EXPERIMENTAL STUDY ON THE ORIGIN OF CREMATED BONE APATITE CARBON

C M Hüls¹ • H Erlenkeuser • M-J Nadeau • P M Grootes • N Andersen

Leibniz Laboratory for Radiometric Dating and Isotope Research, Christian-Albrechts-University, Kiel, Germany.

ABSTRACT. Bones that have undergone burning at high temperatures (i.e. cremation) no longer contain organic carbon. Lanting et al. (2001) proposed that some of the original structural carbonate, formed during bioapatite formation, survives. This view is based on paired radiocarbon dating of cremated bone apatite and contemporary charcoal. However, stable carbon isotope composition of carbonate in cremated bones is consistently light compared to the untreated material and is closer to the δ^{13} C values seen in C₃ plant material. This raises the question of the origin of carbonate carbon in cremated bone apatite. That is, does the isotope signal reflect an exchange of carbon with the local cremation atmosphere and thus with carbon from the burning fuel, or is it caused by isotopic fractionation during cremation?

To study the changes in carbon isotopes (${}^{14}C$, ${}^{13}C$) of bone apatite during burning up to 800 °C, a modern bovine bone was exposed to a continuous flow of an artificial atmosphere (basically a high-purity O_2/N_2 gas mix) under defined conditions (temperature, gas composition). To simulate the influence of the fuel carbon available under real cremation conditions, fossil CO₂ was added at different concentrations. To yield cremated bone apatite properties similar to archaeological cremated bones, in terms of crystallographic criteria, water vapor had to be added to the atmosphere in the oven. Infrared vibrational spectra reveal large increases in crystal size and loss of carbonate upon cremation. The isotope results indicate an effective carbon exchange between bone apatite carbonate and CO₂ in the combustion gases depending on temperature and CO₂ concentration. ${}^{14}C$ dates on archaeological cremated bone apatite may thus suffer from an old-wood effect. Paired ${}^{13}C$ and ${}^{14}C$ values indicate that in addition to this exchange, isotope fractionation between CO₂ and carbonate, and admixture of carbon from other sources such as possibly collagen or atmospheric CO₂, may play a role in determining the final composition of the apatite carbonate.

INTRODUCTION

Bone is a composite of organic (predominantly collagen) and inorganic (mineral) material. The mineral phase is bioapatite, a calcium phosphate close to natural abiotic apatites in a non-stoichiometric composition. An overall range of bioapatite chemistry is given by Skinner (2005) as (Ca, Na, Mg, K, Sr, Pb,...)₁₀(PO₄, CO₃, SO₄,...)₆(OH, F, Cl, CO₃)₂. Cazalbou et al. (2004) use Ca_{8.3µ1.7} (PO₄)_{4.3}(HPO₄ and CO₃)_{1.7}(OH or 0.5CO₃)_{0.3µ1.7} to emphasize the role of structural carbonate for bioapatite properties, e.g. crystal size, crystal strain, solubility, etc. (where **□** indicates a vacancy, i.e. unfilled site, in the structure). Structural carbonate (CO₃²⁻) can be substituted for the phosphate (PO₄³⁻) or the hydroxyl (OH⁻) (B-type carbonate and A-type carbonate, respectively), where differences in the charge state of the substituent are balanced by empty positions in the crystal lattice. Studies on bone apatite, as well as synthetic apatites, have shown that precipitation at body temperature favors the B-type carbonate substitution (Elliott 2002).

Up to 6 wt% CO_3^{2-} (~1.2 wt% C), which originates from blood bicarbonate generated by energy metabolism in the cells, can be substituted into bone apatite as so-called structural carbonate. Although radiocarbon dating of bone apatite is possible in principle, it has been avoided in the past due to the large potential for ion exchange during bone deposition and postburial degradation (Berger et al. 1964; Tamers and Pearson 1965). Recently, a number of studies have shown that ¹⁴C dating of cremated bones (i.e. bones burnt at temperatures >600 °C) gives apparent apatite ages that are in close agreement with those of contemporaneous organic material (Lanting and Brindley 2000; Lanting et al. 2001; De Mulder and Van Strydonck 2004; Van Strydonck et al. 2005; De Mulder et al. 2007; Olsen et al. 2008).

¹Corresponding author. Email: mhuels@leibniz.uni-kiel.de.

