

Cite this: *Mater. Adv.*, 2022,
3, 1713

Potassium–carbonate co-substituted hydroxyapatite compositions: maximising the level of carbonate uptake for potential CO₂ utilisation options†

Duncan A. Nowicki,  ‡*^a Janet M. S. Skakle  ^{ab} and Iain R. Gibson  ^{ac}

CO₂ utilisation is a rapidly growing area of interest aimed at reducing the magnitude of anthropogenic greenhouse gas emissions. We report the synthesis of potassium–carbonate (K–CO₃) co-substituted hydroxyapatites with potassium and carbonate contents ranging from approximately 0.4–0.9 wt% and 3.4–13.0 wt% respectively via an aqueous precipitation reaction between calcium hydroxide, phosphoric acid and either potassium carbonate or potassium hydrogen–carbonate. The incorporated carbonate is situated on both hydroxyl and phosphate sites. A subsequent heat treatment in dry CO₂ at 600 °C allowed for a K–CO₃ co-substituted apatite containing approximately 16.9 wt% CO₃^{2–} to be prepared, amongst the largest carbonate contents that have been reported for such a material to date. Although this work shows that K–CO₃ co-substituted apatites with high levels of carbonate incorporation can be prepared using simple, room temperature, aqueous precipitation reactions with starting reagents unlikely to pose significant environmental risks, testing of these materials in prospective applications (such as solid fertilisers) is required before they can be considered a viable CO₂ utilisation option. A preliminary assessment of the effect of potassium/carbonate substitution on the solubility of the as-prepared compositions showed that increasing carbonate substitution increased the solubility.

Received 1st August 2021,
Accepted 17th December 2021

DOI: 10.1039/d1ma00676b

rsc.li/materials-advances

1. Introduction

The Intergovernmental Panel on Climate Change (IPCC) has projected that if current trends were to continue unabated, greenhouse gases such as CO₂ could become so concentrated in the atmosphere that by the year 2100 the global mean surface temperature of the Earth could be more than 4 °C warmer than it was during the pre-industrial era.¹ Therefore, immediate action must be taken to reduce the magnitude of anthropogenic greenhouse gas emissions.

One route for achieving this is through the capture of CO₂ from large point sources such as fossil-fuel power stations and the subsequent utilisation of this gas in the preparation of

functional materials; in addition to potentially permanently fixing the utilised CO₂, these products could also lead to a reduction in future CO₂ emissions by replacing current more 'CO₂-intensive' alternatives.² For example, CO₂ could be used as a feedstock in the production of several important chemicals such as methanol, urea and salicylic acid^{3,4} or reacted with magnesium-rich brines to synthesise inorganic carbonates such as nesquehonite (MgCO₃·3H₂O) which could find application as a 'green' construction material.⁵ Carbon dioxide could even be reacted with promoter species in the presence of catalysts to form a range of environmentally benign polymers like poly(ethylene carbonate) and poly(propylene carbonate) which could replace conventional plastics.^{6,7}

In a similar manner, CO₂ gas could be utilised in the synthesis of a single-phase potassium–carbonate co-substituted hydroxyapatite. As phosphorus and potassium play important roles in plant growth,^{26–28} we speculate that such K–CO₃ co-substituted apatite materials could prove useful as solid fertilisers whilst allowing for captured CO₂ to be sequestered either in the solid materials themselves or in the soil to which the apatites are applied. However, careful characterisation of the uptake of CO₂ into these materials must be carried out first, to understand *e.g.* the role of reactant chemistry as a source of CO₂, levels of CO₂ incorporation that may be feasible, and the

^a Department of Chemistry, University of Aberdeen, Meston Walk, Aberdeen AB24 3UE, UK. E-mail: dan1@st-andrews.ac.uk, j.skakle@abdn.ac.uk, i.r.gibson@abdn.ac.uk

^b Department of Physics, University of Aberdeen, Meston Walk, Aberdeen AB24 3UE, UK

^c Institute of Medical Sciences, University of Aberdeen, Foresterhill, Aberdeen AB25 2ZD, UK

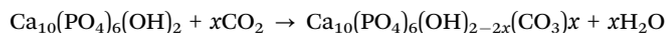
† Electronic supplementary information (ESI) available. See DOI: 10.1039/d1ma00676b

‡ Present address: School of Chemistry, University of St Andrews, North Haugh, St Andrews KY16 9ST, UK.

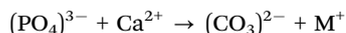


role of thermal treatment in a CO₂ atmosphere to incorporate additional carbon dioxide. Whilst not yet explored in great detail as a potential CO₂ utilisation option, hydroxyapatite has already proven to have application in several fields including aqueous solution remediation, gas detection, agriculture as a fertiliser, ionic conduction and catalysis,^{14–22} and has been studied as a solid sorbent for carbon capture, with a previous study showing stoichiometric hydroxyapatite can adsorb approximately 0.06–0.07 mmol of CO₂ per gram of material when operating at 650–800 °C.^{23,24}

Hydroxyapatite, HA, (Ca₁₀(PO₄)₆(OH)₂) is a widely studied and adaptable calcium phosphate material. Its potential for CO₂ utilisation arises from the fact that the HA crystal structure can accommodate a significant quantity of CO₂ owing to its ability to substitute both hydroxyl (OH[−]) and phosphate (PO₄^{3−}) groups for carbonate (CO₃^{2−}) ions without compromising the crystal structure of the material. This is achieved through chemical substitution mechanisms, termed A-type and B-type substitution respectively.⁸ A-type substitution can be attained by heating stoichiometric HA at high temperature in an atmosphere of CO₂, where a single carbonate ion replaces two hydroxyl groups to allow overall charge balance to be maintained in accordance with the substitution mechanism shown below:



Accordingly, A-type carbonate substitution has a theoretical maximum substitution level, achieved once all hydroxyl groups have been replaced with carbonate (*i.e.* when $x = 1$). The substitution of phosphate groups for carbonate groups can be achieved by the synthesis of a phosphate-deficient apatite ($\text{Ca}/\text{P} > 1.67$) with concomitant carbonate substitution introduced through the water used in the aqueous synthesis and/or the calcining/sintering atmosphere.⁹ To maintain charge balance, this substitution also requires equimolar substitution of carbonate ions for hydroxyl ions. This substitution mechanism therefore has a limit on the maximum level of carbonate substitution as it relies on the limiting A-type substitution. A common synthesis mechanism for substituting carbonate ions for phosphate groups in the hydroxyapatite lattice is by co-substitution of carbonate with a monovalent cation, typically sodium ions^{10,11} but also ammonium¹² or potassium ions.¹³ In these examples, some of the calcium ions are substituted for *e.g.* sodium ions allowing more carbonate to be incorporated into the structure, according to the charge-balance mechanism:



where M⁺ could be Na⁺, NH₄⁺ or K⁺. This gives the substitution mechanism Ca_{10−x}M_x(PO₄)_{6−x}(CO₃)_x(OH)₂. This mechanism focuses on balancing charge on the calcium and phosphate sites of HA, with any carbonate for hydroxyl substitution considered to occur independently. Our previous work showed that high levels of carbonate substitution, approximately 14 and 18 wt%, were achievable when sodium was the monovalent cation involved in the co-substitution, utilising NaCl and CO₂ gas¹⁰ or the more conventional sodium carbonate/sodium

hydrogen carbonate reactants,²⁵ respectively. In these compositions, however, the sodium counter ion does not provide any specific utilisation or benefit for an application of the material. Various methods, such as solid-state reaction,²⁹ hydrothermal reaction,³⁰ hydrolysis³¹ and ion-exchange,³² have been used to synthesise K–CO₃ co-substituted apatites with potassium and carbonate contents reported between approximately 0.4–6.9 wt% and 5.3–12.6 wt% respectively. Whyte *et al.*¹³ synthesised a number of K–CO₃ co-substituted hydroxyapatite materials, with substitution values of $x = 0.1–1.0$, by aqueous precipitation reaction between calcium hydroxide (Ca(OH)₂), phosphoric acid (H₃PO₄) and potassium hydrogen-carbonate (KHCO₃). These compositions were designed in such a way that potassium and carbonate ions would substitute onto calcium and phosphate sites respectively to such an extent that the overall (Ca + K)/(P + C) molar ratio was 1.67. Although the carbonate content of the samples was not directly measured, FTIR analysis suggested that this increased with x as intended. The compositions were then heated for two hours at temperatures ranging from 600 to 1050 °C in either air or CO₂ to test their thermal stabilities. The authors observed that the temperature at which the materials decomposed to produce secondary phases of calcium oxide decreased with increasing values of x (*i.e.* with increasing K⁺/CO₃^{2−} content) and also noted that decomposition occurred at higher temperatures when the samples were heated in an atmosphere of CO₂ compared to when they were heated in air.

The aim of the present work was to investigate the feasibility of using room temperature aqueous precipitation reactions between calcium hydroxide, phosphoric acid and either potassium carbonate or potassium hydrogen-carbonate to prepare several potassium-carbonate (K–CO₃) co-substituted apatite materials with high levels of carbonate incorporation. Potassium carbonate or potassium hydrogen-carbonate were chosen to provide the required K⁺ cations as both of these could theoretically be prepared on an industrial scale by reacting potassium hydroxide with captured CO₂. In some cases, this was followed by a heat treatment in a dry carbon dioxide atmosphere to determine what additional carbonate incorporation could be achieved. The prepared materials were characterised using a variety of techniques including powder X-ray diffraction and FTIR spectroscopy. A preliminary study was performed to assess the effect of the carbonate composition on the solubility of the compositions in an acidic buffer, to simulate the conditions of an acidic soil type.

2. Materials and methods

2.1. Synthesis of samples

Potassium-carbonate (K–CO₃) co-substituted apatites were synthesised at room temperature by aqueous precipitation reaction between calcium hydroxide (Ca(OH)₂, 98% assay, VWR, UK), phosphoric acid (H₃PO₄, 85% solution in water, Merck) and either potassium carbonate (K₂CO₃, ≥99% assay, Sigma-Aldrich, UK) or potassium hydrogen-carbonate (KHCO₃, 99.7% assay, Sigma-Aldrich, UK), following a method described



previously²⁵ that was based on an established precipitation reaction.³³ Briefly, two series of compositions were prepared with K₂CO₃ as the source of potassium ions in Series I and KHCO₃ used in Series II. Compositions based on the design formula Ca_{10-x}K_x(PO₄)_{6-x}(CO₃)_x(OH)₂ were prepared for $x = 0-3$ and the design compositions are provided in Tables S1 and S2 (ESI[†]). Precipitation reactions were maintained at an alkaline pH through the addition of 30 ml of aqueous NH₃ solution. The H₃PO₄ solution was added dropwise to the 'calcium/potassium' suspension of Ca(OH)₂ and K₂CO₃/KHCO₃ over a period of 2 hours, the mixture was aged overnight, filtered and washed with excess water, then the filter-cake was dried in an oven at 90 °C overnight. The dried filter cake was ground to a fine powder which was then subjected to a heat treatment in air at 300 °C for 1 hour using a muffle furnace (Carbolite Gero Ltd, UK), to remove synthesis residuals such as absorbed water; this did not affect the chemical composition of the materials markedly. Powders obtained after this heat treatment were designated 'as-prepared'. Aliquots (c. 0.5 g) of a selected composition were heated in a tube furnace (Carbolite Gero Ltd, UK) under a dry CO₂ gas flow rate of approximately 0.5 dm³ per minute. The temperature in the furnace was ramped from ambient up to the desired heating temperature of 300–700 °C at a rate of 5 °C min⁻¹, held there for one hour and then cooled back to room temperature at the same rate.

2.2. X-ray diffraction analysis

X-ray diffraction (XRD) patterns were collected with a X'Pert Pro diffractometer (PANalytical Ltd, UK), operating at 45 kV and 40 mA with Cu K α radiation ($\lambda = 1.5418 \text{ \AA}$). Data were collected from 15 to 65° 2 θ with a step size of 0.013° and a count time per step of 96 s. Crystalline phases present were identified by comparing obtained patterns with PDF files from the ICDD database. The average crystallite sizes were calculated using the Scherrer equation, using the (002) reflection at approximately 25.8° 2 θ , and with $k = 0.9$. A highly crystalline quartz standard was used to compute the instrumental line broadening. Lattice parameters were determined by simple Rietveld refinement of collected XRD data using the PANalytical software package 'HighScore Plus'.³⁴ Refinements were carried out using the hexagonal space group $P6_3/m$ using hydroxyapatite data reported by Sudarsanan and Young³⁵ as the initial structural model. Only the background function, scale factor, unit cell parameters and peak shape functions were refined.

