

Recycling slaughterhouse waste into fertilizer: how do pyrolysis temperature and biomass additions affect phosphorus availability and chemistry?

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Abstract

BACKGROUND: Pyrolysis of slaughterhouse waste could promote more sustainable phosphorus (P) usage through the development of alternative P fertilizers. This study investigated how pyrolysis temperature (220, 350, 550 and 750 °C), rendering before pyrolysis, and wood or corn biomass additions affect P chemistry in bone char, plant availability, and its potential as P fertilizer.

RESULTS: Linear combination fitting of synchrotron-based X-ray absorption near edge structure spectra demonstrated that higher pyrolysis temperatures decreased the fit with organic P references, but increased the fit with a hydroxyapatite (HA) reference, used as an indicator of high calcium phosphate (CaP) crystallinity. The fit to the HA reference increased from 0% to 69% in bone with meat residue and from 20% to 95% in rendered bone. Biomass additions to the bone with meat residue reduced the fit to the HA reference by 83% for wood and 95% for corn, and additions to rendered bone by 37% for wood. No detectable aromatic P forms were generated by pyrolysis. High CaP crystallinity was correlated with low water-extractable P, but high formic acid-extractable P indicative of high plant availability. Bone char supplied available P which was only 24% lower than Triple Superphosphate fertilizer and two- to five-fold higher than rock phosphate.

CONCLUSION: Pyrolysis temperature and biomass additions can be used to design P fertilizer characteristics of bone char through changing CaP crystallinity that optimize P availability to plants.

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Supporting information may be found in the online version of this article.

Keywords: bone char; biochar; phosphorus; pyrolysis; XANES spectroscopy

INTRODUCTION

Human impact on the phosphorus (P) cycle has turned rock phosphate into a finite resource as it is more rapidly mined than deposited.^{1,2} Despite varying predictions about the depletion time of global P reserves from 50 to 300 years, it is widely accepted that P is a non-renewable resource that requires efficient management.^{3–5} Fertilizer production utilizes 80–90% of the rock phosphate that is mined annually^{5,6} and excessive P fertilizer use has led to pollution of aquatic ecosystems.⁷ Hence, making fertilizer production more sustainable is one of the first steps towards a more efficient P cycle and secured agricultural P supply in future.

One proposed solution is to recover P from organic waste disposal⁸ and convert it into fertilizer. This option includes producing bone char from slaughterhouse waste by means of pyrolysis, the heating of materials under low oxygen conditions. The European Union has banned the use of bone and meat meal in cattle feed preparation due to the risk of spreading bovine spongiform encephalopathy (BSE).⁹ High-temperature pyrolysis may overcome this problem through heat sterilization, as dry heat treatment already at temperatures below pyrolysis temperatures has proven to be an effective method to inactivate BSE prions.¹⁰

Unlike rock phosphate, bone char does not contain toxic metals such as nickel, chrome and cadmium^{11,12} making it a less harmful soil amendment.

Calcium phosphates are the primary constituents of bone. Besides the large mineral structure fraction (60%), bone consists of 40% enamel.¹³ The bone mineral structure is commonly classified as biological apatite. In comparison to geological apatite, such as hydroxyapatite (HA), biological apatite has a smaller crystal size, more carbonate substitutions and a significant OH deficiency, overall resulting in greater solubility.^{14,15}

Pyrolysis modifies the chemical and physical structure of bone. Production conditions including pyrolysis temperature significantly alter the yield and quality of bone char.¹⁶ In addition, the meat:bone ratio can drastically change the end product.¹⁷ The thermal decomposition mainly occurs below 500 °C and ends

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after 750 °C.^{16,18} With increasing pyrolysis temperature, bone char shows higher calcium phosphate (CaP) crystallinity, smaller ash particle size, fewer carboxyl and amide functional groups, more macro-pores and higher metal concentration in comparison to the original feedstock.^{11,12,19} However, the majority of studies about pyrolysis and (co-)gasification of slaughterhouse waste is primarily interested in how production conditions alter its potential as a fuel source²⁰ and optimize tar production.²¹ While there has been an attempt to measure characteristics of the solid product yield,¹² the effect of production parameters on the chemical properties of bone char related to its efficacy as agricultural fertilizer is largely unknown. Although studies have demonstrated the P fertilizer potential of bone char,^{22–25} questions about how pyrolysis temperature and feedstock type change the characteristics of bone char important for the development of effective P fertilizers remain unanswered.

Increasing pyrolysis temperature is assumed to lead to greater P concentrations as shown for animal manures^{26,27} and higher CaP crystallinity, suggesting a decrease in solubility.²⁸ Previous studies have added a carbon source such as coal to the pyrolysis or gasification process of meat and bone meal to prevent stickiness from fatty meat.⁹ Co-pyrolyzing bones with a biomass source could also influence CaP crystal formation and may thereby aid in managing P availability to plants. A concurrent production of biochar from biomass could help making P better available to plants,^{29,30} but no information about co-pyrolysis of biomass with bones has been published.