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588 *C M Hüls et al.*

Cremation of bones results in specific changes in apatite properties such as: total loss of organics; substantial loss of carbonate; and a significant increase in crystallinity and crystal size as well as enhanced friability (Shipman et al. 1984; Stiner et al. 1995). The increased crystal size of burnt apatite, from a few tens of nm in fresh bones to >>100 nm in cremated bones (Enzo et al. 2007; Piga et al. 2008), is thought to preserve the remaining structural carbonate (~0.1 wt%) against ion exchange while the bone is buried in the soil (Lanting et al. 2001).

The generally good agreement between burnt apatite and contemporaneous organic material seems to support this assumption. However, burnt apatite δ^{13} C values are in general depleted compared to unburnt bone apatite (<-21‰ and >-17‰, respectively; Bocherens 2002; Lee-Thorp and Sponheimer 2003). For instance, Olsen et al. (2008) reported δ^{13} C values from a single bone with visual and crystallographic features varying from nearly fresh to fully cremated with a large depletion in δ^{13} C in the cremated part. Surovell (2000), in burning apatite from a 10-kyr-old mammoth at 800 °C to remove possible secondary calcite, found it to contain bomb ¹⁴C, thus clearly indicating carbon exchange during combustion. This raises the question what the isotopic composition of the carbonate in cremated bone apatite is indicative of: the original apatite carbonate, carbon exchange, and/or isotopic fractionation during cremation?

To understand the origin of carbon in cremated bone apatite, we have conducted burning experiments with a modern fresh bovine bone under controlled conditions of variable temperature and composition of the burning atmosphere.

MATERIALS AND METHODS

Pieces (0.5–1 cm thick, 1–2 cm wide, and 1 to ~5 cm long) of a fresh bovine bone (cortical bone from femur) were scraped and defleshed using a scalpel, then burnt at temperatures from 500 to 800 °C under different artificial atmospheres in a flow-through burning reactor (quartz tube with 2 burning chambers; see Figure 1, Tables 1 and 2). To investigate the influence of burning fuel carbon, old (i.e. almost dead) CO₂ (Table 2) was admixed into a pure O_2/N_2 mixture (Table 1). In the first set of experiments, the burning atmosphere (O_2/N_2 with and without CO₂) was directed through a bath of ultrapure water at room temperature before entering the oven ("wet" burning). This procedure simulates a burning atmosphere with some water vapor as for natural burning processes. In a second set of experiments, no moisture was added ("dry" burning). The experimental system is leakproof. It is terminated by a demineralized water buffer to prevent backward diffusion of external gases. CO₂ blanks (no bone sample) were tested trapping the gas after the oven. The blank was <0.007 mg C, applying a CO₂-free O₂/N₂-burning gas flowing at 0.1 L/min under common combustion conditions (500/800 °C) for 0.5 hr. This blank was considered insignificant.

After burning, the sample material was allowed to cool down to <100 °C under continuing gas flow before removal from the oven. The sample material was stored in glass containers under Ar gas to prevent contamination with the atmosphere in the long term. Approximately 5–10 mg of sample material was taken from 2 to 3 bone pieces for analysis with Fourier-transform infrared spectroscopy (FTIR) and ~300 mg for X-ray diffractometry (XRD). The pieces were sampled with a dentist tool on the outside and inside and then powdered in a ball mill. FTIR was performed on a Nicolet 308^{TM} in ATR mode (attenuated total reflection, diamond crystal, 400 to 4000 cm⁻¹ wavelength). Changes in crystallinity were estimated using the splitting factor (SF) of the v_4 vibration mode of PO₄^{3–} between 495 and 750 cm⁻¹ (Weiner and Bar-Yosef 1990). Since no difference is observed between SF values from the outside and inside samples, all measurements from the different experiments were pooled. Single XRD analyses were performed on a Bruker D8-advance with a copper cathode (CuK α) with ~300 mg powdered sample material. The crystallinity index (CI) was calcu-



Figure 1 Experimental layout of cremation tests: A) burning atmosphere (O_2 - N_2 mixture, with and without old CO_2 , is directed through ultrapure water into the burning chamber ("wet" combustion); B) burning atmosphere is fed directly into the burning chamber ("dry" combustion). To prevent the outer atmosphere from diffusing in the system, burning gas is discharged through ultrapure water. During the experiment, the gas flow in the system raises the pressure within the system above atmospheric pressure.