2.3. Chemical composition analysis

The Ca/P molar ratio, potassium content and thus the (Ca + K)/P molar ratio of each sample was determined by energy-dispersive X-ray spectroscopy (EDX) using a Gemini SEM 300 field emission SEM (Zeiss, Germany) equipped with an X-ray detector (Oxford Instruments, UK) operating at an accelerating voltage of 15 kV. A powder compact was coated with carbon prior to analysis. For each sample, three measurements were taken and the results reported as an average alongside the standard deviation. The carbonate contents of samples were determined using a LECO CS744 carbon/sulphur analyser

(LECO Instruments UK Ltd, UK). For each sample, duplicate measurements were made and the mean value reported alongside the standard deviation. It was assumed that all of the carbon detected by the equipment existed as CO₂, which was reasonable as the equipment flooded the combustion chamber with oxygen and also passed the combustion products through an oxidation catalyst.

2.4. FTIR analysis

FTIR spectra were obtained using a Diamond/ZnSe ATR attached to a Spectrum Two™ spectrometer (PerkinElmer, UK). Absorbance spectra were collected at a 2 cm⁻¹ resolution between 4000 and 400 cm⁻¹, averaging 7 scans per sample. Peak deconvolution of the carbonate ν_2 (850–890 cm⁻¹) and ν_3 (1300–1600 cm⁻¹) regions and the phosphate ν_4 (485–700 cm⁻¹) region of the spectra was performed by fitting Gaussian distributions, using the Solver extension in Excel,³⁶ based on peak assignment described elsewhere; a linear background was subtracted prior to deconvolution.

2.5. Dissolution evaluation

The relative solubilities of the different compositions were assessed by incubating as-prepared powder samples (sieved to 53–315 μm size) in an acidic (pH 5.5) buffer solution for 30 minutes and measuring the Ca ion concentration, based on a method described by Mano *et al.*,³⁷ For each composition studied, 30 mg of sample was incubated in 20 ml of 0.08 M sodium acetate–acetic acid buffer solution, prepared as described elsewhere,³⁸ for 30 minutes at 20 °C, under 150 rpm mixing. A 100 μl aliquot was collected for the measurement of Ca ion concentration in the buffer solution. For each composition 3 replicates were prepared.

The calcium ion concentration was determined using a colorimetric assay based on the *ortho*-cresolphthalein complexone (*o*-CPC) method described by Connerty *et al.*³⁹ Briefly, stock solutions of 1.66 mM *o*-CPC (64000-1G, Sigma Aldrich, UK), 0.344 M 8-hydroxyquinoline (8-HQ) (H6878-25G, Sigma Aldrich, UK) and 14.8 M ethanolamine (1008451000, Sigma Aldrich, UK) were prepared. A working solution was prepared by combining these three stock solutions to achieve final concentrations of *o*-CPC and 8-HQ of 0.083 mM and 6.89 mM, respectively, and the concentration of ethanolamine was 0.74 M. A series of Ca²⁺ standards were prepared by diluting a 1000 ppm Ca²⁺ standard (21049-100ML, Sigma Aldrich, UK) in water. In a 96 well plate 300 μl of working solution per well was added to 10 μl of each standard/test sample to be measured, the mixture was allowed to react for 5 minutes, then the absorbance at 575 nm was read on a plate reader (BioTek Synergy HT). For each Ca²⁺ standard duplicate wells were prepared, and for each test sample triplicate wells were prepared.

2.6. Statistical analysis

Comparison of dissolution data from multiple groups ($x = 0-2.0$) was performed with GraphPad Prism (v5.04, GraphPad Software Inc., USA) using one-way analysis of variance (ANOVA) with a *post hoc* Tukey's test. For assessment of differences in dissolution



data for $x = 1.5$ from samples synthesised with two different sources of potassium, a t -test was used. Results are reported as mean \pm SD. Values of $p < 0.05$ are considered to be statistically significant.

3. Results and discussion

3.1. As-prepared apatites

3.1.1. X-ray diffraction (XRD) analysis. XRD patterns of the as-prepared powders of each series are shown in Fig. 1. In Series I (when K_2CO_3 was used), the compositions were phase-pure apatites by XRD for design substitution values up to and including $x = 2.5$; all of the reflections in the collected patterns were comparable to the ICDD standard⁴⁰ for hydroxyapatite. By comparison, the compositions of Series II (when $KHCO_3$ was used) were single-phase apatites only up to $x = 1.5$. Although this study did not explore the reasons for these differing limits, it would seem reasonable to conclude that this disparity was related to the nature of the potassium reagents themselves. For example, perhaps the lower x -value achieved using $KHCO_3$ was a result of HCO_3^- ions having substituted into the lattice or (as $KHCO_3$ is significantly less basic than K_2CO_3 ⁴¹) as a result of a lower reaction mixture pH (the pH of the aged reaction mixtures of Series II tended to be up to a full integer lower than their counterparts of Series I). Also, the HA reflections in all of the collected patterns were quite broad which indicated that each hydroxyapatite phase was composed of very small crystallites as would be expected when using an aqueous precipitation synthesis route.^{42,43} Above these limits of

x , an impurity phase of calcite ($CaCO_3$) was observed to form alongside the apatite phase; diffraction peaks attributable to these secondary phases are explicitly marked in Fig. 1. Crystallite sizes were calculated using the Scherrer equation, using the (002) reflection of HA at approximately $25.8^\circ 2\theta$, and were of the order of 18–29 nm, with some correlation between the design composition and crystallite size, Table S3 (ESI†).

The unit cell parameters of the as-prepared apatites are shown in full in Table S4 (ESI†) and are plotted as a function of the design composition value of x and of the measured carbonate content (described later in Section 3.1.2) of the materials in Fig. 2. In each series, there was a linear contraction in the a lattice parameter and the unit cell volume as the design x -value increased; this linear contraction was also observed when plotted against the measured carbonate contents. There was a linear decrease in the a lattice parameter with increasing measured potassium content for the design compositions between $x = 0$ –1.5, but there was then a loss in any correlation for higher values of Series I (plotted data not shown, but data are provided in Tables S4–S6, ESI†). It is known that B-type carbonate substitution causes the c -axis to expand and the a -axis to contract, whilst A-type carbonate substitution has the opposite effects on the unit cell parameters of HA.⁸ Therefore, although there was no clear trend in the c unit cell parameter in either series with $x/K^+/CO_3^{2-}$ content, the variations in the a lattice parameter and the unit cell volume suggested that the substitution of carbonate ions on phosphate sites had more of an impact on the hydroxyapatite crystal structure, and thus the dimensions of the unit cell of the as-prepared material, than did the substitution of carbonate ions on hydroxyl sites.

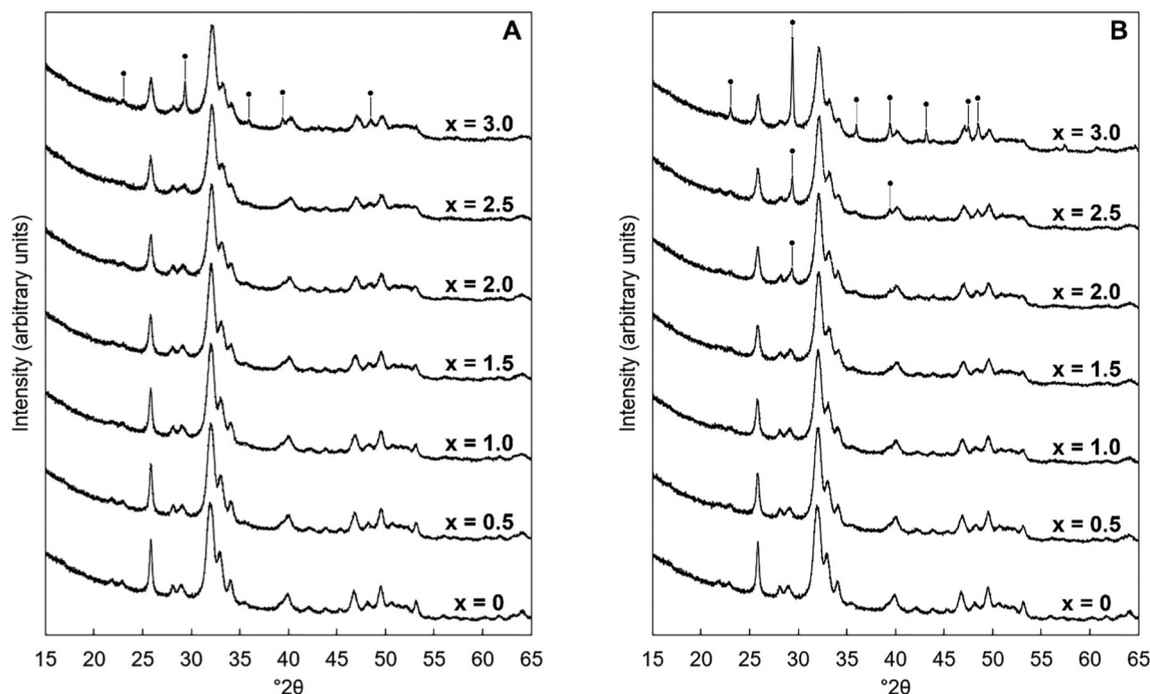


Fig. 1 Normalised ($I_{max} = 100$) XRD patterns between 15 – $65^\circ 2\theta$ of the as-prepared samples of Series I (A) and Series II (B), where $x = 0$ – 3.0 in the design formula $Ca_{10-x}K_x(PO_4)_6-x(CO_3)_x(OH)_2$. Intense reflections corresponding to impurity phases of calcite are marked using ●.



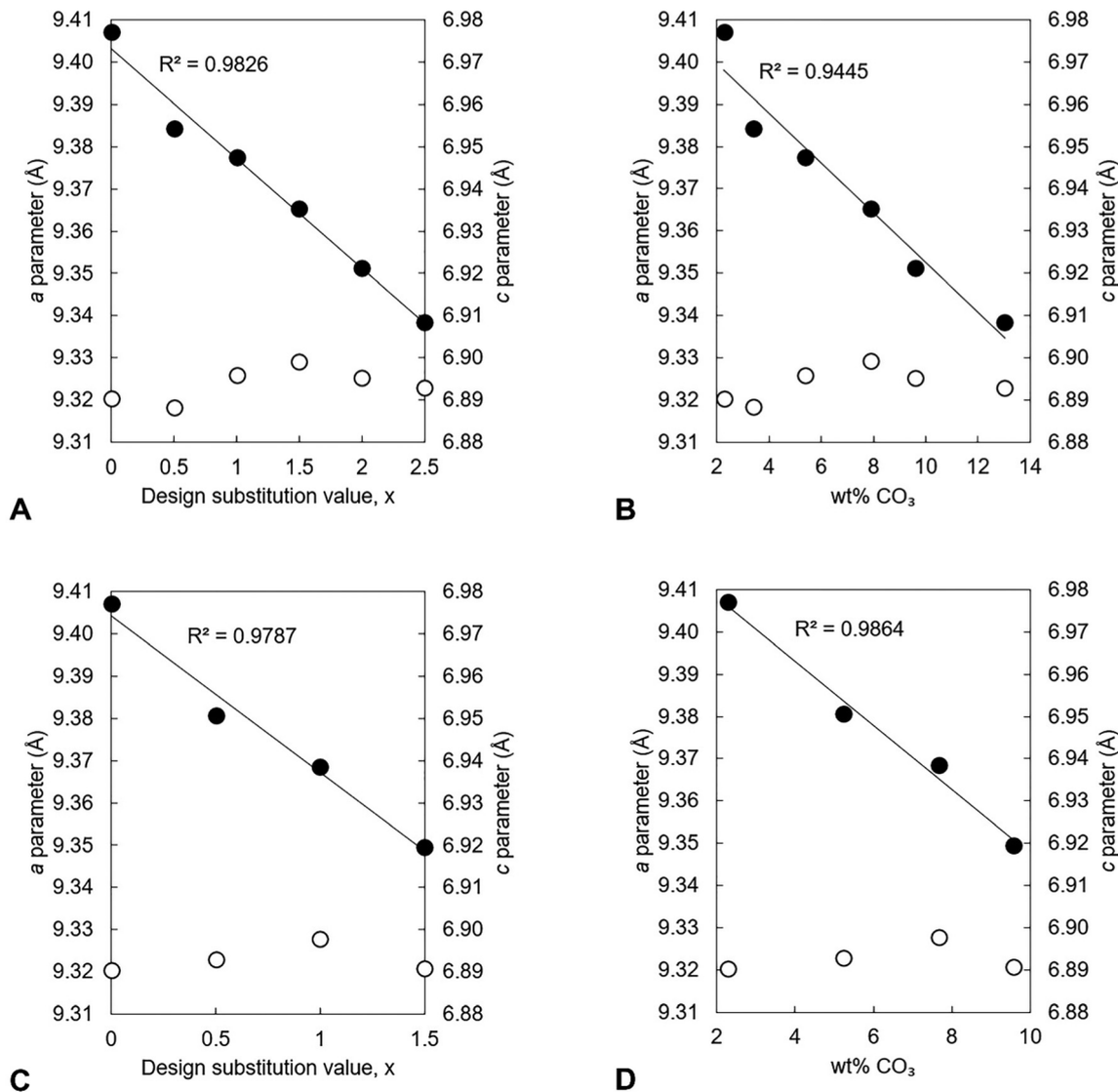


Fig. 2 The a (●) and c (○) lattice parameters of the as-prepared apatites of Series I (A and B) and II (C and D) plotted as a function of the design substitution value x and the measured carbonate content of these materials.

