used to describe each of the calcium phosphate (CaP) powders synthesised in this study (using a standard aqueous precipitation method) are outlined in Table 2. Identical experimental conditions of temperature (60 °C) and pH ≥ 9, were used to synthesise pure nanoscale hydroxyapatite (nHA), nanoscale Sr(1.25%)/Zn(1.25%) hydroxyapatite (Sr/Zn-2.5%nHA), nanoscale Sr(2.5%)/Zn(2.5%) hydroxyapatite (Sr/Zn-5%nHA) and nanoscale Sr(5%)/Zn(5%) hydroxyapatite (Sr/Zn-10%nHA). Sodium hydroxide was used as a buffer to raise the pH ≥ 9 for the co-substitutions, Sr/Zn-2.5%nHA, Sr/Zn-5%nHA and Sr/Zn-10%nHA. The experimental ratios of Ca/P and (Ca + M)/P, (where M = Sr or Zn) were maintained at 1.67, for final comparison.

2.1. Aqueous precipitation synthesis of nHA

nHA powders were synthesised using an acid-base reaction of calcium hydroxide (Ca(OH)$_2$ - Sigma Aldrich) and phosphoric acid (85 wt % H$_3$PO$_4$ - Sigma Aldrich). Ca(OH)$_2$ was suspended in 500 ml of deionised H$_2$O, heated to 60 °C and stirred at 400 rpm for 1 h. An aqueous solution of H$_3$PO$_4$ was prepared by dissolving 85 wt% H$_3$PO$_4$ in 500 ml of deionised H$_2$O. The H$_3$PO$_4$ solution was added to the Ca(OH)$_2$ solution at a rate of 3.5 ml/min using a peristaltic pump. The solution was stirred for a further 2 h and allowed to settle and cool overnight. The pH of the solution was measured at 9.1. The precipitate was washed 3 times using deionised H$_2$O and then dried at 60 °C in an oven for 24 h.

2.2. Aqueous precipitation synthesis of strontium and zinc co-substituted nHA

Starting suspensions were prepared by dissolving calcium nitrate tetrahydrate (Ca(NO$_3$)$_2$·4H$_2$O - puriss.pa Sigma Aldrich) with both strontium nitrate (Sr(NO$_3$)$_2$·4H$_2$O - puriss pa. Sigma Aldrich) and zinc nitrate hexahydrate (Zn(NO$_3$)$_2$·6H$_2$O - Sigma Aldrich) in 500 ml of deionised H$_2$O. The solution was stirred for 1 h at 60 °C and 400 rpm. An aqueous solution of H$_3$PO$_4$ was prepared by dissolving 85 wt% H$_3$PO$_4$ in 500 ml of deionised H$_2$O. The H$_3$PO$_4$ solution was added to the Ca(NO$_3$)$_2$·4H$_2$O solution at a rate of 3.5 ml/min using a peristaltic pump. The pH of the solution was typically measured between 2.0 and 2.5, with the solutions remaining colourless. On addition of the sodium hydroxide to buffer each co-substituted solution to a pH ≥ 10, a white precipitate formed immediately. The solution was stirred for a further 2 h, with the pH maintained at ≥ 10 (by further addition of sodium hydroxide). The solution was allowed to settle and cool to room temperature overnight. The formed precipitate was washed 3 times using deionised H$_2$O and then dried at 60 °C in an oven for 24 h.

Table 2

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Abbreviation used</th>
<th>Chemical formula</th>
<th>Molar quantity (mmol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Synthesised Nanoscale Hydroxyapatite</td>
<td>nHA</td>
<td>Ca$_{10}$(PO$_4$)$_6$(OH)$_2$</td>
<td>50.000</td>
</tr>
<tr>
<td>Synthesised Nanoscale Sr/Zn co- substituted Hydroxyapatite (1.25%/1.25%)</td>
<td>Sr/Zn-2.5%nHA</td>
<td>Ca$<em>{9.75}$Sr$</em>{0.125}$Zn$_{0.125}$(PO$_4$)$_6$(OH)$_2$</td>
<td>45.000</td>
</tr>
<tr>
<td>Synthesised Nanoscale Sr/Zn co- substituted Hydroxyapatite (2.5%/2.5%)</td>
<td>Sr/Zn-5%nHA</td>
<td>Ca$<em>{9.5}$Sr$</em>{0.25}$Zn$_{0.25}$(PO$_4$)$_6$(OH)$_2$</td>
<td>45.000</td>
</tr>
<tr>
<td>Synthesised Nanoscale Sr/Zn co- substituted Hydroxyapatite (5%/5%)</td>
<td>Sr/Zn-10%nHA</td>
<td>Ca$<em>{9}$Sr$</em>{0.5}$Zn$_{0.5}$(PO$_4$)$_6$(OH)$_2$</td>
<td>45.000</td>
</tr>
</tbody>
</table>