Experiment	Temperature	Duration	Weight	Σ gas flow	CO ₂	N ₂	O ₂
nr	(°C)	(hr)	(g)	(mL/min)	(mL/min)	(mL/min)	(mL/min)
Wet combus	stion						
4	800	4.0	6.46	90	0	70	20
11	800	4.4	15.00	70	0	55	15
3	800	3.8	14.99	101	2	79	20
5	800	4.3	8.43	100	10	70	20
6	800	4.0	6.99	70	10	50	10
7	700	4.5	8.02	70	10	50	10
10	650	4.0	8.46	70	10	50	10
8	600	4.5	15.00	70	10	50	10
9	500	4.3	14.99	70	10	50	10
Dry combus	tion						
1	800	4.5	14.96	108	18	80	20
2	800	7.5	14.96	108	18	80	20

Table 1 Conditions for the cremation experiments.

Table 2 ¹⁴C and ¹³C concentrations for bone and CO₂.

Sample name	Sample fraction	Lab ID	Corrected pMC	Uncorrected pMC	δ ¹³ C (‰VPDB) ^a	δ ¹³ C (‰VPDB) ^b
bovine bone	collagen	KIA 30242	106.21 ± 0.29	108.78 ± 0.29	-13.45 ± 0.1	
	apatite	KIA 38371	104.98 ± 0.29	109.54 ± 0.28	-4.07 ± 0.14	-4.35 ± 0.1
old CO ₂	CO ₂	KIA 39049	0.23 ± 0.03	0.23 ± 0.03	-31.95 ± 0.2	-32.60 ± 0.1

^aMeasured with the AMS system.

^bMeasured with conventional mass spectrometry.

lated following Person et al. (1995) using the separation of the peaks corresponding to reflections [211], [112], [300], and [202] (see Figure 2).



Figure 2 A) Part of the FTIR spectra of sample 4 illustrating the calculation of the FTIR splitting factor. A baseline is drawn between 750 and 495 cm⁻¹. The heights of the peaks at ~602 and 565 cm⁻¹ are summed and divided by the height of the valley between them (Weiner and Bar-Yosef 1990). B) Definition of the crystallinity index by XRD using the refraction peaks [202], [300], [112], and [211] (Person et al. 1995).

Sample CO₂ for isotopic measurements (accelerator mass spectrometry [AMS] and gas isotope ratio mass spectrometry [MS]) was liberated from ~1 g of sample material with 4 mL of 60% phosphoric acid at 90 °C in a sealed ampoule reaction system (Nadeau et al. 2001). Before graphitization for subsequent AMS measurements, possible sulfur dioxide was removed by reacting the sample gas

with CuO and silver wool in a quartz tube at 900 °C for 4 hr. The sample CO_2 was graphitized and measured at the Leibniz-Labor AMS system using standard procedures (Nadeau et al. 1997, 1998). Isotope mass spectrometry was done directly on extracted CO_2 on a Finnigan Delta ETM instrument. Stable carbon isotope ratios are given in the δ^{13} C notation versus VPDB standard; errors are 1 standard deviation (Tables 2 and 3).

The carbon in the burnt bone apatite may be from original bone apatite carbonate, bone collagen, and/or burning fuel carbon (old CO_2 in the experiment):

$$C_{\text{HAP-burnt}} = C_{\text{HAP-org}} + C_{\text{col}} + C_{\text{oldCO2}}$$
(1)

where $C_{HAP-burnt}$ is carbon in burnt bone apatite, $C_{HAP-org}$ is carbon from original bone apatite, C_{col} is collagen carbon, and C_{oldCO2} is carbon from old CO₂, admixed during combustion.

Original apatite carbon and collagen carbon have essentially the same ¹⁴C signature, so the contribution of carbon from old $CO_2 M$ can be estimated directly from the measured ¹⁴C concentration of burnt bone apatite by:

$$M = \frac{{}^{14}C_{\text{HAP-org}} - {}^{14}C_{\text{HAP-burnt}}}{{}^{14}C_{\text{HAP-org}} - {}^{14}C_{\text{old}C02}} *100 \ (\%)$$
(2)

Since collagen carbon and the old CO_2 have different $\delta^{13}C$ signatures compared to bone apatite carbon, we use uncorrected ¹⁴C concentrations in percent modern carbon (pMC_{uncorr}) to estimate an assumed carbon exchange.