Therefore, the objectives of this study were (1) to determine how pyrolysis temperature, rendering bone and mixing bone with a biomass source before pyrolysis affect CaP crystallinity; and (2) to determine the relation between these production conditions and the P fertilizer potential of bone char as represented by P solubility and plant availability indicators. We hypothesized that increasing pyrolysis temperature leads to greater P concentrations but higher CaP crystallinity, reducing water-soluble P. The addition of biomass prior to pyrolysis was expected to increase CaP crystallinity.

MATERIALS AND METHODS

Material preparation

Bones with meat residue were collected from Dudley Poultry (Middlesex, NY, USA). Rendered bone meal was purchased from The Espoma Company 1929 (Milville, NJ, USA). Wood chips (80% red maple, 20% sugar maple) were obtained from Robinson Lumber (Owega, NY, USA). Corn stalk and stover came from Cornell Farm Services (Ithaca, NY, USA). Bone with meat residue, wood and corn were oven-dried at 60 °C for 5 days and milled to a particle size < 2 mm. An aliquot of 300 g of each material was individually pyrolyzed at a heating rate of 2.5 °C min⁻¹ and maintained at 220, 350, 550 and 750 °C for 45 min in a 9.93 L stainless steel chamber fitted into a muffle furnace. The chamber was swept with argon gas and had rotating paddles for homogenization. In addition, bone with meat residue mixed with wood or corn and rendered bone mixed with wood at a mass ratio of 1:1 were charred under the same pyrolysis conditions. Char yield was measured.

Samples with pyrolysis temperature of 350 °C and above were ground with mortar and pestle and sieved to < 74 µm particle size for X-ray absorption near edge structure (XANES) spectroscopy and to 74–150 µm particle size to control for texture effects on P solubility in the chemical analyses. Bones and biomass samples dried at 60 and 220 °C were ground by a Thomas Wiley Mill mesh size 60 (<250 µm) for chemical analyses and processed using a

ball grinder for XANES spectroscopy. Possible texture differences of fresh bone or biomass compared to chars may have slightly changed P solubility. To rule out any major effects of specific surface area, a surface analysis using CO₂ (ASAP 2020; Micrometrics Instruments Corporation, Norcross, GA, USA) was performed on those samples showing the greatest changes in P solubility and extractability. Specific surface area was low and showed a slight decrease from 220 °C to 350 °C (see the online supporting material, Table S1).

Rock phosphate (ID, USA) supplied by the Espoma Company 1929 (Milville, NJ, USA) and Greenkeeper's Secret Triple Superphosphate (TSP; T&N, Incorporated, Foristell, MO, USA) were ground to 74–150 µm particle size for chemical analyses.

X-ray absorption near edge structure spectroscopy

Phosphorus K-edge XANES spectroscopy characterization of the char samples was carried out at beam line X19-A of the National Synchrotron Light Source (NSLS) at Brookhaven National Laboratory (Upton, NY, USA) using a monochromator with Si(111) crystals, photon flux of $5 \times 10^{10} \text{ s}^{-1}$ and $2\text{H} \times 1 \text{ V mm}$ beam spot size. A passivated implanted planar silicon detector (Canberra Industries, Meriden, CT, USA) measured the P K-edge fluorescence spectra of the samples in a helium-purged chamber. (See Sato *et al.* for further details on the beam experimental set-up.³¹) Samples were mounted in the center of an acrylic sample holder (diameter of 13 mm, depth of 0.5 mm) and covered with a 5 µm thick polycarbonate Mylar X-ray film. Reference P compounds were obtained from a chemical supplier or synthesized as indicated: hydroxyapatite (HA; Sigma–Aldrich, St Louis, MO, USA), synthesized octacalcium phosphate (OCP) which was verified for its chemical characteristics using X-ray powder diffraction analysis, β -tri-calcium phosphate (TCP; Fluka, St. Louis, MO, USA), aluminium phosphate (AP; Fisher Scientific, Waltham, MA, USA), iron(III) phosphate dihydrate (IPD; Sigma–Aldrich), dibasic calcium phosphate (DCP; Sigma–Aldrich), dicalcium phosphate dihydrate (DCPD; Riedel de Haen, Honeywell, Seelze, Germany), calcium phosphate monobasic (CPM; Fisher Scientific), sodium phosphate dibasic monohydrate (SPDM; Mallinckrodt Baker Inc., XXXXX, XXXXX), phytic acid sodium salt hydrate (PASH; Sigma–Aldrich), 1-naphthyl phosphate (NP; Sigma–Aldrich), diphenyl phosphate (DPP; Sigma–Aldrich) and triphenylphosphine (TPP; Sigma–Aldrich). X-ray energy was calibrated to the P K-edge placing the maximum absorption peak of TCP at 2149 eV as reference energy (E_0). Single scans of all samples were collected at specific energy ranges: 2134–2144 eV (interval step of 0.5 eV), 2144–2164 eV (interval step of 0.1 eV), 2164–2179 eV (interval step of 0.2 eV), 2179–2209 eV (interval step of 1 eV). To increase confidence in results, randomly chosen samples were scanned twice and checked for reproducibility (see the online supporting material, Fig. S1 and Fig. S2).

