Freshly excavated fossil bones are best for amplification of ancient DNA.

Mélanie Pruvost, Reinhard Schwarz, Virginia Bessa Correia, Sophie Champlot, Séverine Braguier, Nicolas Morel, Yolanda Fernandez-Jalvo, Thierry Grange, Eva-Maria Geigl

To cite this version:

HAL Id: hal-00124469
https://hal.archives-ouvertes.fr/hal-00124469
Submitted on 30 Jan 2007

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Classification: Anthropology/Evolution

Freshly excavated fossil bones are best for amplification of ancient DNA

Mélanie Pruvost¹, Reinhard Schwarz¹, Virginia Bessa Correia¹, Sophie Champlot¹, Séverine Braguier³, Nicolas Morel⁴, Yolanda Fernandez-Jalvo², Thierry Grange¹, and Eva-Maria Geigl¹*

¹Institut Jacques Monod, Tour 43, 2, Place Jussieu, 75005 Paris, France. ²Museo Nacional de Ciencias Naturales, José Gutierrez Abascal 2, 28006 Madrid, Spain. ³Musée de Carnac, 10, place de la Chapelle, 56340 Carnac, France. ⁴Musée Vert, 204, avenue Jean Jaurès, 72000 Le Mans, France. ⁵present address: IFBB Gerichtsmedizin und forenische Neuropsychiatrie der Universität Salzburg, Ignaz-Harrer-Str. 79, A-5020 Salzburg, Austria

* Corresponding author: Institut Jacques Monod du CNRS, Université Paris 6 et 7, Tour 43, 2, Place Jussieu, 75251 Paris cedex 05; Tel: +33-1 44 27 57 07; Fax: +33-1-44 27 57 16;
E-mail address: geigl@ijm.jussieu.fr

Number of text pages: 20, Number of figures: 2, Number of tables: 1, Number of supporting tables: 1

Number of words in the abstract: 242, Total number of characters in the paper (including equation, 1 table, 2 figures, references): 42,263

Keywords: ancient DNA/ DNA preservation/ fossil bones/bone diagenesis/fossil preparation/conservation
Despite their enormous potential for phylogeographic studies of past populations, the impact of ancient DNA analyses, most of which are performed with fossil samples from natural history museum collections, has been limited to some extent by the inefficient recovery of ancient genetic material. Here we show that the standard storage conditions and/or treatments of fossil bones in these collections can be detrimental to DNA “survival”. Using a quantitative palaeogenetic analysis of 247 herbivore “fossil” bones up to 50,000 years old and originating from 60 different archaeological and palaeontological contexts we demonstrate that freshly excavated and non-treated, unwashed bones contain six times more DNA and yielded twice as many authentic DNA sequences than bones treated with standard procedures. This effect was even more pronounced with bones from one Neolithic site where only freshly excavated bones yielded results. Finally, we compared the DNA content in the fossil bones of one animal, a c. 3,200 year-old aurochs, that were excavated in two separate seasons 57 years apart. Whereas the washed and museum-stored fossil bones did not permit any DNA amplification, all the recently excavated bones yielded authentic aurochs sequences. We established that during the 57 years, during which the aurochs bones were stored in a collection, at least as much amplifiable DNA was lost as during the previous 3,200 years of burial. This result calls for a revision of the post-excavation treatment of fossil bones to better preserve the genetic heritage of past life forms.
Introduction