3.1.2. Chemical analysis. The potassium content as well as the Ca/P and (Ca + K)/P molar ratios of the as-prepared apatites were measured using EDX and their carbonate contents measured using combustion analysis. The results of this analysis are presented in full in Tables S5 and S6 (ESI[†]) and demonstrate that the chemical composition of these as-prepared materials did not match the design formula. Although potassium ions were indeed present in each of the as-prepared apatites, all of these compositions contained significantly less of these cations than the design amount irrespective of the potassium reagent used in the precipitation reaction, Fig. 3A. In Series I, the measured potassium content of the materials appeared to level out once a certain value of x had been achieved, suggesting that a maximum existed at approximately 0.9 wt% potassium. These lower than expected potassium contents (and the failure to detect any impurity phases using XRD) suggested that, instead of having substituted into the

apatite lattice, most of the potassium cations that were added to the reaction mixture had remained in solution and then been lost during the filtration step. Synthesis of sodium/carbonate substituted hydroxyapatite compositions also observed lower than expected levels of sodium substitution, even though increasing amounts of sodium carbonate reactant added resulted in increasing levels of carbonate substitution.²⁵ NH_4^+ ions are known to be capable of substituting for the calcium ions in hydroxyapatite⁴⁴ and it could therefore be hypothesised that perhaps some competition existed between K^+ ions and NH_4^+ ions for the open calcium sites. C-H-N analysis was performed on several of the as-prepared apatites but this failed to detect the presence of nitrogen in any of the samples (data not shown). Even so, it is possible that nitrogen existed in very small amounts that could not be detected by the equipment or that this element had been removed during the heat treatment in air at 300 °C and so the synthesis of the $x = 0.5$ and $x = 2.5$



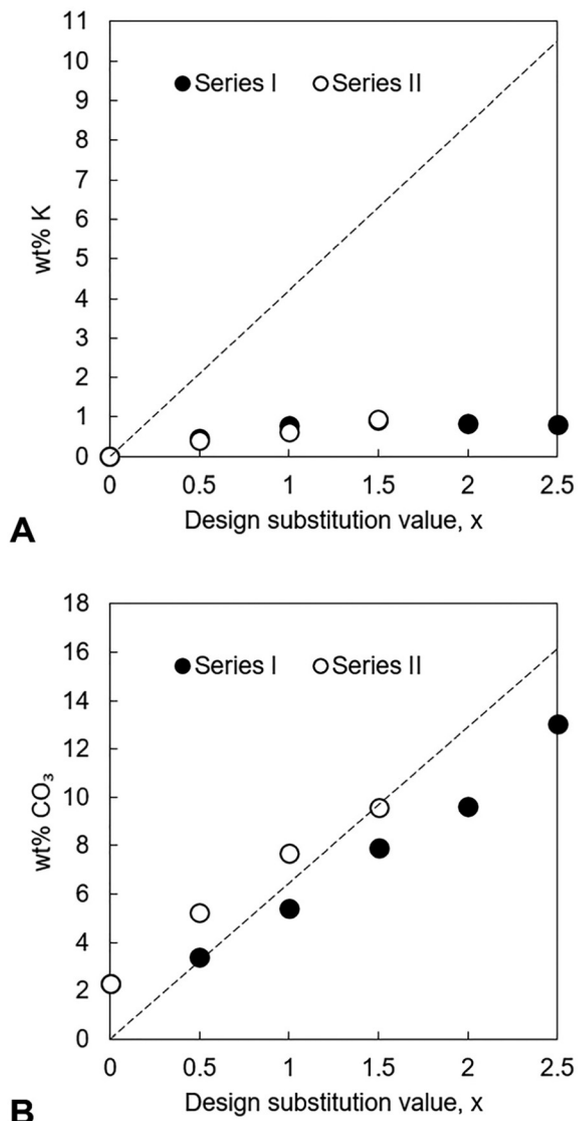


Fig. 3 Measured potassium (A) and carbonate contents (B), in weight percent, of each series of as-prepared apatites plotted as a function of x . The dashed lines represent the theoretical K^+/CO_3^{2-} content at each value of x , calculated from the design composition formula $Ca_{10-x}K_x(PO_4)_{6-x}(CO_3)_x(OH)_2$.

apatites of Series I was repeated but this time without adding NH_4OH solution to the calcium/potassium suspension. Resulting EDX analysis showed that the exclusion of ammonia solution did not lead to the preparation of apatites with larger potassium contents (Table S7, ESI†) and for $x = 2.5$ the potassium content was significantly lower when ammonia solution was excluded. While EDX analysis provides a quantitative measure of the trend in increasing Ca/P molar ratio and K content with increasing substitution, additional analytical quantification by spectroscopy methods such as ICP-OES, would provide more precise quantification of the chemical compositions.

In both series, the measured Ca/P molar ratio increased linearly with x . This signalled that the relative content of phosphate ions in the apatites decreased as the design

substitution value rose in order to accommodate the substitution of carbonate ions onto B-sites as designed. Whereas the measured Ca/P molar ratios of the apatites prepared using $KHCO_3$ were comparable to the design values, this ratio was invariably smaller than designed in the apatites of Series I. The measured (Ca + K)/P molar ratio of every apatite was lower than designed, a result primarily of the much smaller than expected potassium contents. The carbonate content also increased linearly with increasing x in each series, Fig. 3B. Although the greatest CO_3^{2-} content (13.0 wt%) was achieved using K_2CO_3 for $x = 2.5$, the carbonate contents of Series I were smaller than equivalent design compositions (x values) prepared using $KHCO_3$ and also tended to be smaller than was expected according to the design formula. Nevertheless, these results support the concept that a two-step process could be developed where K_2CO_3 or $KHCO_3$ is first produced using captured CO_2 and then used in the precipitation of highly carbonated K- CO_3 co-substituted apatites.

In an effort to increase the maximum degree of carbonation, the synthesis of each series was repeated with CO_2 gas bubbled into the H_3PO_4 solution during the precipitation reaction to create an excess of carbonate ions in the reaction mixture. This gas was bubbled into the H_3PO_4 solution for 30 minutes prior to this solution being added dropwise to the basic suspension and was maintained until the addition of the H_3PO_4 solution was complete. The dissolution of CO_2 gas, and the subsequent evolution of carbonic acid (H_2CO_3),⁴⁵ did not affect the temperature of the H_3PO_4 solution and by extension the reaction mixture to any significant extent. Whilst the bubbling of this gas into the H_3PO_4 solution led to a significant increase in the carbonate content of the apatites of Series I at low values of x , its influence diminished markedly as the design substitution value increased (Table 1). For example, the $x = 2.5$ apatite prepared using K_2CO_3 and with CO_2 gas bubbled into the H_3PO_4 solution contained 14.1 wt% of carbonate, only marginally larger than the 13.0 wt% seen in the $x = 2.5$ apatite of Series I. Instead, the carbonate ions that were introduced into the reaction mixture from the potassium reagent itself (in addition to those present from the dissolution of atmospheric CO_2) were sufficient to reach the high levels of carbonation that were achieved. The differences in the measured carbonate content in the $KHCO_3$ series, with or without gas added during the reaction, were much smaller, for any x value, than the K_2CO_3 series; per mole of potassium, the $KHCO_3$ reactant provides more carbonate to the reaction, so this may explain this observation.

3.1.3. Fourier-transform infrared spectroscopy (FTIR). FTIR spectra of the as-prepared K- CO_3 co-substituted apatites are presented in Fig. 4. These spectra were comparable to those seen in other studies of apatites prepared by aqueous precipitation reaction.^{13,46} The four IR-active phosphate vibrations that are characteristic of hydroxyapatite were observed in each spectrum. These appeared to broaden slightly in each series as the design substitution value increased, which may have been due to decreasing crystallite size along the c -axis.⁴⁷ In addition, a broad band (centred at approximately 1640 cm^{-1})



Table 1 Carbonate contents of as-prepared K–CO₃ co-substituted apatites prepared with K₂CO₃/KHCO₃ and with/without CO₂ gas bubbled into the H₃PO₄ solution during the precipitation reaction, where $x = 0$ –2.5 in the design composition formula Ca_{10–x}K_x(PO₄)_{6–x}(CO₃)_x(OH)₂. When CO₂ gas was used, this gas was bubbled into the H₃PO₄ solution for 30 minutes prior to this solution being added to the calcium/potassium suspension and continued until the addition of the H₃PO₄ solution was complete. Other than this alteration, the procedure used to prepare those apatites was identical to that used for the apatites of Series I and II. Analysis was only performed on samples that were single-phase apatites by XRD analysis

Design x	K ₂ CO ₃		KHCO ₃	
	No CO ₂ gas (series I)	CO ₂ gas	No CO ₂ gas (series II)	CO ₂ gas
0	2.28 ± 0.03	3.82 ± 0.01	2.28 ± 0.03	3.82 ± 0.01
0.5	3.39 ± 0.00	5.68 ± 0.04	5.23 ± 0.04	5.80 ± 0.00
1.0	5.40 ± 0.00	7.98 ± 0.04	7.68 ± 0.04	8.33 ± 0.04
1.5	7.90 ± 0.00	8.93 ± 0.04	9.58 ± 0.04	10.10 ± 0.07
2.0	9.60 ± 0.00	11.00 ± 0.07	N/A	N/A
2.5	13.03 ± 0.04	14.08 ± 0.04	N/A	N/A

attributed to the presence of absorbed water⁴⁸ was visible in each spectrum although this was very weak. The presence of this vibration indicated that neither the oven drying nor the heat treatment in air at 300 °C had fully dried the materials. Vibrations corresponding to the librational and stretching modes of hydroxyl ions were observed in the compositions of both series (at approximately 631 and 3570 cm⁻¹ respectively) at low values of x . Carbonate ν_2 and ν_3 bands were also visible in each of these spectra, Fig. 4A, C, D and F, with contributions corresponding to the substitution of carbonate ions on hydroxyl and phosphate sites, consistent with previous studies.^{8,49–52}