Chemical analyses

Char, rock phosphate and TSP samples were analyzed for plant-available P with a 2% formic acid extraction at a 1:100 w/v ratio. We did not use the standard AOAC methodology (1 mol L⁻¹ neutral ammonium citrate) to estimate P fertilizer efficacy, since testing with high-ash biochars showed that the 2% formic acid fertilizer extractions had a higher correlation with P uptake by rye-grass (*Lolium multiflorum* Lam.) than 1 mol L⁻¹ neutral ammonium citrate or 2% citric acid.²⁷ This should be taken into consideration when comparing the results of this study to the efficacy of other P fertilizers measured by standard AOAC methods. Orthophosphate content of extracts was analyzed with the

ascorbic acid method.³² As the particle size of the bone char samples was relatively small (74–150 μm), it is important to recognize that fine particles have higher extractability.³³ Water-soluble P, calcium (Ca), magnesium (Mg), potassium (K), iron (Fe), sodium (Na) and sulfur (S) were extracted by shaking char samples in deionized water at a 1:150 solid–solution ratio for 16 h and filtering with 2 V Whatman qualitative filter paper.³⁴ The modified dry-ashing method³⁵ was used to obtain total P, Ca, Mg, K, Fe, Na and S. Elemental concentrations were quantified in the extracts by inductively coupled plasma atomic emission spectrometry (ICP-AES Thermo Jarrell Ash 166 Trace Analyzer; Thermo Jarrell Ash Corporation, Franklin, MA, USA). To determine pH, samples were shaken in water at 1:20 solid–solution ratio for 90 min and analyzed for pH (Orion 3 Star; Thermo Scientific Inc., Beverly, MA, USA).

Data analysis

The program ATHENA of the IFFFIT software package³⁶ was used for data processing and analysis of the XANES spectra. After checking the accuracy of the pre- and post-edge parameters of randomly selected scans, spectra were subjected to automatic background correction. Linear combination fitting was performed over the spectral region from 15 eV below to 30 eV above the P absorption edge. To allow for uncertainty, weights of the standards were not fixed to 1. All 13 standards were used in the analysis. A maximum of four standards were allowed to contribute to a fit. First, the groups of organic, inorganic and CaP standards were separately fitted. Standards that never obtained a weight other than 0 were removed from the analysis. The second round of linear combination fitting used the selected organic, inorganic and CaP standards simultaneously. Fits were compared according to their chi-square values. Fits to the standards were qualitatively interpreted. OCP, a precursor of bone apatite,^{37,38} was used as indicator of low CaP bone crystallinity and HA as indicator of high CaP bone crystallinity. In addition, the HA/OCP ratio provided a measure for the degree of bone crystallinity. Lastly, expected curves of the biomass–bone mixtures were calculated arithmetically based on the ratios of total P contents and spectra of the individual materials. They were compared to the measured spectra by plotting them in the same graph.

Statistical analyses were performed with software package JMP Pro 10.

RESULTS AND DISCUSSION

XANES spectral features of standards

CaP standards were selected according to their solubility: CPM > DCPD > DCP > TCP > OCP > HA.^{28,39} Except for TPP, the white line energy (absorption edge) fell approximately at 2149 eV (Fig. 1) and an oxygen oscillation peak occurred around 2166 eV. The CaP standards, other inorganic standards, and organic P standards were distinctively different from one another. The shoulder around 2151 eV was a unique feature for all CaP standards. This secondary peak sharpened and the absorption edge peak widened for more crystalline CaP references such as HA and OCP, which indicates lower solubility and higher thermodynamic stability.^{40,41} For HA, OCP and TCP, a tertiary sharp peak appeared at 2159 eV. CaP standards with a CaHPO₄ such as DCP, DCPD and CPM also have a more rounded peak at the same energy range. This observation is in line with the P K-edge XANES spectra of Sato *et al.*³¹ and Ingall *et al.*⁴² but not with the data published