2.3. Characterisation of the nHA and co-substituted Sr/Zn-nHA powders

X-ray Diffraction (XRD) of the samples was carried out using a Bruker D8 Discover Diffractometer fitted with a Gobel mirror. Diffraction scans were obtained using a Cu Ka X-ray radiation (λ = 1.540 Å source, with a tube voltage of 40 kV and tube current of 40 mA. Each diffraction scan was recorded at 20 values from 5° to 70°, in 0.04° stepped increments, with a scan dwell time of 10 s for each increment.

Fourier Transform Infrared (FTIR) spectroscopy scans were obtained for each sample using a BIORAD FTS 3000MX Excalibur series instrument with a PIKE Diffuse Reflectance Infrared Fourier Transform Spectroscopy (DRIFTS) accessory. Samples were studied in absorbance mode from 4000 to 400 cm$^{-1}$, at a resolution of 4 cm$^{-1}$, with 20 scans per sample.

X-ray Photoelectron Spectroscopy (XPS) was performed using a Kratos Axis Ultra DLD Spectrometer. Spectra were analysed using monochromated Al Ka X-rays (hv = 1486.6 eV (eV)) operating at 10 kV and 15 mA (150 W). During analysis, a hybrid lens mode was used (electrostatic and magnetic) with a 300 µm × 700 µm analysis area. Wide energy survey scans (WESS) were taken at a pass energy of 200 eV, with high resolution spectra recorded at a pass energy of 20 eV. A Kratos charge neutraliser system with a 300 µm × 700 µm analysis area. The statistical analysis for the TEM data was performed with ImageJ software was utilised to measure the dimensions of 30 nanocrystals from each sample across 6 different TEM images. The peak area of the most intense spectral lines for each elemental species was used to determine the percentage atomic concentration.

Transmission Electron Microscopy (TEM) observations were carried out using a JEOL 2100f microscope operated at 200 kV. Bright-field images were recorded from the powders supported on carbon films for size analysis. ImageJ software was utilised to measure the dimensions of 30 nanocrystals from each sample across 6 different TEM images. The results are reported as the mean ± standard deviation.

2.4. Statistical analysis

The statistical analysis for the TEM data was performed with GraphPad Prism version 5.00 for Windows, GraphPad Software, San Diego California USA. One way analysis of variance (ANOVA) was used
to compare mean values in order to determine equivalence of variance between pairs of samples. Significance between groups was determined using the Bonferroni multiple comparison test. A value of P < .05 was taken as statistically significant.

3. Results

3.1. XRD

The comparative XRD patterns for the nHA, Sr/Zn2.5%nHA, Sr/Zn5%nHA and Sr/Zn10%nHA are shown in Fig. 1(a)–(d). The diffraction pattern for the nHA material is given in Fig. 1(a) and closely correlates to the International Centre for Diffraction Data (ICDD) file # 00-09-0432 for hydroxyapatite (HA). The peak positions and intensities recorded clearly indicate that the nHA material contains no detectable impurity phases. Fig. 1(b)–(d) for the Sr/Zn2.5%nHA, Sr/Zn5%nHA and Sr/Zn10%nHA samples, respectively, show diffraction peaks that are poorly resolved, indicating that each of the synthesised materials are nanoscale (≤ 100 nm), and/or have appreciable levels of substitutions within the nHA lattice, whereby Sr2+ and Zn2+ are substituting for Ca2+. All 3 substituted nHA samples, Sr/Zn2.5%nHA, Sr/Zn5%nHA and Sr/Zn10%nHA exhibit noticeable shifts in XRD peak positions to lesser 2θ values, causing individual peak movement to the left, this is highlighted in Fig. 1 with movement of the 002 peak highlighted.