Our knowledge of past life forms stems mainly from fossils, the only witnesses of extinct species, the phylogenetic analyses of which were boosted by the discovery that DNA is sometimes preserved in fossils (1). In fact, water-soluble DNA has been shown to persist in fossil bones for up to as many as 130,000 years in temperate regions (2). The analysis of this ancient DNA has the potential to provide us with answers to archaeological, palaeontological and anthropological questions, when the classical approaches of these latter disciplines cannot. During bone fossilization, however, DNA is at least partially degraded and chemically modified. Little is known about the modifications of ancient DNA that lead to its preservation. Thus, ancient DNA analysis constitutes an enormous methodological and conceptual challenge for palaeogeneticists. Moreover, despite some spectacular achievements, the failure rate of palaeogenetic investigations is high, since DNA preservation is rare, i.e., numerous fossil samples are analyzed, but few sequences are obtained. For example, the success rate of DNA amplification declines with increasing average temperature in the area from which the fossils originate. Whereas 78% (3) and 62% (52 to 71%) (4) of permafrost samples were reported to be successfully amplified, samples from regions with moderate temperature amplified with a 23 to 67% success rate (5) and from arid, hot climates with a mere 2 to 4% success rate (5). Temperature has indeed been identified as a key factor in DNA preservation (6), but cannot be the only one. All factors influencing chemical reactions (e.g., pH, Eh, irradiation, chemical composition of bone and soil, hydrology) may play a role in a complex fashion that is not as yet comprehended.

It has been shown that bones are locally destroyed by bacteria and fungi (“microscopical focal destruction”) (7) and that diagenetic alteration is localized leaving discrete “fossilizing regions” where fossilization can occur (8). Long-term DNA preservation might be favored within various types of microenvironments with different biological and physicochemical
properties, so-called “molecular niches” (9), which have formed in the bones during fossilization. The particular conditions ruling in these niches must slow down DNA degradation processes (e.g., adsorption of DNA to apatite crystallites or clusters of intergrown bone crystals that are not affected by diagenetic changes, such as those described by Trueman et al. (10) and by Salamon et al. (11); complexation of DNA; low local chemical reactivity, particular local pH and ionic conditions etc. (9)). These microenvironmental conditions and the physico-chemical conditions prevailing in the macroenvironment in which the fossilization process takes place must be interdependent, suggesting that any drastic modification of the conditions outside the bones can affect the preservation of the DNA within them. Abrupt changes in the macroenvironmental conditions, such as those which occur during the excavation of fossils and their transfer to museums and natural history collections, might constitute such changes in the physicochemical conditions in the microenvironment and might thus have dramatic consequences on DNA preservation. Therefore, we tested systematically the interdependence of DNA preservation and post-excavation treatment by analyzing bones that had experienced different post-excavation preservation conditions. Here we show how detrimental to DNA survival standard post-excavation treatments can be.
Results

To analyze the influence of standard post-excavation treatments on the preservation of DNA within archaeological bones, called here “fossil bones”, we analyzed in parallel fossil bones from museum collections and freshly excavated bones that were kept after excavation under conditions that resembled as far as possible those in the sediment. To obtain such freshly excavated bones, two of us (EMG, MP) have collaborated closely during the last years with archaeologists and archaeozoologists to ensure that any fossils destined for ancient DNA analysis were handled in a specific way to prevent the growth of microorganisms that would destroy preserved biomolecules and/or their chemical degradation via oxidation and hydrolytic processes. To prevent dissolution and degradation of endogenous DNA and contamination by exogenous DNA we avoided the treatments normally used by archaeologists, such as washing, brushing, and treatment with consolidants and other chemicals. The fossil bones were excavated, handled (including the palaeontological analysis) and stored in an aseptic manner, thus reducing the risk of contamination with human, food-, pet-derived and environmental DNA. We will hereafter call these fossil bones “fresh” ones, as opposed to “old” fossil bones, which had been washed, handled and stored for many years in collections in a dry state at room temperature.

