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Applying isotope analyses of cremated human bones in archaeological research – a review

Abstract

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Numerous experiments have recently been conducted on burnt bones in order to develop methods of isotope analysis which would be useful in archaeological research. Since the results of these studies are not yet widely known, this review presents their potential applications in investigations of human remains from cremation burials. Radiocarbon dating of burnt osteological materials is discussed, including problems related to the "old wood effect". Also considered is the analysis of light stable isotopes, i.e. δ^{13} C, δ^{15} N and δ^{18} O, which is unsuitable for palaeodietary determinations, but useful as a source of information about certain parameters of funeral pyres. Tracing geographical origins and human mobility is possible by means of the analysis of strontium isotope ratio 87 Sr/ 86 Sr. Since an understanding of high-temperature-induced transformations of bone structure and chemical composition is important for these considerations, a detailed account of the processes is given as an introduction.

Key words: isotopes, cremated human bones, radiocarbon dating, funerary practices, migration

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1. Introduction

The application of isotope analyses in archaeological research is a landmark achievement of the cooperation between the humanities and the natural sciences. This remark is easily justified by pointing out the significance of δ¹⁴C radiocarbon determinations, which have remained the first-choice method of absolute dating for over 70 years. Furthermore, the analyses of δ^{13} C and δ^{15} N isotopes are currently widely used in studies of dietary preferences of past populations. Another success was the development of protocols for measuring the ratio of strontium isotopes 87Sr and 86Sr, which allows the detection of geographical origins and for the possible migrations of selected individuals to be traced. The ability to provide answers to such basic questions as to when, who, what, wherefrom and where-to, was the primary drive which led to isotope analyses becoming a valued investigative tool in archaeology, and one which has also aroused interest among the Polish scientific community (e.g. Belka *et al.* 2018; Szczepanek *et al.* 2018; Wadyl (ed.) 2019; Linderholm *et al.* 2020).

The headway of isotope analyses of unburnt human skeletal remains attracted the attention of scholars concerned with those periods, when cremation was the common funerary rite. The first and foremost question was the following: are these analyses equally valuable in the study of osteological material from cremation burials? It is known that the process of burning causes fragmentation, warping and the shrinkage of bones, all of which reduces the pool of data about the deceased. However, it was noted that the high degree of fragmentation of burnt remains may not necessarily result from cremation itself, but instead be the effect of mortuary rituals, post-depositional processes and the exploration of urn vessels (Harvig *et al.* 2012; Harvig 2015, 44, fig. 3.1). In general, it was observed that cremated

remains may provide more information than poorly-preserved skeletal material from inhumation graves (Williams 2015, 259–261; cf. Thompson 2015a, 1–3). Recent decades have also brought progress in research on possible burning-related alterations of the chemical composition of bones (Snoeck 2014, 11–26). As a result, burnt osteological material became a valuable source of physico-chemical knowledge to be used in archaeology and forensic science (Harbeck *et al.* 2011; Schmidt and Symes (eds.) 2015; Ubelaker 2009; 2015; Mamede 2018). The current state-of-the-art allows the above-mentioned question to be answered and this constitutes the main objective of this paper.

2. Heat-induced transformations of the chemical composition of bone tissue

To assess the possible applications of isotope analyses to burnt osteological material, it is important to understand the transformations of bone structure and chemical composition that result from exposure to high temperatures.

Bone is made up of three basic components: 1) water, 2) the organic phase, mainly collagen (ca 90%) as well as proteoglycans, non-collagen proteins, phosphoproteins and lipids, and 3) the inorganic matrix called bone apatite or bioapatite (LeGeros 1991; Lee and Einhorn 2001). Due to the destructive impact of the temperature of the funerary pyre, only the inorganic phase, which constitutes ca 60% of bone mass, is useful in most studies of burnt remains.