3.2. FTIR

FTIR spectra of the synthesised nHA and substituted nHA powders are shown in Fig. 2(a)–(d), respectively. Typical P-O stretching vibrations are clearly seen between 1200 and 900 cm⁻¹, while O-P-O bending vibrations can be observed between 610 and 400 cm⁻¹ for each of the different materials [13]. O-H stretching and librational bands are also clearly visible around 3570 cm⁻¹ and 632 cm⁻¹, respectively [14,15]. In addition to these vibrational bands, carbonate peaks (CO3²⁻) can also be observed between 1550 and 1400 cm⁻¹ and around 870 cm⁻¹ [16]. The carbonate peak positions may indicate that these different materials (both nHA and substituted nHA) have CO₃²⁻ ion substitutions at both the OH (A-site) and PO₄³⁻ (B-site) positions [13].

3.3. XPS

Figs. 3–6 show typical XPS wide energy survey scans (WESS) and high resolution scans recorded as B.E. (e.V.) versus intensity (counts), respectively, for the nHA, Sr/Zn-2.5%nHA, Sr/Zn-5%nHA and Sr/Zn10%nHA powders (namely the Ca2p, P2s, Sr3p and Zn2p peaks for each sample). Tables 3 and 4 quantify the XPS results recorded here. The main peaks recorded for the nHA sample were Ca2p3/2 (347.2 eV), P2p (133.7 eV), P2s (190.9 eV), and O1s (531.6 eV) are all clearly evident. The peak positions are as expected from previous reports, however, the Ca/P ratio was slightly lower than the expected 1.67 for stoichiometric HA, at 1.61 ± 0.04 as highlighted in Table 3[13,15,17,18]. For all 3 substituted samples, the Ca/P ratios were calculated at 1.53 ± 0.06 for Sr/Zn-2.5%nHA, 1.42 ± 0.03 for Sr/Zn-5%nHA and 1.38 ± 0.01 for Sr/Zn-10%nHA, all of which are lower than 1.67 for stoichiometric HA. However, this is as expected and would be a consequence of the substitution of strontium and zinc for calcium within the nHA lattice[19–21].

As all 3 co-substituted samples contain strontium, it should be noted that the spectral envelope observed around 133.5 eV comprises an overlap of the Sr3d₅/₂ (133 ± 0.5 eV), Sr3d₃/₂ (135 ± 0.5 eV) and the P2p (132–133 eV) lines are positioned very close together. Therefore, to quantify the phosphorus and strontium atomic concentration % values in this study for all 3 co-substituted samples, the P2s and Sr3p peaks are used instead of the P2p and Sr3d peaks. The main peaks recorded for all 3 co-substituted samples, shown in Table 4 were therefore Ca2p3/2 (346.5–347.0 eV), P2s (190.0–190.3 eV), O1s (530.5–531.0 eV), Sr3p3/2 (268.3–268.8 eV), Zn2p3/2 (1021.7–1022.0 eV) and Na1s (1071.0–1071.5 eV). The Sr3p3/2 peaks confirm the presence and successful substitution of strontium for calcium within each co-substituted sample and the Zn2p3/2 peaks confirm the presence and successful substitution of zinc for calcium within each co-substituted sample.

3.4. TEM

The TEM images shown in Fig. 7 confirm that the nHA, Sr/Zn-2.5%nHA, Sr/Zn-5%nHA and Sr/Zn-10%nHA samples produced were all on the nanoscale, with the nHA crystals measuring 99.5 ± 12.9 nm in length and 33.8 ± 7.8 nm in breadth, with an aspect ratio of...
Fig. 2. FTIR spectra for nHA and co-substituted nHA samples.

Fig. 3. XPS spectra for nHA sample.

Fig. 4. XPS spectra for Sr/Zn2.5%nHA sample.
3.09 ± 0.75, as reported in Table 5. The Sr/Zn-2.5%nHA, Sr/Zn-5%nHA and Sr/Zn-10%nHA samples all have nanocrystals, which are much smaller in size (both in length and breadth) than the nHA powder, as reported in Table 5, with typically lengths < 40 nm and breadths < 11 nm. The analysis of the TEM results, as highlighted in Fig. 8, show that there is a statistical difference in the length and breadth of the nHA, when compared with the co-substituted materials. In addition, within the co-substituted materials significant differences are observed in the length of the crystals between all samples. With respect to the aspect ratio, there was a significant difference between the nHA and the Sr/Zn-5%nHA and Sr/Zn-10%nHA samples (and between all of the different co-substituted materials). However, no significant statistical difference was observed in the aspect ratio between the nHA and the Sr/Zn-2.5%nHA sample.