An example of the peak deconvolution of the OH libration and PO₄ ν_4 region is shown in Fig. 5A and B for the design composition of $x = 1.5$ of Series I and Series II, respectively, and the peak positions (numbered from 1a–5a from the highest to the lowest wavenumber) and areas of the phosphate ν_4 region, including the OH libration, are listed in full in Table S8 (ESI†). Assignment of vibrations in carbonate-containing apatites can be challenging as many parameters that are specific to a given study, such as chemical composition, preparation method, monovalent ion content, thermal treatment temperature and atmosphere can influence the peak positions and intensity of carbonate and phosphate vibrations, so definitive assignment based on literature data must be made with caution. In addition to the expected three PO₄ vibrations (peaks 2a, 3a and 4a) at approximately 604–601, 582–576 and 565–561 cm⁻¹, a peak at approximately 545–540 cm⁻¹ (peak 5a) was assigned to HPO₄ groups in the apatite structure.⁵³ The peak area of this vibration appeared to decrease slightly with increasing x value in both Series I and II. Peak deconvolution established a peak with decreasing intensity at 631–626 cm⁻¹ (peak 1a) as the value of x increased, which is assigned to hydroxyl libration. The three peaks of the triply degenerate bending mode of the PO₄ ν_4 region did not show any strong trend in peak position or peak area with increasing substitution or between the two series, although there appeared to be a slight shift to higher wavenumber of the ν_{4b}

vibration (peak 3a) from 576 cm⁻¹ for $x = 0$ up to 582 cm⁻¹ for $x = 2.5$. The carbonate ν_2 region produced a very broad band with poor definition of individual vibrations and peak deconvolution required 7–8 peaks to achieve a reasonable fit, although for the highest level of substitution, $x = 2.5$, Series I, a reasonable fit to 3 peaks was possible. The most intense peaks from the fit were at approximately 877–875, 870–868 and 863–848 cm⁻¹, which are consistent with assignments to A-type (for hydroxyl), B1-type (for phosphate) and B2-type (for phosphate) carbonate substitution, respectively,⁵¹ but with a small downshift. A vibration in the region 863–861 cm⁻¹ has been assigned to labile carbonate, HPO₄ groups and B2-type carbonate, depending on the composition, so care should be taken in the assignment of this peak in this broad, poorly resolved ν_2 region. Peak deconvolution of the carbonate ν_3 regions produced a fit comprising 6 peaks, with examples for the design composition of $x = 1.5$ of Series I and Series II shown in Fig. 5C and D. The peak positions and areas (arbitrarily numbered from 1b–6b from the highest to the lowest wavenumber) of the carbonate ν_3 region from the spectra of all single-phase as-prepared compositions in Series I and II are listed in full in Table S8 (ESI†). Three peaks (2b, 3b and 5b) are dominant in the substituted samples, in terms of their peak areas, at 1492–1486 cm⁻¹, 1456–1447 cm⁻¹ and 1412–1401 cm⁻¹; the $x = 0$ sample produced low intensity vibrations in this region and the peak positions were all closer in peak areas, except the peak at 1417 cm⁻¹ which dominated, and peak positions were typically 10–20 cm⁻¹ higher than the positions of corresponding peaks in the substituted samples. Peak 2b (1492–1486 cm⁻¹) can be assigned to A-type substitution and peak 5b (1412–1401 cm⁻¹) can be assigned to B-type substitution.^{8,49–52,54–56} Assignment of peak 3b (1456–1447 cm⁻¹) to literature data does not have such a clear consensus, with studies assigning this to both A-type substitution^{49,51,56} and B-type substitution.^{50,54,55} In our own recent study, A-type carbonate substituted hydroxyapatites produced peaks at approximately 1532 and 1465 cm⁻¹ with similar peak areas.⁵⁷ The peak area of the peak in the 1557–1523 cm⁻¹ region in the current study contributes a much smaller amount to the total peak area of the ν_3 region than the peak at 1456–1447 cm⁻¹, Table S8 (ESI†), suggesting that the latter peak reflects some contribution of A-type carbonate in this region but relates more to B-type carbonate. Weaker bands (peaks 1b, 4b and 6b) at 1557–1523 cm⁻¹, 1424–1419 cm⁻¹ and 1393–1355 cm⁻¹, show some small changes in peak position and area with increasing substitution and between Series I and II, but the most notable was the shift to lower wavenumbers of peak 1b, from approximately 1565 cm⁻¹ for $x = 0$ to 1523 cm⁻¹ for $x = 2.5$. Studies on Na-containing apatites have described distinct A-type vibrations, with bands at 1540 and 1449 cm⁻¹ assigned to type-A1, and type A2 to bands at 1569 and 1507 cm⁻¹.⁵⁸ Peak 1b could be assigned to type A1, with some intensity also within the peak 3b. No clear vibration was identified at approximately 1473 cm⁻¹ which is commonly assigned to B-type substitution, nor a peak at approximately 1565 cm⁻¹ which is commonly assigned to A-type substitution. Yoder *et al.* produced a potassium–carbonate substituted hydroxyapatite using KHCO₃ which had measured potassium



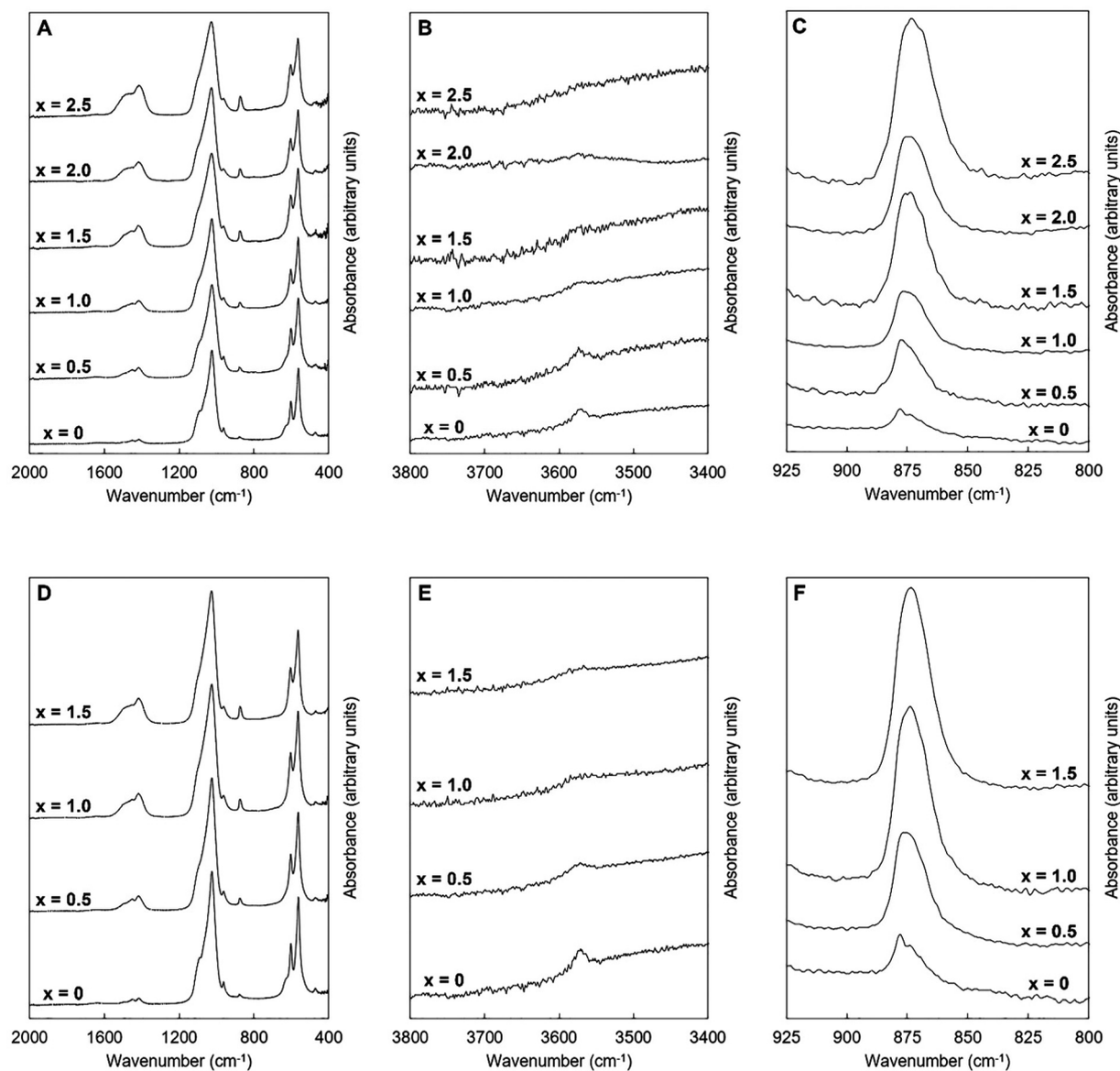


Fig. 4 Normalised ($A_{\max} = 100$) FTIR absorbance spectra of the single-phase as-prepared apatites of Series I (A–C) and II (D–F), where $x = 0$ –2.5 in the design formula $\text{Ca}_{10-x}\text{K}_x(\text{PO}_4)_6-x(\text{CO}_3)_x(\text{OH})_2$.

(0.51 wt%) and carbonate ion (5.99 wt%) contents that were comparable to $x = 0.5$ of Series II in this study (Table S6, ESI[†]) and, without peak deconvolution, identified carbonate vibrations at 1490 and 1540 cm^{-1} that were assigned to A-type and 1417 and 1451 cm^{-1} that were assigned to B-type carbonate.⁵⁴ A band at 1387 cm^{-1} has been assigned to carbonate in a water compartment, described as within water trapped in pores within grains, or water that is surface adsorbed⁴⁹ and, as a weak vibration from the presence of water was visible, this is feasible. No vibrations characteristic of calcite or aragonite were observed at ~ 713 or 700 cm^{-1} in any of these spectra, confirming the absence of a calcium carbonate impurity in these samples;⁵⁹ a peak at 714 cm^{-1} was observed in the spectrum of the $x = 3.0$ sample of Series I (data not shown), which showed an impurity of calcite by XRD (Fig. 1). Overall it

appears that increasing levels of carbonate and potassium substitution have little impact on the vibrations present and their relative contributions to total carbonate in the as-prepared samples of Series I and II.

3.1.4. Dissolution evaluation. To identify if increasing the amount of potassium/carbonate substitution would affect the solubility of hydroxyapatite, powder samples of the Series I as-prepared compositions for $x = 0.5$ to 2.0, Series II composition with $x = 1.5$, and an unsubstituted composition ($x = 0$) were incubated in an acidic (pH 5.5) buffer for 30 minutes, then the Ca^{2+} ion concentration in the buffer solution measured using a colorimetric assay, Fig. 6. Increasing the design composition in Series I, resulting in an increase in carbonate content, and to a lesser extent an increase in potassium content from $x = 0.5$ to 1.5 (Tables S5 and S6, ESI[†]), resulted in an increase in solubility



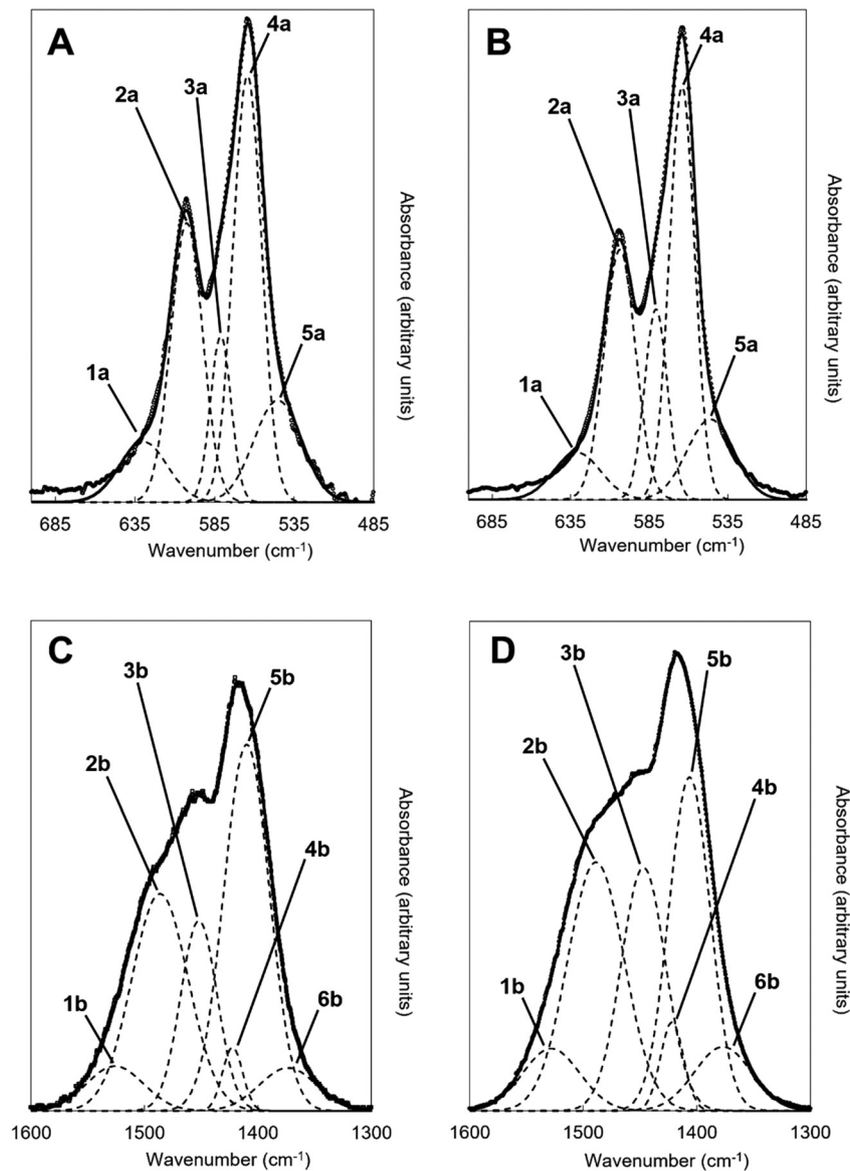


Fig. 5 Deconvolution of the (A and B) OH_1 and $\text{PO}_4 \nu_4$ and (C and D) $\text{CO}_3 \nu_3$ regions of the FTIR spectra of the as-prepared $x = 1.5$ compositions for (A and C) Series I (K_2CO_3) and (B and D) Series II (KHCO_3) potassium-carbonate co-substituted samples showing the various vibrations that contribute to the observed spectra; y-axis is an arbitrary scale. Dashed lines correspond to the individual fitted peaks and the solid line is the corresponding model spectra, with the raw data presented as isolated data points.