RESULTS

Chemical and Mineralogical Changes

Chemical and mineralogical changes of the bone occurring during the burning (<600 °C) and cremation (>600 °C) experiments were documented by FTIR and XRD analysis (Figures 3, 4). In general, the resulting material changes are similar to those found in real archaeological cremated bone (VIRI 4), e.g. loss of organics and carbonate with a main hydroxylapatite phase (HAp) plus a fraction of carbonated hydroxylapatite (cHAp) in conjunction with changes in crystallinity reflected by a change in SF from ~3 to >5 and in CI from <0.4 to >0.8 (from fresh to cremated).

However, a significant difference with respect to carbonate substitution is observed between materials cremated in the wet or dry conditions. Material cremated (>600 °C) under wet conditions shows a large depletion in carbonate content in general with only a fraction of carbonate substituted for PO_4^{3-} (B-type, C content ~0.1–0.4 wt%; Table 3), which is similar to archaeological cremated bones. On the other hand, material cremated under dry conditions showed a higher carbon content (~0.7 wt%, experiments 1 and 2) with carbonate substitution for PO_4^{3-} as well as for OH⁻ (B-type and A-type, respectively). Previous work also suggested a higher A-type carbonate substitution when water is excluded from the mineralization environment (LeGeros et al. 1969). Interestingly, SF for dry-combusted material (SF ~3.5) would indicate a smaller crystallite size if the relationship to crystallite size of Trueman et al. (2004) is applied, whereas CI (CI~1.3) clearly indicates an increase, which is also confirmed by transmission electron microscopy (TEM) of wet and dry cremated material (unpublished data).

Table 3 Sp	litting	factor (SI	F), crystallinity index (CI), ¹⁴ C	, and $\delta^{13}C$ value	s from cremat	ed bone apatit	e. Carbon exe	change with old	CO ₂ is estimated
using ¹⁴ C a	nd ¹³ C							, , ,	
Experiment nr	Temp. (°C)	Old CO ₂ (vol%)	SF CI (wt%)	ent Corrected pMC	Uncorrected pMC	δ ¹³ C (‰VPDB) ^a	δ ¹³ C (‰VPDB) ^b	C exchange with old CO ₂ ¹⁴ C indicated (%)	C exchange with old $CO_2^{13}C$ indicated (%)
Combustio	n withc	but CO_2 , v	wet						
4	800	0	$7.18 \pm 0.62 \ (n = 5) \ 1.1 \ 0.1$	103.98 ± 0.32	107.78 ± 0.33	-7.40 ± 0.18			
11	800	0	6.98 ± 0.32 $(n = 5)$ 1.1 0.1	102.73 ± 0.42	105.22 ± 0.42	-13.36 ± 0.29			
Combustio	n with	CO ₂ , wet							
ŝ	800	1.7	6.78 ± 0.22 $(n = 4)$ 1.1 0.1	69.63 ± 0.27	70.31 ± 0.27	-20.32 ± 0.19		35.9	56.9
5	800	10	$6.83 \pm 0.41 \ (n = 4) \ 1.1 \ 0.2$	54.27 ± 0.28	54.36 ± 0.29	-24.18 ± 0.15		50.5	70.4
9	800	14.3	$6.65 \pm 0.42 \ (n = 5) \ 1.2 \ 0.2$	50.29 ± 0.31	50.39 ± 0.31	-23.93 ± 0.2		54.1	69.5
9	800	14.3	0.1	39.74 ± 0.22	39.66 ± 0.22	-26.05 ± 0.11		63.9	76.9
7	700	14.3	$6.69 \pm 0.65 \ (n = 6) \ 1.1 \ 0.3$	62.81 ± 0.24	63.51 ± 0.24	-19.64 ± 0.18		42.1	54.5
7	700	14.3	0.3	67.29 ± 0.23	68.15 ± 0.23	-18.86 ± 0.17		37.9	51.8
10	650	14.3	$6.52 \pm 0.30 \ (n = 5) \ 1.0 \ 0.4$	48.89 ± 0.19	49.07 ± 0.19	-23.25 ± 0.08		55.3	67.1
8	600	14.3	$5.14 \pm 0.50 \ (n = 5) \ 0.3 \ 0.8$	82.84 ± 0.29	85.06 ± 0.29	-12.05 ± 0.25	-12.05 ± 0.1	22.4	27.9
8	600	14.3	0.7	84.28 ± 0.24	86.72 ± 0.25	-11.04 ± 0.18		20.9	24.4
6	500	14.3	$3.72 \pm 0.23 \ (n = 5) \ 0.1 \ 0.8$	94.20 ± 0.29	97.31 ± 0.30	-9.10 ± 0.34	-9.61 ± 0.1	11.2	17.6
6	500	14.3	0.4	94.03 ± 0.33	97.52 ± 0.34	-7.18 ± 0.20		11.0	10.9
Combustio	n with	CO ₂ , dry							
1	800	16.7	$3.50 \pm 0.48 \ (n = 6) \ 1.3 \ 0.7$	22.15 ± 0.15	21.92 ± 0.15	-30.05 ± 0.17	-30.51 ± 0.1	80.2	90.9
1	800	16.7	0.8	26.07 ± 0.14	25.84 ± 0.14	-29.24 ± 0.11		76.6	88.1
7	800	16.7	$3.63 \pm 0.42 \ (n = 4) \ 1.2 \ 0.7$	15.67 ± 0.14	15.47 ± 0.13	-31.09 ± 0.35	-31.79 ± 0.1	86.1	94.6
^a Measured wi ^b Measured wi	th the A th conve	MS system ational mat	s spectrometry.						