4. Discussion

4.1. XRD

The XRD diffraction patterns for all 3 co-substituted samples, Sr/Zn2.5%nHA, Sr/Zn5%nHA and Sr/Zn10%nHA in Fig. 1(b)–(d), show definite peak shifts to the left indicating that the diffraction peaks have shifted to slightly lower 2 theta (2θ) values, in comparison to the nHA sample. This is due to Sr2+ ions with a larger ionic radius (1.12Å) replacing Ca2+ ions with a smaller ionic radius (0.99Å). It can also be observed in the 002 peaks in Fig. 1 that as the wt% substitution of

<table>
<thead>
<tr>
<th>Sample</th>
<th>Ca/P</th>
<th>(Ca + Sr)/P</th>
<th>(Ca + Zn)/P</th>
<th>(Ca + Sr + Zn)/P</th>
<th>Sr/(Sr + Ca)</th>
<th>Zn/(Zn + Ca)</th>
<th>Sr + Zn/(Sr + Zn + Ca)</th>
</tr>
</thead>
<tbody>
<tr>
<td>nHA</td>
<td>1.61 ± 0.04</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>0.04 ± 0.01</td>
<td>0.02 ± 0.01</td>
<td>0.06 ± 0.01</td>
</tr>
<tr>
<td>Sr/Zn-2.5%nHA</td>
<td>1.53 ± 0.06</td>
<td>1.50 ± 0.04</td>
<td>1.47 ± 0.04</td>
<td>1.54 ± 0.05</td>
<td>0.05 ± 0.01</td>
<td>0.03 ± 0.01</td>
<td>0.08 ± 0.01</td>
</tr>
<tr>
<td>Sr/Zn-5%nHA</td>
<td>1.55 ± 0.01</td>
<td>1.45 ± 0.04</td>
<td>1.63 ± 0.04</td>
<td>0.11 ± 0.01</td>
<td>0.05 ± 0.01</td>
<td>0.05 ± 0.01</td>
<td>0.15 ± 0.02</td>
</tr>
<tr>
<td>Sr/Zn-10%nHA</td>
<td>1.55 ± 0.01</td>
<td>1.55 ± 0.01</td>
<td>1.56 ± 0.06</td>
<td>1.63 ± 0.05</td>
<td>0.04 ± 0.01</td>
<td>0.02 ± 0.01</td>
<td>0.06 ± 0.01</td>
</tr>
</tbody>
</table>

Fig. 5. XPS spectra for Sr/Zn5%nHA sample.

Fig. 6. XPS spectra for Sr/Zn10%nHA sample.

Table 3

Comparative XPS ratios for standard nHA and substituted nHA powders (± standard deviation).
strontium increases, from Fig. 1(b)–(d), so does the level of shifting to the left and lower 2θ values increase, suggesting that gradually larger amounts of strontium have been successfully substituted into the nHA with each sample. The peaks also exhibited significant broadening with increasing Sr2+ content as can be seen in Fig. 1(b)–(d). Again, this result would be expected as increased Sr2+ substitution increases the strain within the lattice [3,22,23]. The XRD peak broadenings found in the 3 co-substituted samples may also indicate the successful substitution of Zn2+ ions for Ca2+ ions within the nHA lattice. However, peak positions would be expected to shift very slightly to higher 2θ values, indicating the substitution of the smaller Zn2+ ion for the Ca2+ ion within the HA crystal lattice, causing a shrinkage in unit cell parameters. A peak shift to the right and higher 2θ values cannot be observed in Fig. 1 but this could be due to the shift to the left and lower 2θ values caused by the Sr2+ substitution within the same samples, masking any subsequent peak shift to the right caused by substituted Zn2+ ions. Previous research substituting HA with zinc alone has produced XRD diffraction patterns with these noticeable peak shifts to the right and higher 2θ values, which could be used to confirm along with XPS data that Zn2+ ions have been successfully substituted within the nHA lattice however the larger amounts of Sr2+ ions substituted within the nHA lattice conceal any peak shift or increase in 2θ value associated with Zn2+ substitution [10,24,25].

4.2. FTIR

The FTIR peak positions observed for the nHA material for the P-O stretching and O-P-O bending vibrations were similar to those previously known for HA, as were those observed for the O-H librational and stretching modes. No peaks indicative of HPO4$^{2-}$ functional groups were detected for the nHA material (typically peaks for HPO4$^{2-}$ are seen around 1120 and 580 cm$^{-1}$) [17,19].