We obtained evidence for the detrimental effect of standard excavation treatments with fossil bones when we compared the PCR amplification success rate from a large-scale study of 247 herbivore bones, from 600 to 50,000 years old, from various depositional contexts of 60 different archaeological and palaeontological contexts in Northern and Southern Europe, the Near and Middle East and the Arabian peninsula (Table S1 and Pruvost et al., in preparation). We amplified the hypervariable region of the mitochondrial DNA using uracil-DNA-glycosylase-coupled quantitative real-time PCR (UQPCR (12, 13)) and evaluated the PCR and cloning products obtained from at least two independent fossil bone extractions. We
found that 46% “fresh” fossils yielded authenticated amplification products, whereas 18% “old” fossils yielded these products (see table S1). The difference was statistically significant (p(Chi2)=0.001). Furthermore, using UQPCR we could measure the quantity of DNA molecules. We could estimate that the number of maximal 153 bp-long molecules that could be amplified was on average about six times higher in the “fresh” fossils than in the “old” ones (see table S1). Again, despite the wide variations in DNA quantity from various bone samples, this difference was statistically significant (p(t-test)=0.043). Interestingly, while it was possible to amplify the larger DNA fragments (201 bp) from 15% of the “fresh” bones, this was the case in only 4% of the “old” bones. To conclude, we have obtained clear evidence that DNA preservation is better in freshly excavated, untreated bones and that post-excavation treatments and/or storage conditions are negatively influencing DNA preservation. Since taphonomic conditions drastically influence DNA preservation, one needs to analyze bones whose fossilization has occurred under comparable taphonomic conditions to clearly establish post-excavation conditions as the cause of DNA degradation.

We therefore studied more comparable situations, i.e, bones that had been collected under various conditions from the same preservation site, Telleilat-Mezraa, a Neolithic site in Turkey. We compared the level of DNA amplification of the hypervariable region of mitochondrial DNA from two different fossil bone groups: the “old” ones had been excavated several years before, had been subsequently brushed with water, dried and stored under light exclusion conditions in collections at room temperature; the “fresh” ones originated from the same archaeological site, but had been recently excavated according to strict protocols designed to optimize recovery of biomolecular evidence. Here, the difference in the success rate was striking: it was possible to amplify DNA from five out of eight “fresh” fossil bones (with quantities of 39,965 to 1,634 molecules/g of bone) as opposed to zero out of eleven “old” fossil bones. Thus, when the analyzed bones came from the same preservation context
suitable post-excavation conditions were important for DNA preservation. The detrimental
effect of post-excavation conditions on DNA preservation was more pronounced than when
multiple bones from multiple sites were analyzed. Yet, these bones belonged to different
individuals that had died in different ways and therefore had not experienced an identical
fossilization history, which is presumably unique for each fossil assemblage from a single
organism.