592 C M Hüls et al.



Figure 3 Selection of FTIR spectra of A) fresh bovine bone and an archaeological cremated bone from the VIRI intercomparison (VIRI sample 4, Naysmith et al. 2007), B) cremated bone material without old CO₂ (wet conditions), C) cremated and burnt bone with old CO₂ from 500 °C–800 °C (wet conditions), and D) cremated bone material with old CO₂ at 800 °C (dry conditions). Typical absorption peaks for organic and inorganic fractions are indicated.



Figure 4 Selection of XRD powder pattern of: A) archaeological cremated bone from the VIRI intercomparison (VIRI sample 4, Naysmith et al. 2007); B) cremated bone material without old CO_2 (wet conditions); C) cremated and burnt bone with old CO_2 from 500 °C–800 °C (wet conditions); and D) cremated bone material with old CO_2 at 800 °C (dry conditions). Typical XRD peaks for hydroxylapatite crystal planes are indicated in the upper panel.

Another chemical difference is observed between material cremated under wet conditions in comparison to archaeological cremated bone apatite. FTIR of true cremated bone material (i.e. the VIRI 4 sample in Figure 3) shows a prominent vibration intensity around 2012 cm⁻¹, which is absent in the cremated bone material of the experiments discussed here. This specific infrared (IR) frequency is related to the formation of cyanamide-apatite Ca₉(PO₄)₅(HPO₄)(HCN₂), formed during heating of the apatite under an ammonia atmosphere (Dowker and Elliott 1979; Habelitz et al. 2001). Interestingly, this species is observed in cremated bones when cremation can be assumed to have occurred along with all body tissues (skin, muscles, fat, etc.), i.e. body cremation. This specific IR frequency is not observed where the archaeological finds indicate cremation of defleshed bone, e.g. in bone waste from a meal, which became cremated after thrown in a camp fire. In addition, cyanamide-apatite is also indicated by FTIR from cremated bone material studied by Van Strydonck et al. (2010), who used modern bone material together with the attached body tissue for cremation experiments. Regarding these observations, we suggest that the formation of cyanamide-apatite is linked to the cremation of bones with attached body tissue and thus may serve as an indicator for real cremation. Aside from the remaining carbonate, the incorporation of a cyanamide group (HCN $_{2}^{-}$) delivers additional carbon to the resulting cremated apatite.

Isotopic Changes

Apatite carbon isotopes (¹⁴C, ¹³C) were analyzed on the experimentally burnt and cremated bone by AMS. For some samples, a parallel measurement of the apatite δ^{13} C was made with conventional MS (Table 3). Overall, the agreement between AMS and MS δ^{13} C measurements is high (n = 5, R = 1; $\delta^{13}C_{AMS} = 0.99 \,\delta^{13}C_{MS} + 0.17$).