Bioapatite is a crystal form of calcium hydroxyphosphate, which strongly reassembles the naturally occurring mineral hydroxyapatite (Ca₅(PO₄)₃(Cl/F/ OH)₂). However, in contrast to the latter, it is more susceptible to non-stoichiometric transformations of chemical compounds, which enables the incorporation of ions present in the blood into the structure of bioapatite (Figure 1). Essential in this respect are the carbonate ions (CO₃²⁻), which substitute nearly all of the hydroxyl groups (OH⁻) and part of the phosphates (PO₄³⁻) – called A-type carbonates (exchangeable with OH⁻) and B-type carbonates (exchangeable with PO₄³-) respectively (LeGeros 1969; 1991; Astala et al. 2005; Wopenka and Pasteris 2005; Snoeck 2014, 12; Madupalli et al. 2017). This process of substitution is highly relevant to the structure of the inorganic matrix of bone. The ions (A- and B-type carbonates) are not of the same shape and size, leading to deformations of bioapatite crystallinity during the exchange. For this reason, bioapatite is built up of smaller crystals than hydroxyapatite (it is less crystallized), which in turn leads to its enhanced susceptibility to various chemical processes, including diagenetic ones (Wopenka and Pasteris 2005; Snoeck 2014, 12; Snoeck et al. 2014a; Mamede et al. 2018).

High temperatures strongly alter both bone structure and their chemical composition. At first, the chain of transformation involves dehydration and the destruction of the organic phase, including the disintegration of collagen fibres. At the same time, the inorganic matrix undergoes a progressive loss of

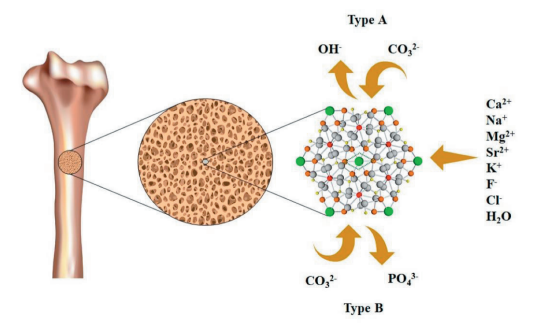


Fig. 1. The transformation of chemical compounds in bone apatite (after Mamede et al. 2017, modified by B. Wolska)

carbonates (CO₃²⁻), a rise of hydroxyl groups (OH⁻), and even the formation of new mineral forms, i.e. NaCaPO4, NaCl and KCl (Van Strydonck et al. 2005; Zazzo et al. 2009; Snoeck 2014, 39-94; Snoeck et al. 2014a; Mamede et al. 2018, Fig. 2). However, above all, these processes are accompanied by a gradual increase of bone apatite crystallinity, resulting in a high resilience to chemical weathering (Stiner et al. 1995; Lebon et al. 2010; Quarta et al. 2013; Thompson 2013; 2015; Snoeck 2014, 16, 89, 123; Snoeck et al. 2014a; Mamede et al. 2018). This property makes it similar to two-hydroxyapatite - the main component of the enamel of teeth which are considered the most robust elements of the human skeleton (LeGeros 1991; Hoppe et al. 2003; Chai et al. 2009; Muller et al. 2019). Consequently, burnt bone is highly resistant to postdepositional chemical transformations, which is significant from the perspective of physico-chemical and osteological analyses. It should be noted that this only pertains to fully calcined bones, that have been exposed to temperatures higher than 600°C (Snoeck et al. 2014a).

Isotopes (gr. "equal place") are variants of chemical elements that have the same atomic number but a different mass number (Jones and Atkins 2016, 11). Based on their stability, they are divided into stable, e.g. ¹³C, ¹⁵N, ¹⁸O, ⁸⁶Sr, ⁸⁷Sr and radioactive, such as ¹⁴C. In the life cycle of humans and animals, isotopes accumulate in the organic phase and inorganic matrix of bones. For instance, ¹⁴C infiltrates organisms during the digestion of ¹⁴CO₂, which is an element of the food consumed (Jones and Atkins 2016, 1034). Similarly acquired are strontium isotopes ⁸⁷Sr and ⁸⁶Sr, present in ingested water and plant/animal tissues, in amounts dependant on the local geological background (Beard and Johnson 2000; Bentley 2006; Slovak and Paytan 2012; Snoeck *et al.* 2015).