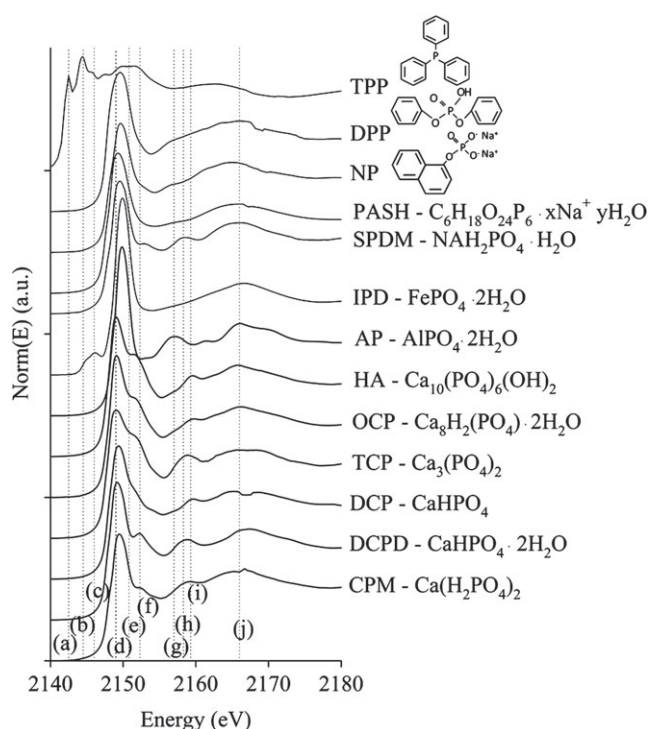


Figure 1. Phosphorus K-edge XANES spectra for P standards species. The dotted lines indicate energy levels that characterize unique spectral features for different P species: (line a) triphenylphosphine (TPP), (line b) TPP, (line c) iron(III) phosphate dehydrate (IPD), (line d) absorption edge, (line e) octacalcium phosphate (OCP) and hydroxyapatite (HA), (line f) calcium phosphate (CaP) species, (line g) aluminium phosphate (AP), (line h) sodium phosphate dibasic (SPDM), (line i) HA and OCP, (line j) oxygen oscillation.

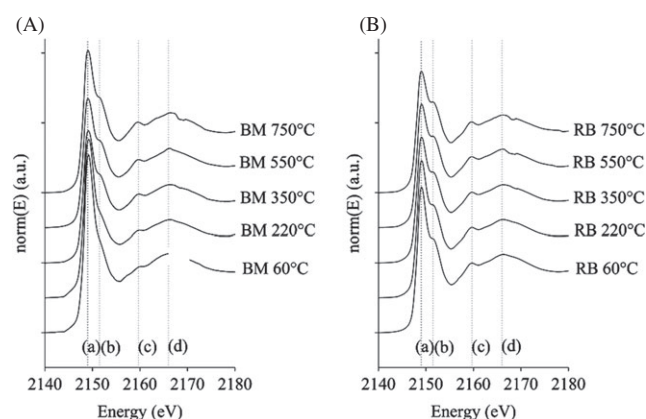


Figure 2. Phosphorus K-edge XANES spectra for bone with meat residue (BM) and rendered bone (RB) at different pyrolysis temperatures. The dotted lines indicate energy levels that characterize unique spectral features for different P species: (line a) absorption edge, (line b) octacalcium phosphate (OCP) and hydroxyapatite (HA), (line c) OCP and HA, (line d) oxygen oscillation.

by Peak *et al.*,⁴⁰ which do not show a tertiary peak at 2159 eV for DCP. This could be attributed to the difference between our sample synthesized by Sigma–Aldrich and a natural DCP sample (monetite) used by Peak *et al.*⁴⁰

IPD was characterized by a pre-absorption edge peak at 2146 eV. AP had a secondary broad peak at 2157 eV. SPDM showed two small peaks in between the absorption edge and oxygen

Table 1. Linear combination fitting results of different feedstock mixtures dried or pyrolyzed at different temperatures

FS ^a	Temp (°C)	OCP (%)	HA (%)	IDP (%)	PASH (%)	DPP (%)	χ^2
BM	60	71.9	–	15.9	22.6	–	7.09
BM	220	59.2	–	11.3	36.4	–	3.13
BM	350	63.1	27.9	2.8	9	–	0.38
BM	550	38.1	53.7	–	9.7	–	0.30
BM	750	33.2	69.3	3.7	–	–	1.26
RB	60	83.4	19.7	2.1	–	–	1.27
RB	220	61.7	39.1	2.6	–	–	0.54
RB	350	17.3	81.1	3.1	–	–	0.07
RB	550	4.2	93.3	4.2	–	–	0.07
RB	750	6.2	94.9	–	–	–	0.05
BW	220	67.3	–	12.1	8.2	23.1	5.13
BW	350	90.7	–	1.3	–	12.7	1.14
BW	550	96.9	–	0.4	–	6.2	0.71
BW	750	89.2	11.6	1.9	–	–	0.36
BC	220	60.7	–	11.2	37.6	–	4.41
BC	350	83.2	–	7.6	14.6	–	2.46
BC	550	99.1	–	5.3	–	–	0.88
BC	750	100	3.8	4.3	–	–	2.01
RBW	220	88.8	11.3	2.5	–	–	0.59
RBW	350	64.1	36.5	1.5	–	–	0.44
RBW	550	26.9	72.3	3	–	–	0.26
RBW	750	33.2	68.5	–	–	–	0.16