under acidic conditions, with a significant increase in the calcium ion concentration of the buffer solution for design compositions with $x = 1.5$ and 2.0 ($p < 0.05$), compared to the unsubstituted $x = 0$ HA, Fig. 6A. Comparing the two sources of potassium carbonate, the $x = 1.5$ composition prepared with KHCO_3 (Series II) resulted in a higher calcium ion concentration of the buffer solution than the $x = 1.5$ composition prepared with K_2CO_3 (Series I), Fig. 6B. The former had a higher measured carbonate content (9.58 wt%) than the latter (7.9 wt%), while both had comparable potassium contents (Tables S5 and S6, ESI[†]), which supports the view that the carbonate content of the materials correlates with the solubility. Plotting the calcium ion concentration in the buffer solution against the measured

carbonate contents of the co-substituted samples from Fig. 3 produced a linear fit with $R^2 = 0.986$, (data not shown). Xiong *et al.* studied the effect of surface charge of HA nanoparticles on the P release of hydroxyapatite nanoparticles as potential alternatives to triple super phosphate (TSP) fertilisers (monocalcium phosphate) in an acidic soil (pH 4.7) and observed that sustained release of phosphate was dependent on the HA nanoparticles; an electronegative surface charge resulted in the HA nanoparticles having a slower and more sustained release of phosphorus than the TSP fertiliser.²² Our data, while preliminary, suggests that modifying the composition of hydroxyapatite nanoparticles by potassium/carbonate co-substitution is another method to control the solubility, and



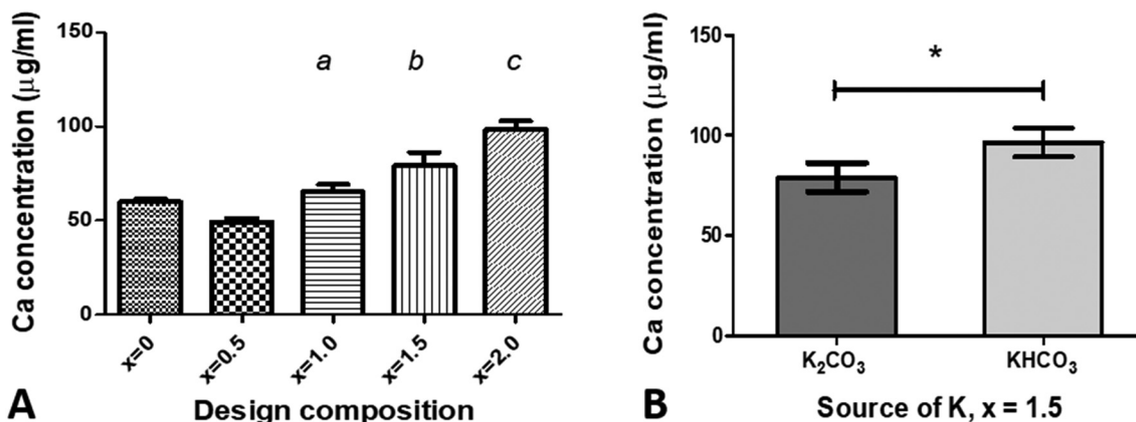


Fig. 6 Measurement of Ca^{2+} ion concentration in the sodium acetate–acetic acid buffer after incubation of samples for 30 minutes, for (A) various compositions of x using K_2CO_3 as the source of potassium, and (B) a composition of $x = 1.5$ using two different sources of potassium, K_2CO_3 and KHCO_3 . Data are presented as the mean \pm SD ($n = 3$). Statistical differences, based on $p < 0.05$, are noted as: (a) significantly different from $x = 0.5$; (b) significantly different from $x = 0, 0.5$ and 1.0 ; (c) significantly different from $x = 0, 0.5, 1.0$ and 1.5 ; and * – significantly different between K source.

therefore the phosphorus release, with the additional potential of a concomitant release of potassium ions (albeit it in much smaller quantities than would have been predicted from the design formula).

3.2. Heated apatite(s)

3.2.1. X-ray diffraction (XRD) analysis. The $x = 2.5$ apatite of Series I was subjected to several heat treatments in dry CO_2 to test if the degree of carbonate substitution in the material could be increased. This composition was chosen as it exhibited the largest carbonate content of all of the as-prepared apatites. XRD patterns of samples heated at 300–700 °C are presented in Fig. 7. Whilst the material remained as a phase-pure apatite after it was heated at 300–600 °C, a secondary phase of calcite was produced after heat treatment at 700 °C. This impurity phase would likely have initially formed as CaO due to the calcium-rich (*i.e.* Ca/P molar ratio > 1.67) nature of the composition,⁶⁰ but due to the CO_2 atmosphere of the heat treatment underwent carbonation to form calcite. The formation of this secondary phase demonstrated that this K– CO_3 co-substituted apatite was significantly less thermally stable than stoichiometric HA, which has been shown to be stable in dry or moist air up to approximately 1200 °C.⁶¹ This disparity was to be expected and can be explained by examining the differing mechanisms which govern the thermal decomposition of stoichiometric HA and CHAs. Whereas stoichiometric HA is known to decompose as a result of a loss of channel hydroxyl groups above a certain temperature *via* an oxyhydroxyapatite intermediate, CHAs are understood to decompose upon heating due to the loss of substituted carbonate groups as CO_2 . Whilst the loss of hydroxyl groups has been reported to begin at approximately 1000 °C in air,⁶² the loss of carbonate groups from the HA lattice typically begins at much lower temperatures.^{63,64}

The lattice parameters of the K– CO_3 co-substituted apatites after being heated in dry CO_2 at 300–600 °C are shown in full in Table S9 (ESI[†]) and plotted as a function of the heat treatment

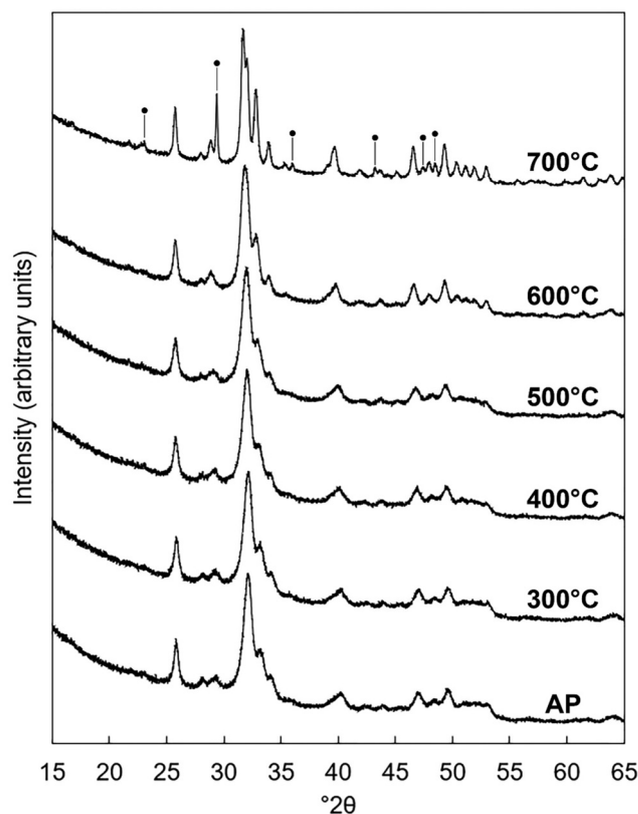


Fig. 7 Normalised ($I_{\text{max}} = 100$) XRD patterns of the $x = 2.5$ composition of Series I as-prepared (AP) and after being heated for one hour in dry CO_2 at 300–700 °C. Intense reflections corresponding to an impurity phase of calcite are marked using ●.

temperature in Fig. 8. Both the a and c unit cell parameters (as well as the unit cell volume) of the heated samples tended to be larger than the as-prepared material, with the magnitude of these expansions proportional to the temperature at which the heat treatments were performed. These trends can be explained by considering two scenarios. The first is that CO_3^{2-} ions had



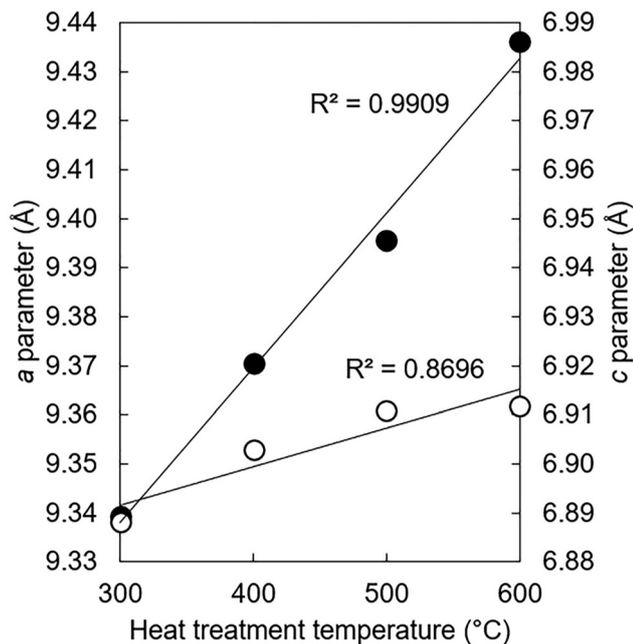


Fig. 8 a (●) and c (○) unit cell parameters of the $x = 2.5$ apatite of Series I after this material had been subjected to a heat treatment in dry CO_2 at 300–600 °C.

substituted during the heat-treatment exclusively for the relatively smaller hydroxyl groups and oriented mostly parallel to the ab plane, producing the observed expansions in the a parameter and the unit cell volume in accordance with Vegard's law.⁶⁵ Although expansions in the c unit cell parameter are not usually associated with this type of carbonate substitution, the expansions were relatively small and so could be accounted for by a non-parallel component in the orientation of the substituted carbonate groups. A-type carbonate substitution is usually associated with higher temperature treatments in a CO_2 atmosphere for long durations of time, but appreciable A-type substitution can occur at lower temperatures and for shorter time periods.⁵⁷ Alternatively, carbonate ions may have simultaneously substituted onto both hydroxyl and phosphate sites. Here, the relative expansions of a and c could be used as evidence that the heat treatment had brought about significantly more CO_3^{2-} substitution on the A-sites than it did the B-sites. Unfortunately, the quality of the diffraction patterns of these nanoscale samples did not allow for a full structure refinement (such as atomic coordinates and site occupancy) to be performed, which is a similar problem to diffraction data from bioapatites.⁶⁶

3.2.2. Quantitative carbonate analysis. The results of combustion analysis performed on the $x = 2.5$, Series I samples that had remained as phase-pure apatites after the heat treatments in dry CO_2 are shown in Table 2. Each of the heated samples contained more carbonate than in the as-prepared sample. There was a consistent increase in the carbonate content of the apatite as the temperature increased from 300–600 °C, which suggested that over this temperature range, the carbonate ions previously incorporated into the apatite had been

Table 2 Measured carbonate content of the $x = 2.5$ apatite of Series I as-prepared (AP) and after being heated for one hour in dry CO_2 at 300–600 °C

Temperature (°C)	CO_3^{2-} content (wt%)
AP	13.03 ± 0.04
300	14.78 ± 0.18
400	16.20 ± 0.07
500	16.88 ± 0.11
600	16.93 ± 0.18

preserved, likely as a result of the CO_2 atmosphere in which the heat treatments were performed. After it had been heated in dry CO_2 at 600 °C, the apatite material contained approximately 16.9 wt% of carbonate. We believe that this may be the largest carbonate content that has been observed for a single phase K- CO_3 co-substituted apatite to date, with other such materials having been reported to contain between 4.95–12.6 wt% of carbonate.^{29–32,54}

Self-evidently, one of the key performance indicators as to the viability of a given CO_2 utilisation option is the quantity of carbon dioxide that can be stored. Table 3 demonstrates that the equivalent quantity of carbon dioxide incorporated during the synthesis of this apatite was of the same order of magnitude, but lower than, a number of other widely proposed utilisation options. However, it should be kept in mind that all sectors must be decarbonised if net-zero is to be achieved and it is to be expected that some options will perform better than others with regards to this indicator.