The above-mentioned heat-induced alterations of bone tissue bear heavily on studies of its isotopic composition. Poor preservation of extractable collagen (with a nitrogen value lower than 0,8%; see Brock et al. 2010) precludes the possibility of analysing this particular fraction, as is usually done in the case of unburnt bones (e.g. skeletal burials). Bone apatite is subjected to significant deformations as a result of the gradual loss of carbonates and increased number of hydroxyl groups, what corresponds with the loss and increase (respectively) of certain chemical elements (isotopes). These undergo further transformations, such as exchange and/or incorporation/fractioning during the process of cremation or subsequent residence of the remains in the soil (see below). All in all, it seems justi-

fied to say that burning causes irreversible changes in the original isotopic composition of bone. Nevertheless, even in this form, it is still a source of data that is extremely valuable in archaeological research.

3. Isotope analyses in studies on burnt human bones

3.1. Radiocarbon dating

Radiocarbon dating of burnt osteological remains was first conducted at the end of the 20th century. During the procedure, comparable ¹⁴C dates were obtained for burnt bone and charcoal derived from the same context, opening up the possibility to determine the age of archaeological materials in which no organic matter was present (Lanting and Brindley 1998). This revolutionary discovery spurred further research (Lanting and Brindley 2000; Lanting et al. 2001; Van Strydonck et al. 2005; Naysmith et al. 2007), yet this did not fully consider the potential differences between the age of bones and the fuel used for their cremation. As it transpired, this has major relevance for radiocarbon dating: the process leading to the "old wood effect" takes place when a bone is burnt in the presence of much older wood - the latter "conveying" a portion of its age, resulting in the artificial oldening of osteological material (Olsen et al. 2008; 2013; Hüls et al. 2010; Zazzo et al. 2012; Snoeck et al. 2014b). This phenomenon is best illustrated by the results of an experiment in which a contemporaneous pig trotter was burnt using ca 480-year-old wood, and its remains obtained a date of 326 ± 22 BP (Snoeck et al. 2014b). A similar but opposite effect was observed in a test where "old" archaeological bone and modernday wood were subjected to burning (Zazzo et al. 2012). These studies have shown that 14C dates do not necessarily indicate the actual age of burnt remains, but the age of the fuel used.

The above-mentioned findings reflect certain chemical processes that take place in bone during its exposure to high temperatures, e.g. of a fireplace or funeral pyre. Specifically, of crucial importance is the origin of the carbon isotopes present in the carbonate fraction of burnt bone apatite, since surely some of them are not part of the original composition (hence the "old wood effect"). Four mutually non-exclusive possibilities were proposed: 1) contamination of the carbonate fraction of bone apatite by atmospheric carbon, 2) exchange of carbon between the carbonate fraction of bone apatite and the gas released by the cremation fuel, 3) incorporation of organic carbon

released during the combustion of organic matter, i.e. skin, muscle, fat, 4) incorporation of organic carbon released during the combustion of collagen as well as other proteins and lipids, and 5) heat-induced fractioning of the carbonate fraction of bone apatite (Figure 2) (Zazzo *et al.* 2009; Snoeck *et al.* 2014b).

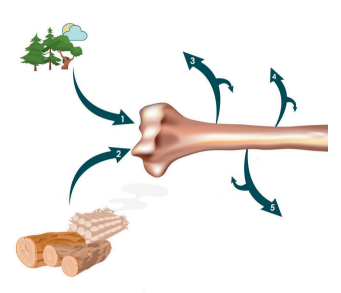


Fig. 2. Sources of CO_2 that probably affect the isotopic composition of the carbonate fraction of bone apatite during heating/burning: 1) the atmosphere, 2) the burnt fuel, 3) the burnt organic matter, i.e. skin, muscle, fat, marrow 4) the burnt collagen, 5) the burnt carbonate fraction of bone apatite. The arrows indicate the possible directions of carbonate and/or CO_2 flow, and the conditions in which the fractioning of carbon occurs (after Snoeck 2014, 97, fig. 5.1., modified by K. Kij)