Finally, we obtained final and conclusive evidence from the analysis of exceptional fossil
material that shared the same diagenetic history but had experienced different post-excavation
histories. We analyzed ribs from an individual aurochs excavated in two different campaigns,
the first in 1947 (14) and the second in 2004. The fossil bones that were excavated in 1947
from a deep karstic crevice in Pontvallain (Pays de la Loire, France) had been stored in the
collections of the natural history museum of Le Mans (Musée Vert), France. In 2004, the
crevice was reinvestigated and 120 additional bones and teeth were recovered. Direct skin-to-
fossil contact was carefully avoided, and the specimens were immediately stored at –20°C.
The assemblage of the well-preserved bone fragments belonged to a single, adult individual.
One of the “2004 fossil bone” pieces perfectly refitted a “1947 fossil bone” fragment of the
hip of the aurochs (Figure 1). The genetic analysis of both types of fossils was carried out on
the same skeletal elements (ribs) to minimize preservation differences due to anatomical and
local geochemical differences. No PCR amplification of the bovine mitochondrial D-loop was
obtained from the ten samples from the shafts of two ribs excavated in 1947, despite
numerous attempts and a negligible inhibitory effect of the extracts. In contrast, the nine
samples from the shafts of three ribs excavated in 2004 yielded, with a success rate of 100%,
a 153 bp and a 201 bp amplification fragment of the bovine mitochondrial D-loop using
UQPCR. The quantity of DNA that was amplified from the various extracts, varied from 1 to
511 molecules/amplification reaction (average of 61 molecules/reaction ± 55.5).
From the fossil bones of the aurochs in Pontvallain we obtained an aurochs sequence that is identical to two of the sequences obtained from two older British aurochs fossils (7,500 and 11,900 years (15); see Table 1). The phylogenetic position of the retrieved sequence proves its authenticity. Thus, our study demonstrates that ancient DNA that has been preserved for thousands of years in fossil bones can be degraded relatively quickly when the bones are removed from the preserving conditions of their original setting. This degradation is not the consequence of any differences in the fossilization process, but is clearly due to changes in the macroenvironment and/or the standard handling and storage procedures in natural history collections. We then analyzed whether the different post-excavation treatments of these bone samples had left any hallmarks of distinctive morphological changes using light microscopy and environmental scanning electron microscopy with backscattered electron detectors (ESEM), which makes it possible to characterize and quantify the histological changes that occur during bone diagenesis (16, 17). Under the light microscope, the fossil bones showed no signs of human or carnivore activity. ESEM analysis of bone sections revealed that the shape and distribution of internal porosity was very similar in all analyzed specimens, with no apparent difference in texture between the fossil bones recovered in 1947 and those in 2004 (see Figure 2). Histological traits were identical in all samples and they all showed extensive bacterial attack. Neither type of fossil bone showed any cracking or exfoliation, either on the surface or at the sections. The absence of any such alterations indicates that there were no differences in the weathering stages, in the humidity/dehydration or dryness, or in compacting or deformation between the two types of fossil bones (see Figure 2B, D). No traits of specific preparation and/or conservation treatment (e.g. chemical solutions to clean the fossils or act as preservatives/consolidants (18)) were detectable. The elemental composition as revealed by wavelength and energy dispersive X-ray spectrometrical analysis (WDS and EDS, respectively) provided homogeneous spectra of the calcium and phosphate composition in
both the 1947 and the 2004 fossil bones. In summary, there is no microscopic indication that samples obtained from excavations in 1947 and those from 2004 underwent different diagenetic processes that may suggest differences in their taphonomic history. Nor are there microscopic differences between the two types of fossil bones that could be attributed to particular treatments or conditions during storage of the fossil bones after their excavation from the site of Pontvallain in 1947.

Discussion

DNA preservation occurring in post-mortem bone must be influenced by many different parameters. Approximately three different preservation phases can be distinguished. During the first diagenetic phase, the bone undergoes bacterial putrefaction. This is a rapid, complex and multi-component process. We estimate that the putrefaction phase can cause a fifteen-fold decrease in the quantity of amplified DNA, since we were able to measure $1.5 \times 10^6$ molecules/g of fresh bone and only $1 \times 10^5$ molecules/g of bones from a recent (c. 20 year-old) bone that had completed the putrefaction phase. This bone was a naturally and manually defleshed humerus from the carcass of a Batina zebu.

If, at the end of the putrefaction phase, the conditions are favorable for long-term preservation of organic matter, the bone will enter diagenetic phase 2 and DNA degradation will continue mainly on a chemical basis. One of the major DNA degradation pathways is depurination (19). This degradation reaction probably follows a first-order kinetics model in which $[A]/[A_0] = e^{-kt}$, where $[A]/[A_0]$ is the fraction of remaining material, $k$ the degradation rate and $t$ the time. We estimated the fraction of remaining material by comparing the quantity of PCR amplifiable DNA contained in both, (i) the recent post-putrefaction bones, and (ii) the fossil bones from the 3,200 year-old Pontvallain aurochs. From the freshly excavated fossil bones of the latter, we amplified 100-fold less than from the post-putrefaction bones, i.e., an
average of $2,547 \pm 5,835$ mitochondrial DNA molecules per gram of bone. This indicates that the degradation rate of DNA in the aurochs bones during burial was about 90% per 2,000 years.