3.2.3. FTIR analysis of heated samples. FTIR spectra of the potassium-carbonate co-substituted apatite material ($x = 2.5$, Series I) after it had been subjected to the heat treatments in dry CO_2 are displayed in Fig. 9. The heat treatments caused the broad band at approximately 1640 cm^{-1} , attributable to the presence of absorbed water, to disappear. Additionally, heating the apatite improved the resolution of the contributions corresponding to A and B-type carbonate substitution in the ν_3 and ν_2 carbonate regions,⁵² Fig. 9B and C. As for the as-prepared samples, deconvolution of the OH_L and $\text{PO}_4 \nu_4$ region revealed the expected 3 peaks (peaks 2A–4A) related to the PO_4 vibrations in apatites, with no significant change in position or relative peak area with increasing heat treatment temperature

Table 3 A comparison of various CO_2 utilisation options in terms of the quantity of carbon dioxide incorporated. The apatites referenced here correspond to the $x = 2.5$ apatite of Series I, which contained approximately 13 wt% carbonate as-prepared and 16.9 wt% CO_3^{2-} after being heated for one hour at 600 °C in dry CO_2

Utilisation option	CO_2 utilised (mmol CO_2 per g material)	Ref.
As-prepared K- CO_3 apatite	2.3	N/A
Heated K- CO_3 apatite (600 °C in dry CO_2)	2.7	N/A
Calcium carbonate, CaCO_3	9.9	67
Magnesium carbonate, MgCO_3	11.8	68
Nesquehonite, $\text{MgCO}_3 \cdot 3\text{H}_2\text{O}$	7.3	5
Methanol, CH_3OH	31.1	3
Acetic acid, CH_3COOH	16.6	3
Benzoic acid, $\text{C}_6\text{H}_5\text{COOH}$	8.2	3



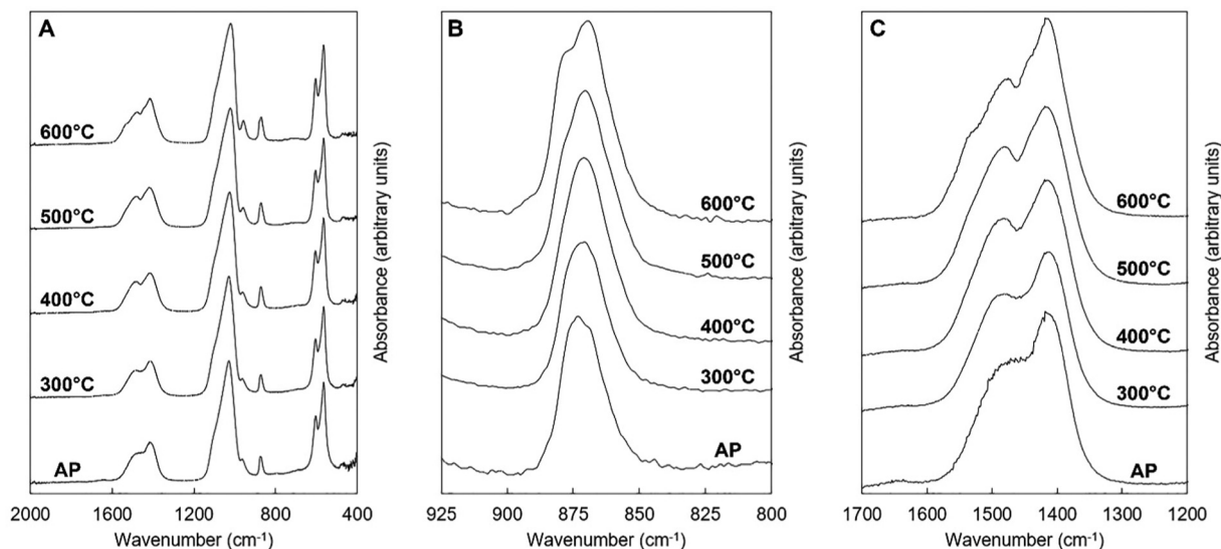


Fig. 9 Normalised ($A_{\max} = 100$) FTIR spectra of the $x = 2.5$ apatite of Series I as-prepared (AP) and after being subjected to heat treatments in dry CO_2 at 300–600 °C.

Table 4 The deconvoluted peak positions (cm^{-1}) and relative peak areas (%) from the FTIR spectra of the OH_L and phosphate ν_4 (1A–5A), carbonate ν_3 (1B–6B) and carbonate ν_2 (1C–4C) regions of the $x = 2.5$ apatite of Series I as-prepared (AP) and after being heated for one hour in dry CO_2 at 300–600 °C. Peaks are arbitrarily numbered from 1A–5A, or 1B–6B, or 1C–4C, from high to low wavenumbers. Due to rounding peak areas to whole numbers the sum of the peak areas will not always equal 100%

	Peak position (cm^{-1}) and area (%)				
	AP	300 °C	400 °C	500 °C	600 °C
$\text{OH}_L/\text{PO}_4 \nu_4$					
1A	626 (8%)	627 (8%)	623 (7%)	619 (7%)	N/A
2A	603 (28%)	604 (28%)	604 (28%)	604 (28%)	604 (36%)
3A	582 (14%)	582 (16%)	582 (18%)	582 (17%)	580 (17%)
4A	565 (38%)	564 (39%)	564 (38%)	564 (43%)	564 (43%)
5A	546 (12%)	546 (9%)	547 (8%)	541 (5%)	544 (5%)
$\text{CO}_3 \nu_3$					
1B	1523 (13%)	1529 (12%)	1523 (15%)	1539 (9%)	1529 (19%)
2B	1492 (19%)	1497 (17%)	1492 (18%)	1498 (25%)	1495 (10%)
3B	1455 (23%)	1465 (19%)	1455 (16%)	1479 (6%)	1472 (12%)
4B	1424 (12%)	1442 (7%)	1424 (4%)	1446 (14%)	1448 (5%)
5B	1401 (27%)	1410 (35%)	1401 (37%)	1410 (39%)	1418 (41%)
6B	1372 (7%)	1377 (10%)	1372 (11%)	1368 (6%)	1379 (12%)
$\text{CO}_3 \nu_2$					
1C	876 (41%)	877 (29%)	878 (29%)	879 (23%)	880 (23.5%)
2C	868 (51%)	869 (60%)	871 (26%)	872 (29%)	871 (50%)
3C	N/A	N/A	866 (35%)	866 (37%)	865 (27.5%)
4C	858 (8%)	858 (10%)	857 (9%)	857 (11%)	857 (8%)

(Table 4 and the example for 500 °C in Fig. 10C). Increasing the temperature from 300 to 600 °C did, however, affect the OH_L vibration (peak 1A) which shifted to lower frequency and eventually could not be fitted in the spectrum from the 600 °C treatment. The peak at 546 cm^{-1} (peak 5A) in the as-prepared sample, assigned to HPO_4 , also decreased in intensity as the temperature increased, which would be consistent with loss

of hydroxyl groups on heating. Peak deconvolution of the carbonate ν_3 region of the heated samples showed some clear differences from the as-prepared samples, Table 4 and graphically for the sample heated at 500 °C in Fig. 10A; this also used a six-peak fit (peaks 1B–6B). The main peaks assigned to A-type carbonate substitution in the as-prepared samples, at 1523 and 1492 cm^{-1} in the $x = 2.5$ sample (peaks 1B and 2B), remain in all heated samples with comparable contributions (by % peak area) to the total ν_3 region. The peak at $1401\text{--}1410 \text{ cm}^{-1}$ (peak 5B) that was assigned to B-type substitution in the as-prepared samples was again a dominant peak, but appeared to shift considerably to 1418 cm^{-1} after heating at 600 °C. A peak at $1415\text{--}1417 \text{ cm}^{-1}$ (peak 4B) has been assigned to B-type substitution in many studies, so the similarity in peak area of the peak in the $1401\text{--}1418 \text{ cm}^{-1}$ region supports that this is a shift as a result of heating to 600 °C. The peak at approximately $1446\text{--}1455 \text{ cm}^{-1}$ (peak 3B) that appears in as-prepared samples decreased in contribution with heating. For samples heated at 500–600 °C a peak at $1472\text{--}1479 \text{ cm}^{-1}$ appeared which is commonly reported in carbonate-substituted apatites and is assigned to B-type carbonate. This peak increased in relative peak area at the expense of the decreasing peak at $1446\text{--}1465 \text{ cm}^{-1}$. While the resolution of the ν_3 region improves with heating the total band is very broad and in this region, where different studies have assigned vibrations in the region $1445\text{--}1455 \text{ cm}^{-1}$ to both A and B-type substitution, care should be taken to assign the different deconvoluted peaks. What is clear is that heating the as-prepared $x = 2.5$ sample has resulted in some structural change with changes in peak position and relative intensity in the $1400\text{--}1480 \text{ cm}^{-1}$ region. From analysis of these heated samples the vibration at $1368\text{--}1379 \text{ cm}^{-1}$ (peak 6B) is not likely due to carbonate in a water environment, as proposed by Káflak *et al.*;⁴⁹ the relative peak area % increased slightly after heating in CO_2 and the water bending vibration at approximately 1640 cm^{-1} that was visible in



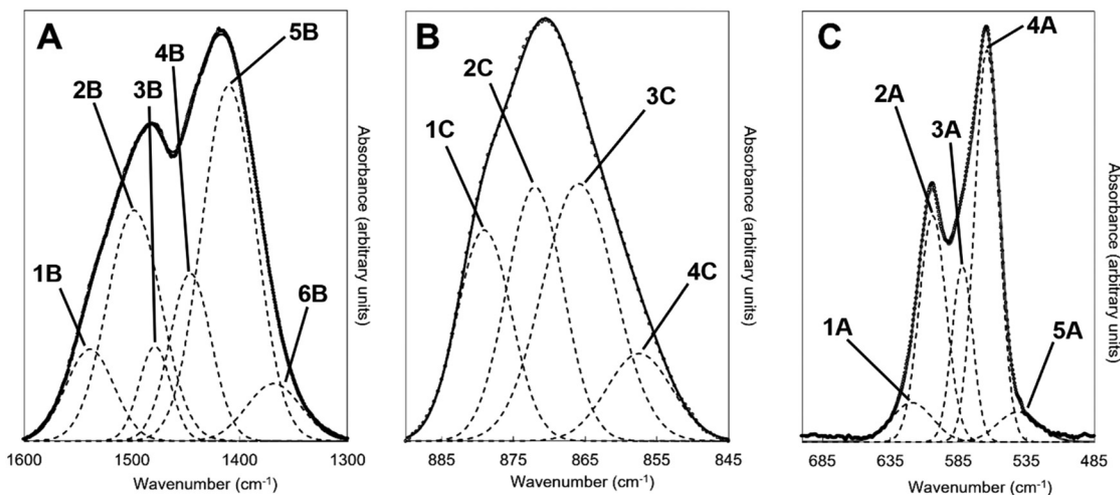


Fig. 10 Deconvolution of the (A) $\text{CO}_3 \nu_3$, (B) $\text{CO}_3 \nu_2$ and (C) $\text{PO}_4 \nu_4$ regions of the FTIR spectra of the $x = 2.5$, Series I potassium–carbonate co-substituted hydroxyapatite sample heated to 500°C in CO_2 , showing the various vibrations that contribute to the observed spectra; y-axis is an arbitrary scale. Dashed lines correspond to the individual fitted peaks and the solid line is the corresponding model spectra, with the raw data presented as isolated data points.

the as-prepared samples disappeared. A band at 1384 and 1386 cm^{-1} was assigned to A-type carbonate in sodium–carbonate substituted apatites.⁵⁰ Heating the as-prepared samples also improved the resolution of the carbonate ν_2 region and the peak deconvolution was successful with 4 peaks, or 3 peaks for the AP and the 300°C heat treatment, Table 4 and the example for 500°C in Fig. 10B. Peak 1C at $880\text{--}876\text{ cm}^{-1}$ can be assigned to A-type carbonate substitution and shows a small shift to higher frequency with increasing heat treatment, and between 300 to 600°C there is only a small decrease in the contribution of this peak to the total peak area of the ν_2 region. A peak (peak 2C) at $872\text{--}868\text{ cm}^{-1}$ is observed at all conditions and between 400 and 600°C a second peak (peak 3C) can be fitted at $856\text{--}855\text{ cm}^{-1}$; these can be assigned to B1- and B2-type carbonate, with a small downshift to lower frequency,⁵¹ and their summed relative peak area shows a small increase with increasing heat treatment temperature. A peak (peak 4C) at $857\text{--}858\text{ cm}^{-1}$ is present before and after all of the heat treatments and represents about $8\text{--}11\%$ of the total peak area. The presence of this peak somewhat mirrors the peak at $1368\text{--}1379\text{ cm}^{-1}$ in the carbonate ν_2 region, and this could be attributed to a second A2-type carbonate band. It is known that carbonate environments in sodium-bearing carbonate substituted hydroxyapatites are influenced by the presence of the monovalent sodium ions, so the presence of potassium in the materials described here may also have a local influence on the carbonate environments.