In reality, only four possibilities are relevant (2-5), the first one being less feasible due to the low CO₂ content (0,04%) in the atmosphere. The third and fourth were confirmed only for the exchange of carbon between bone apatite and collagen. Soft tissues (skin, muscles, fat) undergo relatively rapid combustion (see Fairgrieve 2008, 44-45, tab. 3.3, 3.4.) and for this reason they do not have a substantial influence on the final chemical composition of the carbonate fraction of burnt bone apatite (Snoeck et al. 2014b). At the same time, since collagen is close to apatite and subjected to gradual degradation up to 600°C, it has more potential for mutual chemical interaction (Hüls et al. 2010; Van Strydonck et al. 2010; Zazzo et al. 2012; Snoeck et al. 2014b; 2016a). It is not responsible, however, for the incorrect radiocarbon dating of burnt bones, because both these fractions (collagen and apatite) share a similar (although not identical) 14C age (Snoeck 2014, 121; Snoeck et al. 2014b). Experiments have also confirmed the fractioning of carbon isotopes, occurring between bone apatite and CO2 released by burning organic matter and combustion gases (Zazzo *et al.* 2009; Hüls *et al.* 2010; Van Strydonck *et al.* 2010; Harbeck *et al.* 2011; Snoeck *et al.* 2014b; 2016a). Nevertheless, the isotopic composition of carbon in burnt bone is mostly influenced by the process of the exchange with the burning environment, i.e. an exchange with carbon emitted by the fuel (Table 1). It is precisely this phenomenon which is mainly responsible for the "old wood effect" (Olsen *et al.* 2008; 2013; Hüls *et al.* 2010; Zazzo *et al.* 2012; Snoeck *et al.* 2014b; 2016a).

Table 1. Research results indicating an increased exchange of carbon between the burnt fuel and the carbonate fraction of bone apatite, what is the main reason of the "old-wood" effect.

Authors	Research material	Carbon exchange with old fuel
Hüls <i>et al</i> . 2010	Modern bovine bone	36% – 86%
Zazzo et al. 2012	Archaeological human and animal bones	48% – 91%
Snoeck et al. 2014	Modern animal bones	39% – 95%

Summarizing, the carbon present in the carbonate fraction of burnt bone apatite has three possible sources: endogenic apatite carbon, endogenic carbon present in collagen, and fuel-derived carbon. In light of this knowledge, is it practical to date burnt remains using radiocarbon? Despite certain limitations, in most cases the method delivers relevant information. However, the following circumstances should be considered:

- 1) The remains of the deceased might have been cremated some-time after their death, using young wood as fuel. The result of the ¹⁴C analysis will indicate the approximate date of the burning, not the actual moment of the individual's death.
- 2) The remains of the deceased might have been cremated using also old wood as fuel. The result of the ¹⁴C analysis will indicate the approximated age of the wood, not the actual moment of the individual's death.

Both these conditions may be difficult to assess for cremated materials from archaeological contexts, with the consequence that the radiocarbon dating of singular samples will deliver unambiguous data. The interpretative field can be narrowed by obtaining a series of dates for several graves, or in the case of bi-ritual cemeteries, by additionally comparing values acquired for skeletal and cremation burials. Such radiocarbon research has recently been caried out on numerous (1428) samples from cremation and inhumation burials discovered at archaeological sites in Belgium. The results have shown cultural changes in funerary practices from the Mesolithic period to the Middle Ages (Capuzzo *et al.* 2020).