The 3 co-substituted samples appear to have broader P-O stretching bands located around 1099, 1043 and 964, and the O-P-O bending bands at 603 and 565, which could be due to both Sr2+ and Zn2+ substituting for calcium in the nHA lattices. There is also increased CO3$^{2-}$ substitution, in all 3 co-substituted samples, which previous literature has highlighted as an expected outcome when Zn2+ substitutes for Ca2+ in the nHA lattice [25,26]. In addition to these results, O-H stretching bands located around 3570 cm$^{-1}$ and O-H librational bands around 632 cm$^{-1}$ reduce in intensity for all 3 co-substituted samples in comparison to the nHA sample. This de-hydroxylation supports the fact that substitution of Sr2+ and Zn2+ ions for Ca2+ disrupts the nHA lattice.

![Fig. 7. TEM images of nHA and co-substituted nHA samples.](image-url)
Table 5
Dimensions of nanocrystals (± standard deviation) measured using TEM analysis and processed using ImageJ software.

<table>
<thead>
<tr>
<th>Sample</th>
<th>nHA</th>
<th>Sr/Zn-2.5%-nHA</th>
<th>Sr/Zn-5%-nHA</th>
<th>Sr/Zn-10%-nHA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length (nm)</td>
<td>99.5 ± 12.9</td>
<td>31.4 ± 6.5</td>
<td>23.0 ± 3.0</td>
<td>17.6 ± 3.3</td>
</tr>
<tr>
<td>Breadth (nm)</td>
<td>33.8 ± 7.8</td>
<td>9.5 ± 1.5</td>
<td>10.0 ± 1.5</td>
<td>10.2 ± 1.4</td>
</tr>
<tr>
<td>Aspect ratio</td>
<td>3.09 ± 0.75</td>
<td>3.40 ± 0.96</td>
<td>2.36 ± 0.47</td>
<td>1.75 ± 0.33</td>
</tr>
</tbody>
</table>

4.3. XPS

The Ca/P ratio for the nHA sample was 1.61 ± 0.04, slightly lower than the expected 1.67 for stoichiometric HA. All 3 co-substituted samples had further reduced Ca/P ratios, 1.53 ± 0.06 for Sr/Zn-2.5% nHA, 1.42 ± 0.03 for Sr/Zn-5% nHA, and 1.38 ± 0.01 for Sr/Zn-10% nHA. This reduction in calcium content for each co-substituted sample would suggest successful substitution of strontium and zinc for calcium within the co-substituted nHA lattices, also as the wt% of strontium and zinc increases from 2.5%, 5% and 10%, the Ca/P ratio decreases, suggesting gradually more is substituted into the nHA, replacing calcium.

The (Ca + Sr)/P ratios for each co-substituted sample, found in Table 3 are as follows; 1.59 ± 0.05 for Sr/Zn-2.5%nHA, 1.50 ± 0.04 for Sr/Zn-5%nHA and 1.55 ± 0.01 for Sr/Zn-10%nHA. It can be seen that for each sample the ratio is larger for (Ca + Sr)/P in comparison to the Ca/P ratios alone, providing further evidence that for each co-substituted sample strontium has been successfully substituted for calcium within the nHA lattice. Again, the 3 (Ca + Zn)/P ratios for the co-substituted samples, Table 3 are as follows; 1.56 ± 0.06 for Sr/Zn-2.5%nHA, 1.47 ± 0.04 for Sr/Zn-5%nHA and 1.45 ± 0.04 for Sr/Zn-10%nHA. This would again confirm that zinc has successfully substituted for calcium within the nHA lattice as amounts of the starting strontium nitrate and zinc nitrate hexahydrate are increased.

Small amounts of sodium were also detected in the XPS spectra analyses, with the Na1s peak located between 1071.0 and 1071.5 eV for all 3 co-substituted samples. This was caused by the use of sodium hydroxide as a buffer to increase the pH above 10 during each precipitation reaction and has been found to encourage the substitution of zinc for calcium within the nHA lattice. It should be noted that XPS is a surface analytical technique and only penetrates the uppermost surface (1–10 µm) of each sample.