^a Feedstock (FS) includes bone with meat residue (BM), rendered bone (RB), bone with meat residue and wood (BW), bone with meat residue and corn (BC), rendered bone and wood (RBW). P species are octacalcium phosphate (OCP), hydroxyapatite (HA), iron(III) phosphate dehydrate (IPD), phytic acid sodium salt hydrate (PASH), diphenyl phosphate (DPP).

oscillation peak: one similar to the CaP shoulder of more soluble CaP compounds at around 2151 eV and one peak at 2158 eV. The AP spectrum was consistent with other studies, while the post-absorption edge spectrum of IPD was smoother than in other studies.^{31,40,42,43}

The main absorption edge peaks of PASH, NP and DDP shifted to the right by 0.2–0.5, which was also reported by Brandes *et al.*⁴⁴ While the shape and width of the absorption edge peak slightly differed among these three organic phosphate standards, they had no other distinctive features. The spectrum of PASH was similar to the scans of other phytic acid species from previous studies.^{44,45} The white line energy peak of TPP occurred at 2144.5 eV, which is still in the pre-edge of all other reference compounds. TPP also showed a secondary peak before its major absorption peak at 2142.5 eV. There was no oxygen oscillation peak because TPP was the only compound with no oxygen bound to P. This standard was included to identify P atoms in heterocycles.

Chemical structures of phosphorus in bone char

In the case of rendered bone and bone with meat residue, increasing pyrolysis temperature led to widening of the absorption edge peak at 2149 eV, sharpening of the CaP shoulder at 2151 eV and the peak characteristic for apatite-like CaP minerals at 2159 eV (Fig. 2), demonstrating an increase in CaP crystallinity. This is consistent with results from X-ray diffractometry.¹¹ At lower temperatures, the spectra of the OCP standard resembled those of rendered

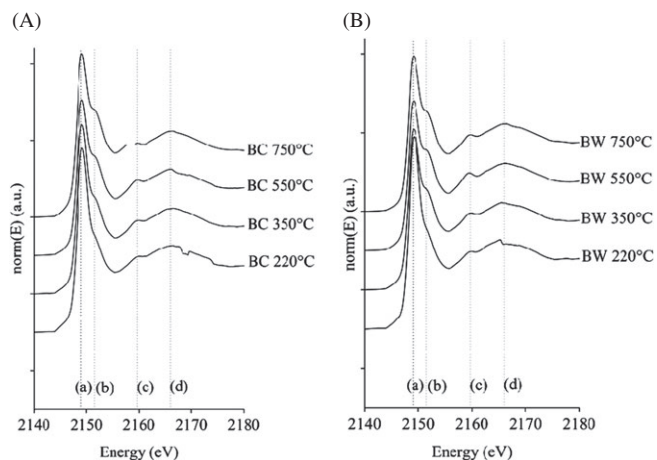


Figure 3. Phosphorus K-edge XANES spectra for mixtures of bone and biomass: (A) bone with meat residue and corn biomass (BC) and (B) bone with meat residue and wood biomass (BW). The vertical dotted lines indicate energy levels that characterize unique spectral features for different P species: (line a) absorption edge, (line b) octacalcium phosphate (OCP) and hydroxyapatite (HA), (line c) OCP and HA, (line d) oxygen oscillation (XANES spectra for pure corn and wood biomass shown in supplementary online Fig. S6).

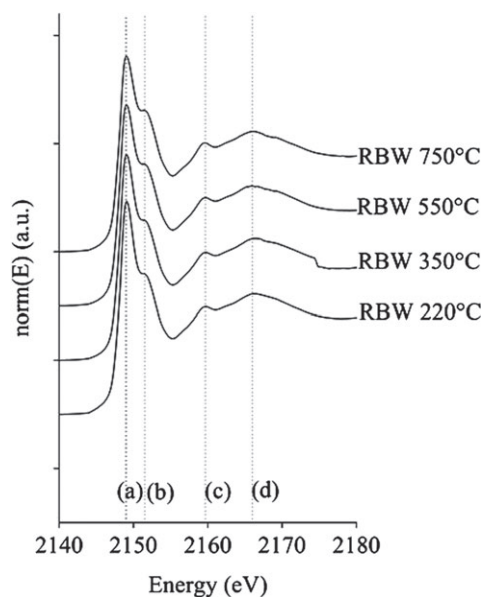


Figure 4. Phosphorus K-edge XANES spectra for mixtures of rendered bone and wood biomass: The vertical dotted lines indicate energy levels that characterize unique spectral features for different P species: (line a) absorption edge, (line b) octacalcium phosphate (OCP) and hydroxyapatite (HA), (line c) OCP and HA, (line d) oxygen oscillation.

bone and bone with meat residue spectra with a higher degree of fit than HA as shown by Fig. 2 and the linear combination fitting results (Table 1). However, with an increase in temperature the P spectral properties of bone with meat residue move from 0% to 69% match with HA and those of rendered bone from 20% to 95% match with HA (Table 1). It is important to note that the linear combination fitting results of OCP and HA should not be interpreted quantitatively, but that they can be used to evaluate overall trends. HA used in this study was synthesized and consequently more crystalline than biological apatite. OCP has a similar chemical structure as HA but also contains water in its crystal lattices.⁴⁶ As