3.2. Light stable isotopes ¹³C, ¹⁵N and ¹⁸O

Isotopes of light elements present both in the organic phase and inorganic matrix of bone, such as carbon, nitrogen and oxygen, have been used in bioarchaeological studies to reconstruct the dietary preferences of past communities. Their successful applications to unburnt skeletal remains led to an interest in using a similar analysis for cremated osteological material. Unfortunately, however, this was found to be unsuitable for such determinations (DeNiro *et al.* 1985). The advantages of investigating stable isotopes in burnt bone only became clear in the last decade, although they turned out to be unrelated to palaeodietary research (Snoeck *et al.* 2014b; 2016a).

Isotope signatures of δ^{13} C, δ^{15} N and δ^{18} O preserve their original values only up to 200°C (DeNiro et al. 1985; Harbeck et al. 2011; Schurr et al. 2015). Changes occurring above this threshold were identified experimentally, with varying degrees of influence on the final isotopic composition. The most recent results indicate a rapid loss of light isotopes in collagen and apatite, except for structural carbon δ^{13} C, the level of which decreases to 700-800°C but starts to increase with a further rise of temperature (Harbeck et al. 2011; Snoeck et al. 2016a). This general reduction is directly related to the heat-induced alteration of bone composition and structure, above all the combustion of the organic phase and loss of carbonates (CO₃²⁻) from the inorganic matrix, accompanied by the processes of exchange, fractioning and incorporation of carbon from different sources (see 3.1., Fig. 2). The current state of research shows that palaeodietary analyses in burnt bone are not possible (DeNiro et al. 1985; Lanting et al. 2001; Van Strydonck et al. 2005; Olsen et al. 2008; 2013; Zazzo et al. 2009; 2013; Harbeck et al. 2011; Snoeck et al. 2014b; 2016a; cf. Schurr et al. 2015).

It has recently been pointed out that stable light isotopes may be used in studies oriented at reconstructing the cremation ritual, serving as proxies of the conditions in the funeral pyre during the burning of the deceased (Snoeck *et al.* 2016a).

This method was developed in the course of experiments concentrated on the isotopic variance of Δ^{13} C and Δ^{18} O ($\Delta = \delta$ of the sample – δ unburnt apatite), instead of their absolute values (Snoeck et al. 2016a). To confirm that the conditions of cremation have a meaningful bearing on this variance, a test was arranged that simulated possible different circumstances of the ritual - burning of a sample on firewood (with fuel), also with the addition of C₃ and C₄ plants (or their products, such as maize). Burning in laboratory conditions (without fuel) served as reference (Snoeck et al. 2016a). It was observed that bones cremated on firewood have less carbon δ^{13} C and oxygen δ^{18} O, and that the values of oxygen δ^{18} O are lower when the exchange of carbon δ^{13} C between the bone and the fuel intensifies. Moreover, bones containing cyanamide – an organic compound identified by FTIR and expressed in relation to the amount of phosphorus in the sample (CN/P) – usually have higher values of carbon δ^{13} C with a slight difference in oxygen δ^{18} O presence. The addition of C₄ plants during the cremation also has an impact on the isotopic composition of samples, resulting in considerably enhanced levels of δ^{13} C and δ^{18} O (Snoeck et al. 2016a). All these observations are related to complex processes of exchange, fractioning and incorporation of the described elements, which to a varied extent (still unassessed) shape the final isotopic composition of burnt bone apatite. Tracing this composition in individual samples allows recognizing remains cremated under different conditions. Three possibilities were outlined:

- 1) identification of remains burnt in the presence of firewood and those cremated without its use (probably a rare occurrence in archaeological materials).
- 2) identification of remains burnt using C₃ firewood and C₄ plant fuel.
- 3) identification of remains burnt on larger or smaller funeral pyres, in different places (pyres located in landscape positions with higher or lower oxygen availability) and the positioning of the body during the cremation (on/in the pyre).

Considering these remarks, a comparison of the results of δ^{13} C and δ^{18} O isotope analysis of remains from a single cemetery allows the pinpointing of those individuals who were cremated in different circumstances than others (i.e. their isotopic values do not correlate with the cemetery "baseline"). Further investigations may address the possible reasons for such exceptional treatment, which can be a consequence of customs specific for a given community, or a sign suggesting that the deceased in question was cremated in a location different than the place of burial. In the lat-

ter case, an analysis of strontium isotope ⁸⁷Sr and ⁸⁶Sr ratio is a supplementary investigative method.