The isotopic signals measured in the burnt and cremated bone apatite samples document large and significant changes in the apatite carbonate (Table 3). Low ¹⁴C and ¹³C concentrations are seen in samples cremated at 800 °C with old CO₂. The wet cremation with a CO₂ concentration of 14.3 vol% gave ¹⁴C values between ~40 to 50 pMC (Table 3, experiment 6). Using Equation 2 and uncorrected ¹⁴C values, an apparent carbon exchange with the old CO₂ of about 54–64% is indicated. These results agree with recent findings of Van Strydonck et. al. (2010), who ran cremation experiments under more natural conditions using modern bone with remaining tissue and old coal at similar temperatures. Lower old CO₂ concentrations during cremation result in a slightly lower carbon exchange of 50% and 36% (Table 3, experiment 5, CO₂ ~10 vol%; experiment 3, CO₂ ~1.7 vol%). At temperatures ≤600 °C, estimated carbon exchange is lower, ~22% at 600 °C and 11% at 500 °C, which is in accordance with the lower crystallographic changes.

The largest carbon exchange is indicated by ¹⁴C measurements on apatite cremated at 800 °C under dry conditions (Table 3, experiments 1 and 2). For experiment 1, ¹⁴C values of 21.92 and 25.84 pMC indicate an exchange of 80.2% and 76.6%, respectively. Experiment 2 ran longer, ~7.5 hr instead of 4.2 hr and the measured ¹⁴C value of 15.47 pMC indicates an even larger carbon exchange of ~86%. The larger carbon exchange can be explained by carbonate substituted for OH⁻ (A-type) and for PO₄³⁻ (B-type), while wet-cremated apatite shows carbonate only at the PO₄³⁻ position (B-type). This fact is also seen in higher carbon contents of dry compared to wet cremated apatite (carbon weight fractions ~0.7 to 0.8 versus ~0.1 to 0.2 wt% C). It is therefore quite evident that changes in cremation conditions, such as availability of water vapor, may strongly affect carbon exchange.

Measured ¹⁴C values from burnt bone apatite thus imply a significant carbon exchange with CO_2 from the burning atmosphere, which, in the case of a natural cremation, would come from the burning fuel (e.g. wood). Since the wood used for cremation may partly be old, cremated bones could also suffer from an old-wood effect. Depending on the circumstances of the cremation (temperature,

596 C M Hüls et al.

duration, composition of the burning atmosphere, and composition/age of the fuel [nearly fresh or old wood]), an age effect of 50–100 yr may be possible, which could get even larger when a significant amount of "old" fuel is used (e.g. peat, coal).

If we assume only apatite carbonate and old CO_2 as carbon sources and ignore isotopic fractionation, we can also use the large difference in $\delta^{13}C$ values between the original bone apatite and the old CO_2 (about -4‰ and -32‰, respectively) to estimate carbon exchange. As shown in Table 3, the carbon exchange calculated from $\delta^{13}C$ is significantly larger than the exchange based on ¹⁴C. This discrepancy indicates that other factors, such as isotopic fractionation and carbon derived from collagen or post-burning contamination, need to be considered.

Two wet-cremation experiments (4 and 11; Table 3), run without old CO_2 loading in order to address this issue, show depleted values for both ¹³C and ¹⁴C. Collagen and apatite carbonate have similar ¹⁴C and different ¹³C concentrations, so admixture of collagen carbon will cause only a ¹³C depletion, and cannot explain the observations. Fractionation will affect both isotopes and, if we assume the usual mass difference dependence, we can use the measured $\delta^{13}C$ values to calculate the ¹⁴C values to expect. For experiments 4 and 11, $\delta^{13}C$ values are depleted by 3.3‰ and 9.3‰, respectively, relative to the original bone apatite, which translates to a predicted decrease of 0.73 and 2.04 pMC in ¹⁴C, significantly less than the observed decrease of 1.76 and 4.32 pMC. Thus, fractionation alone cannot explain the observations.