Estimation of the carbonate content from the FTIR data of the as-prepared compositions and the heat-treated $x = 2.5$ samples was performed and compared to the carbonate content (wt%) measured from combustion analysis data, Tables 1 and 2. A number of approaches have been reported in the literature for quantifying the carbonate content in apatites from the FTIR data, including determining carbonate to phosphate ratios from peak intensities or from peak areas, using isolated peaks or specific regions, and also using ν_2 or ν_3 carbonate regions

and ν_3/ν_1 or ν_4 phosphate regions of the spectra.⁶⁹ As the deconvolution of the carbonate ν_2 region of the as-prepared samples was limited, and the poorly defined and overlapping ν_3/ν_1 phosphate regions results in assumptions being made, the carbonate to phosphate ratios determined were based on (i) the total ν_3 carbonate peak area divided by the total ν_4 phosphate peak area excluding the OH_L peak, and (ii) the peak intensity ratio (PIR) of the carbonate band at $\sim 1415\text{ cm}^{-1}$ (in practise $1401\text{--}1418\text{ cm}^{-1}$) divided by the peak intensity of the ν_4 phosphate band at $\sim 604\text{ cm}^{-1}$.⁴⁷ The measured carbonate contents (wt%) by combustion analysis of all the as-prepared and heated samples were then plotted against the peak area ratios of the $\text{CO}_3 \nu_3$ and $\text{PO}_4 \nu_4$ regions, Fig. 11A, and the peak intensity ratios of the $\text{CO}_3 \nu_3$ vibration at approximately 1410 cm^{-1} and the $\text{PO}_4 \nu_4$ vibration at approximately 603 cm^{-1} , Fig. 11B, with linear fits applied to the data. Both methods provide good linear fits to the data (R^2 values of 0.978 and 0.939 , respectively), even though this combines data using two different potassium carbonate reactants and also as-prepared and heat treated samples. This confirms that the total carbonate content of samples increases with both increasing substitution design level, and with increased temperature of heat treatment. In terms of the distribution of carbonate ions over hydroxyl and phosphate sites, so A-type and B-type, the ratio of the peak areas assigned in the ν_2 region is commonly used to describe a B/A ratio, with this being extended when including duplicate bands (B1 and B2, and A1 and A2) have been assigned.⁵¹ If the peak at 857 cm^{-1} is not included, the $(\text{B1} + \text{B2})/\text{A1}$ peak area ratio, based on the assignment described above, shows an increase with increasing heat treatment temperature, from 1.2 , 2.1 , 2.1 , 2.9 and 2.9 for AP, 300 , 400 , 500 and 600°C (including the 857 cm^{-1} peak, so the $(\text{B1} + \text{B2})/(\text{A1} + \text{A2})$ ratio, provides values of 1.0 , 1.5 , 1.6 , 1.9 and 2.2). The increase in carbonate content of the $x = 2.5$ composition with increasing temperature is therefore associated with a greater amount of



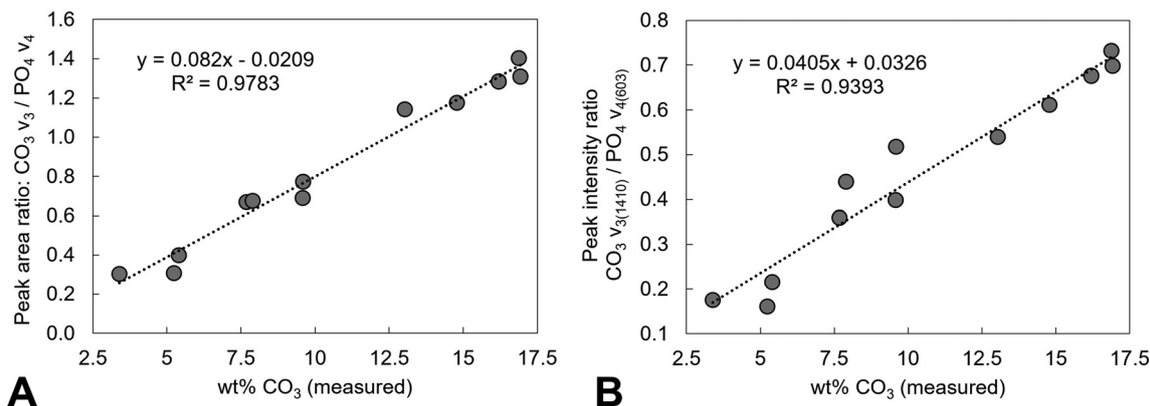


Fig. 11 Analysis of the FTIR data of samples of potassium-carbonate co-substituted hydroxyapatites. The measured carbonate contents (wt%) by combustion analysis of all the as-prepared and heated samples from this study were plotted against (A) peak area ratios of the $\text{CO}_3 \nu_3$ and $\text{PO}_4 \nu_4$ regions (excluding any contribution from an OH vibration at approximately 630 cm^{-1}) and (B) peak intensity ratios of the $\text{CO}_3 \nu_3$ vibration at approximately 1410 cm^{-1} and the $\text{PO}_4 \nu_4$ vibration at approximately 603 cm^{-1} , with linear fits applied to the data.

that carbonate locating on the B-site. The increase in both unit cell parameters of the heated samples with increasing temperature was supportive of addition of carbonate ions in both hydroxyl and phosphate sites, but the unit cell parameter changes could also be associated with the reduction in the hydroxyl and HPO_4 peak intensities observed in the FTIR data, and/or rearrangement of the planar carbonate groups on their respective sites, as the samples are heated. The heated samples do not significantly improve the crystallinity of the samples, with temperatures above $750 \text{ }^\circ\text{C}$ typically required to achieve this, which limits the analysis of the structure by Rietveld refinement of the diffraction data.

4. Conclusions

K- CO_3 co-substituted apatite materials with potassium and carbonate contents that ranged from approximately 0.4–0.9 wt% and 3.4–13.0 wt% respectively were prepared by aqueous precipitation reaction between $\text{Ca}(\text{OH})_2$, H_3PO_4 and either K_2CO_3 or KHCO_3 . A subsequent heat treatment in dry CO_2 at $600 \text{ }^\circ\text{C}$ allowed for a potassium-carbonate co-substituted apatite containing approximately 16.9 wt% CO_3^{2-} to be prepared, believed to be amongst the largest carbonate contents that have been reported for such a material to date. Dissolution experiments showed that the solubility of these materials increased with increasing potassium/carbonate substitution. Whilst this work demonstrates that K- CO_3 co-substituted apatites with high levels of carbonate incorporation can be prepared using simple, room temperature, aqueous precipitation reactions with starting reagents unlikely to pose significant environmental risks, comprehensive testing of these materials in prospective applications (e.g. as solid fertilisers) would be required before they can be considered a viable CO_2 utilisation option.

Conflicts of interest

The authors have no conflicts to declare related to this study.

Acknowledgements

The authors would like to acknowledge the University of Aberdeen and the Royal Commission for the Exhibition of 1851 for providing financial support as well as Mr Colin Taylor, Mr John Still and Dr Brian Hutton for their assistance in collecting some of the experimental data presented here.

References

- 1 IPCC, *Climate Change 2014: Synthesis Report. Contribution of Working Groups I, II and III to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change* [Core Writing Team, ed. R. K. Pachauri and L. A. Meyer, IPCC, Geneva, Switzerland, 2014, p. 151.
- 2 IPCC, *IPCC Special Report on Carbon Dioxide Capture and Storage. Prepared by Working Group III of the Intergovernmental Panel on Climate Change*, ed. B. Metz, O. Davidson, H. C. de Coninck, M. Loos and L. A. Meyer, Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA, 2005, p. 442.
- 3 M. Aresta and A. Dibenedetto, Utilisation of CO_2 as a Chemical Feedstock: Opportunities and Challenges, *Dalton Trans.*, 2007, 2975–2992.
- 4 E. I. Koysoumpa, C. Bergins and E. Kakaras, The CO_2 Economy: Review of CO_2 Capture and Reuse Technologies, *J. Supercrit. Fluids*, 2018, **132**, 3–16.
- 5 J. Morrison, G. Jauffret, J. L. Galvez-Martos and F. P. Glasser, Magnesium-Based Cements for CO_2 Capture and Utilisation, *Cem. Concr. Res.*, 2016, **85**, 183–191.
- 6 R. Muthuraj and T. Mekonnen, Recent Progress in Carbon Dioxide (CO_2) as Feedstock for Sustainable Materials Development: Co-Polymers and Polymer Blends, *Polymer*, 2018, **145**, 348–373.
- 7 M. Tamura, K. Ito, M. Honda, Y. Nakagawa, H. Sugimoto and K. Tomishige, Direct Copolymerization of CO_2 and Diols, *Sci. Rep.*, 2016, **6**, 24038.