3.3. Strontium isotope ratio 87Sr/86Sr

Geographical origins and the mobility of people have been intensely studied topics in archaeology. Informative in this matter is an analysis of strontium isotopes ratios *FSr/*6Sr – an element assimilated by the human organism during the ingestion of food and water, which becomes incorporated in the structure of bones in a manner similar to calcium. Among the isotopes discussed in this paper, strontium is the only one resistant to high temperatures, a fact known from the early 1990's (Grupe and Hummel 1991). In the last decade, the method was applied to remains from cremation graves, confirming its usefulness as a tool in research on ancient migrations (Snoeck 2014, 125–137; Harvig *et al.*, 2014; Snoeck *et al.*, 2015; 2016b; 2018; Graham *et al.* 2019).

As an element heavier than 50u, strontium is not subjected to the same chemical processes as the above-mentioned light isotopes during burning. Experiments have indicated that the 87Sr/86Sr value in samples does not change, even after heating in temperatures exceeding 1000°C (Grupe and Hummel 1991; Beard and Johnson, 2000; Harbeck et al. 2011; Snoeck et al. 2015). This is related to a relatively high melting point of strontium carbonate (ca 1500°C), i.e. considerably above the temperatures potentially reached in funeral pyres (McKinley 1994; Ubelaker 2015; Snoeck 2014, 126; Snoeck et al. 2015). Moreover, there is no noticeable process of fractionation between 87Sr and 86Sr during bone combustion (as was it observed in the case of carbon isotopes), because the difference between their atomic masses is low (Harbeck et al. 2011; Snoeck 2014, 126; Snoeck et al. 2015).

Of crucial importance in strontium isotope analysis is the type of material selected for the study. In the case of inhumation burials, the enamel of teeth is usually sampled due to its specific properties: 1) the lack of metabolic activity, which ensures a fixed isotopic composition during the whole human life, 2) a high resilience to post-depositional alteration (see 3.1.), and 3) temporal differences in the formation of individual tooth crowns, allowing the assessment of the individuals' place of residence in particular periods of their lives, between the early childhood and adolescence, depending on the type of tooth studied. Due to the rare occurrence of tooth enamel in cremated materials, a substitute needed to be found. This role was taken by the otic capsule located in the petrous part of

the temporal bone which is composed of very dense bone tissue that is resilient to chemical processes (Jeffery and Spoor 2004; Jørkov *et al.* 2009; Harvig *et al.* 2014). Its morphology remains unchanged even under high temperatures and for this reason it is often discovered among burnt human remains. Experimental research confirmed the credibility of this material in studies on the geographical origins of cremated individuals (Harvig *et al.* 2014). However, it is important to note that analyses of the petrous portion of the temporal bone reveal information about the place of residence in very early childhood, as this bone forms at a prenatal age and does not change after the second year of life (Jeffery and Spoor 2004; Jørkov *et al.* 2009).

To ascertain the whereabouts of individuals in later stages of their lives, additional samples have to be taken from other parts of the human skeleton (e.g. the long bones or cranium), which had been rebuilt during the whole life due to progressing ossification. This process involves the embedding in bone of new bioapatite crystals, including strontium isotope values characteristic for the local geological background. However, ossification is a non-uniform process and proceeds differently in particular types and structures of bones. It is also determined by the metabolic level of the individual. For these reasons, the analysis of the strontium isotope composition in such samples is less straightforward as it cannot be related to a specific age of humans, (contrary to the case of the petrous part of the temporal bone or the enamel of teeth, whose period of formation in the human organism is wellresearched), but only to a time interval of 10-20 years before their death (Jørkov et al. 2009; Szostek et al. 2015; Snoeck et al. 2015). Moreover, the possible intensive mobility of individuals or several year-long residence in new places have to be considered, which significantly influences the strontium isotope composition in the skeletal tissue, further complicating proper interpretation (Slovak and Paytan 2012, 748-749). Consequently, it is advised to conduct the analysis using samples taken from different elements of the cremated remains.