Table 2. Chemical characteristics of the sample: pH, total P, total Ca, formic-P by unit mass and total P, and water-P (means and standard deviation; $n = 3$)

FS ^a	Temp. (°C)	pH	Total-Ca (g Ca kg ⁻¹)	Total-P (g P kg ⁻¹ char)	Formic-P (g P kg ⁻¹ char)	Formic-P (g kg ⁻¹ P) ^b	Water-P (g kg ⁻¹ P) ^b	Mass yield (% initial)
BM	60	6.3 ± 0.0	58.7 ± 6.9	32.6 ± 3.6	28.1 ± 3.1	861 ± 96	156 ± 15	100
BM	220	6.7 ± 0.0	57.3 ± 3.3	31.3 ± 1.6	15.0 ± 3.1	480 ± 100	102 ± 6	90
BM	350	8.4 ± 0.8	151.7 ± 6.5	83.4 ± 3.6	67.5 ± 3.0	810 ± 36	12 ± 1	38
BM	550	10.6 ± 0.0	194.1 ± 3.2	104.1 ± 1.4	89.8 ± 0.9	862 ± 09	29 ± 1	25
BM	750	11.0 ± 0.0	204.1 ± 5.7	109.5 ± 0.3	98.1 ± 1.9	895 ± 18	30 ± 1	26
RB	60	7.3 ± 0.0	183.4 ± 10.5	85.9 ± 4.7	79.1 ± 2.5	922 ± 29	20 ± 2	100
RB ^c	220	6.8 ± 0.0	189.4 ± 6.0	87.9 ± 0.4	85.0 ± 4.0	967 ± 45	20 ± 1	–
RB	350	7.5 ± 0.0	270.6 ± 7.2	127.1 ± 1.3	119.6 ± 7.0	941 ± 55	3 ± ND ^e	68
RB	550	9.2 ± 0.1	307.2 ± 5.4	139.9 ± 4.4	138.2 ± 1.4	988 ± 10	5 ± ND ^e	60
RB	750	10.2 ± 0.1	337.1 ± 8.2	153.2 ± 2.2	147.0 ± 4.7	959 ± 30	3 ± ND ^e	54
BW ^d	60	–	29.9 ± 3.4	16.3 ± 1.8	14.1 ± 1.6	861 ± 96	156 ± 15	100
BW	220	6.5 ± 0.0	27.8 ± 1.5	15.4 ± 0.9	14.7 ± 1.2	951 ± 79	203 ± 10	94
BW	350	8.3 ± 0.1	72.1 ± 2.7	39.7 ± 1.0	30.6 ± 2.1	770 ± 53	15 ± 1	39
BW	550	10.1 ± 0.1	95.4 ± 3.7	53.1 ± 2.5	45.7 ± 2.9	861 ± 54	37 ± 3	30
BW	750	10.5 ± 0.0	106.3 ± 5.1	56.4 ± 1.5	55.1 ± 3.5	977 ± 63	27 ± 2	28
BC ^d	60	–	30.4 ± 3.4	16.6 ± 1.8	14.2 ± 1.6	855 ± 95	163 ± 15	100
BC	220	6.7 ± 0.0	25.4 ± 1.5	14.3 ± 0.6	13.2 ± 1.3	920 ± 88	240 ± 10	101
BC	350	9.6 ± 0.1	69.8 ± 2.4	37.8 ± 1.6	31.3 ± 1.1	829 ± 30	18 ± 1	6
BC	550	10.3 ± 0.0	100.6 ± 2.0	55.7 ± 1.7	50.2 ± 0.3	901 ± 06	44 ± 2	23
BC	750	10.4 ± 0.0	110.1 ± 2.7	58.1 ± 1.0	53.1 ± 2.7	913 ± 47	43 ± 2	17
RBW ^d	60	–	92.3 ± 5.3	43.0 ± 2.3	39.6 ± 1.2	922 ± 29	20 ± 2	100
RBW	220	6.6 ± 0.0	102.1 ± 5.3	47.4 ± 1.6	47.8 ± 0.6	1010 ± 13	34 ± 1	91
RBW	350	7.2 ± 0.1	224.4 ± 4.2	102.4 ± 1.7	102.1 ± 3.4	997 ± 33	6 ± 1	51
RBW	550	8.8 ± 0.0	218.7 ± 3.7	100.9 ± 3.8	101.9 ± 2.9	1010 ± 29	5 ± ND ^e	44
RBW	750	10.1 ± 0.0	258.3 ± 4.5	116.6 ± 2.6	119.0 ± 5.1	1021 ± 44	4 ± ND ^e	41
RP	–	6.6 ± 0.0	224.0 ± 4.2	84.8 ± 1.4	23.7 ± 0.5	279 ± 6	4 ± 1	–
TSP	–	3.2 ± ND ^e	161.0 ± 5.2	205.3 ± 1.8	194.0 ± 4.1	945 ± 20	100 ± ND ^e	–

^a Feedstock (FS) include bone with meat residue (BM), rendered bone (RB), bone with meat residue and wood biomass (BW), bone with meat residue and corn biomass (BC), rendered bone and wood biomass (RBW), rock phosphate (RP), and Triple Superphosphate (TSP) fertilizer.