- 8 R. Z. LeGeros, O. R. Trautz, E. Klein and J. P. Legeros, Two Types of Carbonate Substitution in the Apatite Structure, *Experientia*, 1969, **25**(1), 5–7.
- 9 I. R. Gibson and W. Bonfield, Novel Synthesis and Characterization of an AB-type Carbonate-Substituted Hydroxyapatite, *J. Biomed. Mater. Res.*, 2002, **59**(4), 697–708.
- 10 D. A. Nowicki, J. M. Skakle and I. R. Gibson, Nano-scale Hydroxyapatite Compositions for the Utilization of CO₂ Recovered Using Post-combustion Carbon Capture, *J. Mater. Chem. A*, 2018, **6**(13), 5367–5377.
- 11 J. A. Stephen, C. Pace, J. M. S. Skakle and I. R. Gibson, Comparison of Carbonate Hydroxyapatite With and Without Sodium Co-Substitution, *Key Eng. Mater.*, 2007, **330-332** I, 19–22.
- 12 M. Vignoles, G. Bonel and R. A. Young, Occurrence of nitrogenous species in precipitated B-type carbonated hydroxyapatites, *Calcif. Tissue Int.*, 1987, **40**(2), 64–70.
- 13 J. Whyte, D. Hadden, I. Gibson and J. Skakle, Synthesis and Stability of Potassium/Carbonate Co-Substituted Hydroxyapatites, *Key Eng. Mater.*, 2008, **361**, 207–210.
- 14 S. Bailliez, A. Nzihou, E. Beche and G. Flamant, Removal of Lead (Pb) by Hydroxyapatite Sorbent, *Process Saf. Environ. Prot.*, 2004, **82**(2), 175–180.
- 15 Y. Feng, J. Gong, G. Zeng, Q. Niu, H. Zhang and C. Niu, *et al.*, Adsorption of Cd(II) and Zn(II) From Aqueous Solutions Using Magnetic Hydroxyapatite Nanoparticles as Adsorbents, *Chem. Eng. J.*, 2010, **162**(2), 487–494.
- 16 L. Luo, Y. Liu, Y. Tan, H. Li, Q. Zhang and K. Li, Room Temperature Gas Sensor Based on Tube-like Hydroxyapatite Modified With Gold Nanoparticles, *J. Cent. South Univ.*, 2016, **23**(1), 18–26.
- 17 R. Liu and L. Rattan, Synthetic Apatite Nanoparticles as a Phosphorus Fertilizer for Soybean (*Glycine Max*), *Sci. Rep.*, 2014, **4**, 5686.
- 18 D. Montalvo, M. J. McLaughlin and F. Degryse, Efficacy of Hydroxyapatite Nanoparticles as Phosphorus Fertilizer in Andisols and Oxisols, *Soil Sci. Soc. Am. J.*, 2015, **79**(2), 551–558.
- 19 B. Singh, S. Kumar, B. Basu and R. Gupta, Enhanced Ionic Conduction in Hydroxyapatites, *Mater. Lett.*, 2013, **95**, 100–102.
- 20 B. R. De Vasconcelos, L. Zhao, P. Sharrock, A. Nzihou and D. P. Minh, Catalytic Transformation of Carbon Dioxide and Methane Into Syngas Over Ruthenium and Platinum Supported Hydroxyapatites, *Appl. Surf. Sci.*, 2016, **390**, 141–156.
- 21 Z. Boukha, M. Kacimi, M. F. R. Pereira, J. L. Faria, J. L. Figueiredo and M. Ziyad, Methane Dry Reforming on Ni Loaded Hydroxyapatite and Fluoroapatite, *Appl. Catal., A*, 2007, **317**(2), 299–309.
- 22 L. Xiong, P. Wang, M. N. Hunter and P. M. Kopitke, Bioavailability and Movement of Hydroxyapatite Nanoparticles (HA-NPs) Applied as a Phosphorus Fertiliser in Soils, *Environ. Sci.: Nano*, 2018, **5**(12), 2888–2898.
- 23 E. Landi, S. Riccobelli, N. Sangiorgi, A. Sanson, F. Doghieri and F. Miccio, Porous Apatites as Novel High Temperature Sorbents for Carbon Dioxide, *Chem. Eng. J.*, 2014, **254**, 586–596.
- 24 O. H. Ojeda-Niño, C. Blanco and C. E. Daza, High Temperature CO₂ Capture of Hydroxyapatite Extracted From Tilapia Scales, *Univ. Sci.*, 2017, **22**(3), 215–236.
- 25 D. A. Nowicki, J. M. S. Skakle and I. R. Gibson, Maximising Carbonate Content in Sodium-Carbonate Co-substituted Hydroxyapatites Prepared by Aqueous Precipitation Reaction, *J. Solid State Chem.*, 2021, **297**, 122042.
- 26 C. Zörb, M. Senbayram and E. Peiter, Potassium in Agriculture– Status and Perspectives, *J. Plant Physiol.*, 2014, **171**(9), 656–669.
- 27 W. T. Pettigrew, Potassium Influences on Yield and Quality Production for Maize, Wheat, Soybean and Cotton, *Physiol. Plant.*, 2008, **133**(4), 670–681.
- 28 L. V. Kochian, Plant Nutrition: Rooting for More Phosphorus, *Nature*, 2012, **488**(7412), 466–467.
- 29 R. M. Verbeeck, E. A. De Maeyer and F. C. Driessens, Stoichiometry of Potassium-Containing and Carbonate-Containing Apatites Synthesized by Solid State Reactions, *Inorg. Chem.*, 1995, **34**(8), 2084–2088.
- 30 T. I. Ivanova, O. V. Frank-Kamenetskaya and A. B. Kol'tsov, Synthesis, Crystal Structure and Thermal Decomposition of Potassium-Doped Carbonated Hydroxyapatite, *Z. Kristallogr.*, 2004, **219**(8), 479–486.
- 31 E. A. P. De Maeyer, R. M. H. Verbeeck and I. Y. Pieters, Effect of K⁺ on the Stoichiometry of Carbonated Hydroxyapatite Obtained by the Hydrolysis of Monetite, *Inorg. Chem.*, 1996, **35**(4), 857–863.
- 32 E. G. Nordström and K. H. Karlsson, Chemical Characterization of a Potassium Hydroxyapatite Prepared by Soaking in Potassium Chloride and Carbonate Solutions, *Bio-Med. Mater. Eng.*, 1992, **2**(4), 185–189.
- 33 M. Akao, H. Aoki and K. Kato, Mechanical Properties of Sintered Hydroxyapatite for Prosthetic Applications, *J. Mater. Sci.*, 1981, **16**(3), 809–812.
- 34 T. Degen, M. Sadki, E. Bron, U. König and G. Nénert, The HighScore Suite, *Powder Diffr.*, 2014, **29**(S2), S13–S18.
- 35 K. Sudarsanan and R. A. Young, Significant Precision in Crystal Structure Details: Holly Springs Hydroxyapatite, *Acta Crystallogr., Sect. B: Struct. Crystallogr. Cryst. Chem.*, 1969, **B25**, 1534–1543.
- 36 J. S. O. Evans and I. R. Evans, Structure Analysis from Powder Diffraction Data: Rietveld Refinement in Excel, *J. Chem. Educ.*, 2021, **98**(2), 495–505.
- 37 T. Mano, K. Akita, N. Fukuda, K. Kamada, N. Kurio and K. Ishikawa, *et al.*, Histological Comparison of Three Apatitic Bone Substitutes with Different Carbonate Contents in Alveolar Bone Defects in a Beagle Mandible with Simultaneous Implant Installation, *J. Biomed. Mater. Res., Part B*, 2020, **108**(4), 1450–1459.
- 38 A. Ito, Y. Sogo, A. Yamazaki, M. Aizawa, A. Osaka and S. Hayakawa, *et al.*, Interlaboratory Studies on In Vitro Test Methods for Estimating In Vivo Resorption of Calcium Phosphate Ceramics, *Acta Biomater.*, 2015, **25**, 347–355.
- 39 H. V. Connerty and A. R. Briggs, Determination of Serum Calcium by Means of Orthocresolphthalein Complexone, *Am. J. Clin. Pathol.*, 1966, **45**, 290–296.



- 40 ICDD. PDF Card No. 9-432. Newton Square, Pennsylvania, USA.
- 41 M. Erbdinger, X. Ni and P. J. Halling, Kinetics of Enzymatic Solid-to-Solid Peptide Synthesis: Synthesis of Z-Aspartame and Control of Acid-Base Conditions by Using Inorganic Salts, *Biotechnol. Bioeng.*, 2000, **72**, 69–76.
- 42 S. N. Danilchenko, A. V. Koropov, I. Y. Protsenko, B. Sulkiocleff and L. F. Sukhodub, Thermal Behavior of Biogenic Apatite Crystals in Bone: An X-ray Diffraction Study, *Cryst. Res. Technol.*, 2006, **41**(3), 268–275.
- 43 M. Sadat-Shojai, M. Khorasani, E. Dinpanah-Khoshdargi and A. Jamshidi, Synthesis Methods for Nanosized Hydroxyapatite With Diverse Structures, *Acta Biomater.*, 2013, **9**(8), 7591–7621.
- 44 Y. Doi, Y. Moriwaki, T. Aoba, M. Okazaki, J. Takahashi and K. Joshin, Carbonate Apatites from Aqueous and Non-Aqueous Media Studied by ESR, IR, and X-ray Diffraction: Effect of NH₄ Ions on Crystallographic Parameters, *J. Dent. Res.*, 1982, **61**(2), 429–434.
- 45 J. Colt, *Dissolved Gas Concentration in Water*, Elsevier, 2nd edn, 2012.
- 46 S. Kannan, J. Ventura and J. Ferreira, Synthesis and Thermal Stability of Potassium Substituted Hydroxyapatites and Hydroxyapatite/ β -Tricalciumphosphate Mixtures, *Ceram. Int.*, 2007, **33**(8), 1489–1494.
- 47 G. Dal Sasso, Y. Asscher, I. Angelini, L. Nodari and G. Artioli, A Universal Curve of Apatite Crystallinity for the Assessment of Bone Integrity and Preservation, *Sci. Rep.*, 2018, **8**(1), 12025.
- 48 J. Reyes-Gasga, E. L. Martínez-Piñeiro, G. Rodríguez-Álvarez, G. E. Tiznado-Orozco, R. García-García and E. F. Brès, XRD and FTIR Crystallinity Indices in Sound Human Tooth Enamel and Synthetic Hydroxyapatite, *Mater. Sci. Eng., C*, 2013, **33**(8), 4568–4574.
- 49 A. Kaflak, A. Ślósarczyk and W. Kolodziejcki, A Comparative Study of Carbonate Bands from Nanocrystalline Carbonated Hydroxyapatites Using FT-IR Spectroscopy in the Transmission and Photoacoustic Modes, *J. Mol. Struct.*, 2011, **997**(1–3), 7–14.
- 50 C. H. Yoder, M. M. Bollmeyer, K. R. Stepien and R. N. Dudrick, The Effect of Incorporated Carbonate and Sodium on the IR Spectra of A- and AB-type Carbonated Apatites, *Am. Mineral.*, 2019, **104**(6), 869–877.
- 51 M. E. Fleet, Infrared Spectra of Carbonate Apatites: ν_2 -Region Bands, *Biomaterials*, 2009, **30**(8), 1473–1481.
- 52 H. Madupalli, B. Pavan and M. M. J. Tecklenburg, Carbonate Substitution in the Mineral Component of Bone: Discriminating the Structural Changes, Simultaneously Imposed by Carbonate in A and B Sites of Apatite, *J. Solid State Chem.*, 2017, **255**, 27–35.
- 53 C. Rey, M. Shimizu, B. Collins and M. J. Glimcher, Resolution-Enhanced Fourier Transform Infrared Spectroscopy Study of the Environment of Phosphate Ions in the Early Deposits of a Solid Phase of Calcium-Phosphate in Bone and Enamel, and Their Evolution with Age. I: Investigations in the ν_4 PO₄ domain, *Calcif. Tissue Int.*, 1990, **46**(6), 384–394.
- 54 C. H. Yoder, N. T. Landes, L. K. Tran, A. K. Smith and J. D. Pasteris, The Relative Stabilities of A- and B-type Carbonate Substitution in Apatites Synthesized in Aqueous Solution, *Mineral. Mag.*, 2016, **80**(6), 977–983.
- 55 A. Antonakos, E. Liarokapis and T. Leventouri, Micro-Raman and FTIR Studies of Synthetic and Natural Apatites, *Biomaterials*, 2007, **28**(19), 3043–3054.
- 56 M. E. Fleet and X. Liu, Coupled Substitution of Type A and B Carbonate in Sodium-Bearing Apatite, *Biomaterials*, 2007, **28**(6), 916–926.
- 57 D. A. Nowicki, J. M. Skakle and I. R. Gibson, Faster Synthesis of A-type Carbonated Hydroxyapatite Powders Prepared by High-Temperature Reaction, *Adv. Powder Technol.*, 2020, **31**(8), 3318–3327.
- 58 M. E. Fleet and X. Liu, Location of Type B Carbonate Ion in Type A–B Carbonate Apatite Synthesized at High Pressure, *J. Solid State Chem.*, 2004, **177**(9), 3174–3182.
- 59 N. V. Vagenas, A. Gatsouli and C. G. Kontoyannis, Quantitative Analysis of Synthetic Calcium Carbonate Polymorphs Using FT-IR Spectroscopy, *Talanta*, 2003, **59**(4), 831–836.
- 60 J. Barralet, J. Knowles, S. Best and W. Bonfield, Thermal Decomposition of Synthesised Carbonate Hydroxyapatite, *J. Mater. Sci.: Mater. Med.*, 2002, **13**(6), 529–533.
- 61 E. Landi, A. Tampieri, G. Celotti and S. Sprio, Densification Behaviour and Mechanisms of Synthetic Hydroxyapatites, *J. Eur. Ceram. Soc.*, 2000, **20**(14–15), 2377–2387.
- 62 C. Liao, F. Lin, K. Chen and J. Sun, Thermal Decomposition and Reconstitution of Hydroxyapatite in Air Atmosphere, *Biomaterials*, 1999, **20**, 1807–1813.
- 63 J. P. Lafon, E. Champion, D. Bernache-Assollant, R. Gibert and A. M. Danna, Thermal Decomposition of Carbonated Calcium Phosphate Apatites, *J. Therm. Anal. Calorim.*, 2003, **72**, 1127–1134.
- 64 K. Tönsuaadu, K. A. Gross, L. Pluduma and M. Veiderma, A Review on the Thermal Stability of Calcium Apatites, *J. Therm. Anal. Calorim.*, 2012, **110**, 647–659.
- 65 A. R. West, *Solid State Chemistry and its Applications: Student Edition*, John Wiley & Sons, 2nd edn, 2014.
- 66 C. Meneghini, M. C. Dalconi, S. Nuzzo, S. Mobilio and R. H. Wenk, Rietveld Refinement on X-ray Diffraction Patterns of Bioapatite in Human Fetal Bones, *Biophys. J.*, 2003, **84**(3), 2021–2029.
- 67 R. Chang, S. Kim, S. Lee, S. Choi, M. Kim and Y. Park, Calcium Carbonate Precipitation for CO₂ Storage and Utilization: A Review of the Carbonate Crystallization and Polymorphism, *Front. Energy Res.*, 2017, **5**, 17.
- 68 J. McCutcheon, I. Power, J. Shuster, A. Harrison, G. Dipple and G. Southam, Carbon Sequestration in Biogenic Magnesite and Other Magnesium Carbonate Minerals, *Environ. Sci. Technol.*, 2019, **53**, 3225–3237.
- 69 E. A. Taylor and E. Donnelly, Raman and Fourier Transform Infrared Imaging for Characterization of Bone Material Properties, *Bone*, 2020, **139**, 115490.