Another matter to be considered when studying strontium isotopes in the context of human mobility is the potential susceptibility of burnt bones to absorption/incorporation of exogenic strontium during their long residence in soil. This issue, important from the viewpoint of archaeology, was investigated in a series of experiments which have shown successive enrichment in Sr of a calcined cow tibia left for 12 months in a strontium-rich (87Sr) solution (Snoeck *et al.* 2015). The subsequent application of acetic acid and an ul-

trasonic bath helped to remove the contamination, including the exogenic strontium isotopes; in effect, a sample with an original ⁸⁷Sr and ⁸⁶Sr composition was obtained (Snoeck et al. 2015). During the same research, an identical procedure (artificial enrichment in Sr and sample pre-treatment) was administered to the enamel of a horse tooth, which at the end of the experiment still exhibited a higher value of exogenic strontium compared to the burnt cow bone. Most likely, this results from the incorporation of the isotope into the crystallite structure of teeth, instead of its absorption onto the surface, as in the case of burnt cow bone. The experiment shows the high resilience of the latter to the diagenesis caused by the high crystallinity of the cremated bone (Snoeck et al. 2015). This points towards an equal (or even slightly higher) credibility of cremated osteological material in strontium isotope analysis, compared to teeth enamel.

In sum, the physico-chemical processes that occur during cremation do not affect the original composition of strontium isotopes in bone, and the possible alterations resulting from protracted burial in soil may be mitigated by diligent laboratory treatment. In this light, burnt osteological remains are reliable material in studies on geographical origins and mobility in communities that cremated their dead.

4. Conclusions

Cremation burials are thought to have been practiced in Polish lands as early as the Mesolithic (Gil-Drozd 2011). This funerary custom proliferated in later times and burnt human bones are frequently discovered during archaeological fieldwork as a result. The paper reviews the possibilities of conducting isotope analyses on such material and shows that the research perspectives that they entail are centred around several issues.

The first is the determination of the absolute chronology of sepulchral sites by means of the radiocarbon dating of burnt bones. It must be remembered, however, that the obtained δ¹⁴C dates will not reflect the actual radiocarbon age of the remains, but of the wood used for the cremation. Consequently, the analysed samples may be oldened if they were burned in the presence of old wood (the "old wood effect"). Such circumstances may have been common in the past, as shown by anthracological studies of charcoal from cremation graves discovered in present-day Poland (Moskal-del Hoyo 2012).

The second is the interpretation of the funerary rite, albeit focused on the cremation procedure itself,

instead of the manner in which the burnt remains were deposited in graves. By comparing the values of $\delta^{13}C$ and $\delta^{18}O$ it is possible to assess whether the corpse was burned in the presence of fuel (usually wood; probably a universal occurrence in archaeological materials) or without it. Also feasible are estimates regarding the size of the funeral pyre and its location in the landscape with higher or lower oxygen availability, as well as the positioning of the body in/on the pyre. Tracing the values of these isotopes in a series of samples from a single cemetery allows us to recognize those individuals who were cremated in different conditions. This opens further avenues of inference.

Another issue is the identification of the geographical origins of people interred in cremation graves by an analysis of the strontium ⁸⁷Sr/⁸⁶Sr ratio, which in certain circumstances is simultaneously linked with attempts to describe the migrations they made during their lifetime. These matters are raised in discussions regarding long-range trade, mobility of labourers/specialists or even of whole communities, and for this reason information about the birthplace of individuals seems to have significance in many areas of research on the past.

On the other hand, isotope analysis of cremated bone is not a suitable method in palaeodietary studies. This is caused by the lack of collagen and the high temperature-induced changes in bone apatite which result in an irreversible alteration of $\delta^{13}C$ and $\delta^{15}N$ values. Other procedures should be sought to investigate the issue of alimentation.

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