4.4. TEM

The TEM results show clearly that as the co-substituted wt% of strontium and zinc increased from 2.5, 5 to 10%, the average particle length for each sample reduced from 31.4 ± 6.5 nm, 23.0 ± 3.0 nm and 17.6 ± 3.3 nm, respectively with the breadth remaining relatively constant at 9.5 ± 1.5 nm, 10.0 ± 1.5 nm and 10.2 ± 1.4 nm, respectively. This corresponds with previous research which showed that nHA substituted with both strontium and zinc produced smaller sized particle samples [27,28]. It has also been reported in previous literature that as the concentration of the substituted ion rises, regardless of 2.5% to 5% for each co-substituted sample the subsequent percentage ratios of atomic concentration also increase accordingly, confirming that as a larger percentage concentration of strontium is added at the beginning of the reaction, a larger amount of strontium is substituted for calcium within the nHA lattice of each sample. A similar trend can be seen with the Zn/(Zn + Ca) ratios again located in Table 3, a clear increase in zinc substitution for each increasing wt% sample can be seen, for Sr/Zn-2.5%nHA, 0.02 ± 0.01, for Sr/Zn-5%nHA, 0.03 ± 0.01 and for Sr/Zn-10%nHA, 0.05 ± 0.01. This would again indicate that as the wt% concentration of zinc is increased from 1.25%, 2.5% to 5% for each co-substituted sample the subsequent percentage ratios of atomic concentration also increase, again confirming that as a larger percentage concentration of zinc is added at the beginning of each synthesis, a larger amount of zinc is substituted for calcium within the nHA lattice of each sample. Again, the Sr + Zn/(Sr + Zn + Ca) ratios for Sr/Zn-2.5%nHA, 0.06 ± 0.01, for Sr/Zn-5%nHA, 0.08 ± 0.01 and for Sr/Zn-10%nHA, 0.15 ± 0.02 agree with the strontium and zinc ratios and show a distinct rise in % atomic concentration, as the starting percentage concentrations of both strontium and zinc are increased. This indicates that a larger amount of both strontium and zinc has been co-substituted into the nHA lattice of each sample, as the starting concentrations of strontium and zinc increase from 1.25% to 2.5% to 5% for each ion. This suggests that the wet precipitation synthesis successfully substitutes larger concentrations of both strontium and zinc within the nHA lattice as amounts of the starting strontium nitrate and zinc nitrate hexahydrate are increased.
whether it is strontium or zinc, the subsequent substituted nHA crystals tend to agglomerate [27,28]. The statistical analysis of the crystal length, breadth and aspect ratio here, highlight that as the concentration of the co-substituted ions increase, it has an obvious effect on the growth of the crystals, (with higher ion substitutions inhibiting crystal growth).

This study has also found that for every sample analysed using TEM, the observed nanocrystals appeared granular in nature, which may indicate a high level of porosity and an increased surface area within both the nHA and substituted nHA materials synthesised. An optimised surface nano-topography could in turn increase adsorption of specific proteins and ultimately improve osteoblast cell adhesion and promote osteoblast proliferation and differentiation [29]. Previous research has also found that nano-sized HA, with a larger surface area, has increased sinterability and densification which can in turn improve its mechanical properties and fracture toughness [2,29].

5. Conclusion

The specific aim of this work was to synthesise nHA and nHA co-substituted with strontium and zinc at varying concentrations, Sr/Zn-2.5%nHA, Sr/Zn-5%nHA and Sr/Zn-10%nHA, using an aqueous precipitation technique. The results indicated that nHA and nHA co-substituted with strontium and zinc were successfully produced at varying concentrations. The nHA materials produced did not contain any other CaP impurity phases and were proven to be nano-sized in scale, as shown by XRD, FTIR, XPS and TEM analyses. Further comparison did indicate that crystallinity did vary as the amount of Sr2+ and Zn2+ increased within the co-substituted samples, with the degree of structural order decreasing as Ca2+ ions were progressively replaced with either larger Sr2+ or smaller Zn2+ ions with increasing dehydroxylation observed within the co-substituted samples, which further highlighted disruption of the HA lattice. Also, the particle morphology of the co-substituted nHA samples changed with increased co-substitution, with the particles steadily reducing in length.

Very few previous studies have investigated the synthesis of nHA materials co-substituted with Sr2+ and Zn2+. Therefore, the successful co-substitution of both strontium and zinc within nHA, as demonstrated successfully here, could provide a more bioactive bone substitute material. However, there is a need for further detailed in vitro and in vivo studies to confirm the efficacy of these co-substituted materials.

Acknowledgements

The authors would like to acknowledge the Ph.D. funding provided by the Department for the Economy (Northern Ireland) to support this work.

References

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