^b Formic-P and water-P are expressed in g kg⁻¹ total P.

^c Mass yield of RB at 220 °C was not recorded.

^d Nutrient characteristics of these samples were calculated as an arithmetic mean from the two materials it was composed of and pH content was not measured.

^e No standard deviation detected (ND)

a consequence, disordered and less crystalline bone apatite could show more resemblance to the OCP reference. Therefore, we will interpret a high OCP fit as bone apatite with low crystallinity and a high HA fit as bone apatite with high crystallinity (Fig. 3).

The rendered bone spectra showed a 37% higher fit with HA than bone with meat residue produced at 750 °C, but even at lower temperatures spectra of rendered bone char better resembled those of the HA reference (Table 1). Rendering removes fatty meat residue, leaving a purer source of CaP mineral showing a higher resemblance with HA even without charring. Moreover, less contamination of fatty meat could facilitate CaP crystal formation at a faster rate during pyrolysis (Fig. 4).

Biomass additions reduced CaP crystal formation during bone pyrolysis. At 750 °C, corn reduced P resembling HA in bone with meat residue by 95%, and wood decreased HA-type structures in bone with meat residue by 84% and in rendered bone by 28% (Table 1). There are several mechanisms that could explain the reduction of CaP crystal formation. First, the addition of corn or wood biomass resulted in an increase of carbon and organic acids. Adsorption and chemical bonding of these molecules onto CaP crystal surfaces could have blocked sites acting as nuclei

for continued crystal growth.^{37,47,48} Second, metal ions from the biomass sources could have decreased crystal formation through impeding the growth of nuclei or poisoning the substrate resulting in heterogeneous nucleation.⁴⁴ Strong metal inhibitors of CaP crystallization are Cu²⁺, Zn²⁺ and Mg²⁺,^{49,50} none of which were present in high amounts. Cu²⁺ and Zn²⁺ were below the detection limit of the ICP-AES and higher Mg concentration did not correlate with reduced crystal growth (see the online supporting material, Figs. S3–S5). Future studies should focus on identifying the mechanism responsible for reduced crystal growth.

The organic P standards PASH and DPP constituted a small part of the spectra from bone with meat residue alone and mixed with wood or corn biomass, but not of the spectra from rendered bones and only at low pyrolysis temperatures (Table 1). This indicates that rendering bone removes most organic P, being part of the fatty meat residue, and that with increasing pyrolysis temperature organic P seems to be broken down and transformed into inorganic P. NP and TPP did not show fits with any of the samples and DPP only matched with spectra of the original feedstock (Table 1). These results suggest that no heterocyclic P forms were generated through pyrolysis.

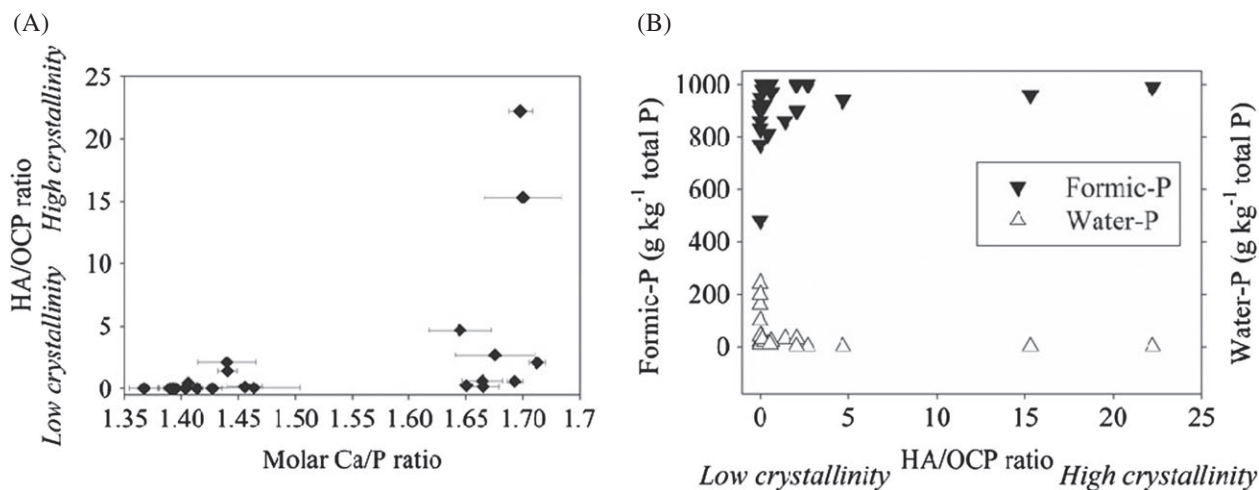


Figure 5. Calcium phosphate (CaP) crystallinity relationships: (A) relation between two indicators of CaP crystallinity molar calcium/phosphorus (Ca/P) ratio and hydroxyapatite/octacalcium phosphate (HA/OCP ratio), (B) effect of HA/OCP ratio as indicator of CaP crystallinity on water-soluble and formic acid extractable P (individual Ca and P concentrations shown in supplementary Fig. S3).