Figure 5 gives the measured ¹⁴C (uncorrected for ¹³C fractionation) and δ^{13} C values of the burnt and cremated apatite samples. All samples show a deviation from a mixing line between original apatite carbonate and old CO₂ isotopic composition with consistently lower δ^{13} C values. This difference is



Figure 5 ¹⁴C (uncorrected for ¹³C) versus δ^{13} C. The black dotted line represents the mixing line between old CO₂ and bone apatite carbonate without fractionation. The gray field is defined by mixing lines with a ¹³C fractionation of -3‰ at 800 °C and -4‰ at 500 °C, respectively, for an isotope system of CO₂ to CaCO₃ in equilibrium (Bottinga 1969; Scheele and Hoefs 1992). The ¹⁴C/¹³C end-members for collagen, apatite, and old CO₂ are indicated.

apparently lower at temperatures <600 °C at which changes in crystal properties, carbonate loss, and estimated carbon exchange are also low, and high at temperatures >600 °C with a large exchange between CO₂ and apatite. This is as expected from thermal decomposition (Zazzo et al. 2009) and from studies of the calcite-CO₂ system in thermodynamic equilibrium at high temperatures (Bottinga 1969; Scheele and Hoefs 1992) that report a calcite ¹³C depletion relative to CO₂ of ~4‰ at 500 °C and 3‰ at 800 °C. Assuming this relation to hold for our apatite system, the apatite carbonate in equilibrium with the old CO₂ should have δ^{13} C values of -35% to -36% (800 to 500 °C), which gives a mixing field closer to, but still less negative, than the observed values. Use of the depleted carbonate values as end-members gives a relative reduction of the ¹³C-calculated exchange values in Table 3 of about 10%, bringing them closer to, but still not in agreement with, the ¹⁴C-based values. For the dry cremation with CO₂ at 800 °C (experiments 1 and 2), the change is particularly important as the reduction in the ¹³C-based exchange is especially large here and brings it relatively close to the ¹⁴C-based values.

Although contamination with atmospheric CO_2 during the cremation experiment can be excluded in the sealed combustion system, a post-burning contamination of an activated apatite formed by carbonate loss during combustion is possible after removal of the sample from the combustion system, especially during sample grinding for AMS CO_2 extraction. Isotopically light laboratory air depleted in ¹⁴C (such as from the use of dry ice in the lab rooms) may have contributed to the residual ¹⁴C depletion observed for samples 4 and 10 after fractionation correction. Unfortunately, we do not yet have isotope measurements (¹⁴C, ¹³C) of laboratory atmospheric CO_2 to quantify this possibility. Clearly, more tests are needed to rule out the possibility of post-burning/cremation contamination or to understand fully the dynamics of the cremation process.

CONCLUSIONS

Infrared spectroscopy (FTIR) and X-ray diffractometry (XRD) confirm the complete loss of organics, loss of carbonates, and significant crystallographic changes in cremated bone apatite. They also reveal a difference in carbonate substitution between apatites cremated at 800 °C wet: water available, CO_3^{2-} only substituted for PO_4^{3-} (B-type, being the main type of carbonate substitution in bone apatite); and dry: no water, B-type with additional substitution of CO_3^{2-} for OH⁻ (A-type). Dry cremation showed a much lower carbonate loss.

¹⁴C concentrations and δ^{13} C values of burnt and cremated apatite indicate significant carbon exchange with CO₂ in the combustion atmosphere—up to 64% (wet cremation) and 86% (dry cremation), both based on ¹⁴C values—which varies with CO₂ concentration and duration. Archaeological cremated bone apatite may thus contain a significant amount of carbon originating from the burning fuel, and their ¹⁴C dates may thus suffer from an old-wood effect.

 δ^{13} C values generally yield a much larger calculated carbon exchange than the ¹⁴C measurements if any isotope fractionation between CO₂ and apatite carbonate is left out of consideration. The discrepancies between the results obtained from paired isotope values show that the isotopic changes of apatite carbonate during burning and cremation are not governed by carbon exchange between apatite carbonate and the combustion gases alone. Temperature-dependent isotope fractionation between combustion CO₂ and apatite carbonate, admixture of organic (collagen) carbon, and postcremation contamination with atmospheric CO₂ may also have influenced the results.

ACKNOWLEDGMENTS

Thanks are due to the Leibniz team for sample preparation and AMS analysis. The discussions and collaboration with Mark Van Strydonck and Mathieu Boudin as well as the comments and suggestions by the reviewers are gratefully acknowledged.

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