Phosphorus concentration, solubility and availability

All samples demonstrated total P enrichment with increasing pyrolysis temperature (Table 2). P vaporization and particulate transfer to air is generally low in comparison to N, S, C and K,^{51–54} resulting in a greater P concentration in bone chars produced at higher temperatures. Ca was enriched to a higher degree than P. This increased the molar Ca/P ratio at high pyrolysis temperatures (Table 2). Other studies have obtained a similar relationship between temperature and Ca/P ratio and have found that an increase in Ca/P ratios is coupled with higher HA contents,⁵⁵ suggesting higher CaP crystallinity. These results are in line with our XANES data. A threshold at a molar Ca/P ratio of 1.55 seems to divide the bone fertilizers in two groups with higher versus lower CaP crystallinity as represented by HA/OCP ratio (Fig. 5A).

Formic-P (g kg⁻¹ total P) also increased with temperature (Table 2), suggesting that bone chars produced at higher pyrolysis temperature provided larger proportions of plant-available P. However, the opposite was true for water-P (g kg⁻¹ total P), which decreased with an increase in production temperature (Table 2). Even at lower temperatures, formic-P was substantially higher than water-P (Table 2). The water-soluble P fraction may be an under-estimate of the amount of P available for plants, while the formic acid-extractable P fraction may be an over-estimate of immediate available P.⁵⁶ Overall, amounts of formic-P in bone char reached 147.0 ± 4.7 g kg⁻¹. This P concentration is five times greater than in Idaho rock phosphate, 2.5 times greater than in GAFSA rock phosphate²¹ and only 24% lower than the formic-P in TSP fertilizer measured as 194.0 ± 4.1 g kg⁻¹ (Table 2). It should be recognized that possible changes in particle size may have influenced P extractability³⁴ with increasing pyrolysis temperatures, even though a systematic change in size cannot be confirmed and surface area rather decreased.

HA/OCP ratios showed a significant negative correlation with water-P and a significant positive correlation with formic-P (Table 3), resulting in lower water-P and higher formic-P (Fig. 5B). Previous studies confirm that more crystalline CaP such as HA are more stable and, therefore, have lower solubility in water than OCP.^{57–59} On the other hand, HA has a higher pH and might be more acid-soluble than other CaP structures. This may explain the increase in formic-P with temperature. As wood and corn additions led to lower crystallinity, water-P increased and formic-P

Table 3. Spearman's correlations of hydroxyapatite/octacalcium phosphate (HA/OCP) ratio as indicator of calcium phosphate (CaP) crystallinity, formic-P (% of total-P), water-P (% of total-P), pH and pyrolysis temperature

Variable	By variable	Spearman's ρ	Prob > $ \rho $
HA/OCP	Formic-P	0.55	0.008
HA/OCP	Water-P	-0.77	< 0.0001
HA/OCP	pH	0.48	0.024
Temperature	HA/OCP	0.45	0.037
Temperature	Formic-P	0.19	0.403
Temperature	Water-P	-0.34	0.116

decreased. Rendering slaughterhouse waste prior to pyrolysis led to higher total P and formic-P, but had lower water-P concentration in the chars (Table 2).

CONCLUSION

P K-edge XANES analysis and chemical P extractions showed that pyrolysis temperature increased CaP crystallinity, total P content and formic-P, but decreased P solubility in water. Organic P forms disappeared during pyrolysis in favor of inorganic P forms and no aromatic P was detectable with the methods used in this study. Rendering slaughterhouse waste before pyrolysis resulted in a purer mineral P source with higher total P contents and higher CaP crystallinity. Mixing wood or corn biomass with bone largely reduced CaP crystal formation during pyrolysis. Hence, all three variables – pyrolysis temperature, rendering and biomass addition – can be used to manipulate P characteristics of a bone-based fertilizer that are desirable for a particular farming system. Effects of varying surface area or particle sizes were not specifically investigated, but may have a significant effect on P availability in addition to changes in chemistry. For site-specific application, recommendations would need to recognize the changes in P dissolution and adsorption depending on soil pH. Future research should therefore test a variety of bone chars in diverse cropping systems and soil types to determine which bone fertilizers perform best in different settings.

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SUPPORTING INFORMATION

Supporting information may be found in the online version of this article.

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