The fact that we did not obtain any PCR product from other ribs belonging to the same individual but which has been unearthed 57 years ago and subsequently stored in the natural history collection, indicates that at least 99% of the DNA was degraded during this period, which corresponds to diagenetic phase 3. This means that the degradation rate was at least 70 times faster during the 57 years after excavation than during the ~3,200 year-long burial phase. The corresponding degradation rate of 90% per 30 years is comparable to that described for recent fox teeth, which had been autoclaved after the animal’s death and conserved during the first 30 years in a museum (20) and for which a DNA degradation rate of 90% per 15 years can be estimated. The slower DNA degradation rate of 90% per 2,000 years estimated for the buried aurochs bones from Pontvallain compares well with the depurination rate of DNA in solution obtained by Lindahl and Nyberg (19) applied to a 150 nucleotide-long molecule at a temperature of 5 – 10°C (temperature in the burial environment, measured during excavation) and at neutral pH (see calculations in Material and Methods). According to Lindahl and Nyberg’s measurements (19), however, an increase of 15°C in the average temperature (which we assume to be around 20-25°C in a museum) would be enough to accelerate the degradation rate 16 times. Furthermore, additional modifications of the pH and the ionic strength could further raise the DNA degradation rate to 70-fold. A decrease in the pH from 7.4 to 6.4 increases the depurination rate 3.3 times (19). Washing of the aurochs bones, which were buried in sediment of pH 7.5, with tap water, which today typically has a pH of around 5.5, is likely to decrease the pH but to an unknown extent, since the mineral part of the fossil bone has probably retained some of the buffering capacity of the bioapatite of fresh bone. A decrease in the salt content as a consequence of washing could also be
responsible for an increase in the DNA degradation rate since depurination is seven-fold faster when the concentration of NaCl is decreased from 0.1 M to 0 M (19). Finally, washing of the fossil bone could have dissolved the most soluble parts of the DNA. Thus, the rapid degradation of DNA observed in the aurochs bones after their excavation is compatible with the effects of the standard washing procedures for fossil bones that are routinely used by archaeologists and palaeontologists, combined with an elevation in temperature in the storage room.

Bone is a very heterogeneous tissue with unevenly distributed biological and physicochemical properties and bacterial attack will not be homogeneous. Furthermore, local differences in physicochemical properties should also influence long-term DNA preservation and, indeed, we and others (21) have observed local heterogeneity in DNA preservation within a bone. We were able to ascertain that the differences in DNA preservation observed with the Pontvallain aurochs bones were not due to such local heterogeneity in DNA preservation, since the preservation of DNA proved to be similar within each series of bone samples from each category (three “fresh” and two “old” ribs) and different between categories.

Whatever the exact causes, our results show very clearly how detrimental to the preservation of DNA in fossil bones standard treatments are. When classifying the 247 “fossil” herbivore bones analyzed, which all experienced differences in the diagenetic phases 1 and 2, into two categories according to the post-excavation treatment (“standard” versus “special”), the PCR success rate for the bones excavated under special conditions was twice that of those treated “normally”. Moreover, these standard excavation and storage conditions reduce the quantity of DNA by a factor of about six. This is particularly detrimental when the quantity of DNA contained in the bones is already low, as in the case of most fossil bones. In fact, treatments that result in a six-fold reduction in the DNA quantity would cause the failure
of PCR amplification in 3 of the 15 PCR-positive fossil bones excavated under special conditions (see table S1). This would be sufficient to reduce the percentage of PCR-positive bones amongst the “fresh” bones to a level not significantly different from that observed in the bone sample excavated under standard conditions. Thus, the average decrease in the quantity of amplifiable DNA could be sufficient to explain the difference in the PCR success rate.

The effect of the post-exavcation treatment was found to be even more pronounced in bones preserved in the same burial site (Telleilat-Mezraa). These bones had undergone a different diagenetic phase 1 and a similar diagenetic phase 2 and had then experienced a different post-excavation phase 3. Here we observed a striking difference in the PCR success rate amongst the “fresh”, recently excavated fossil bones (63%) compared to the ones excavated under standard conditions (0%).

Finally, when we compared the fossil bones from the same animal and the same burial site (the aurochs from Pontvallain), which had experienced differences in diagenetic phase 3 only, we also obtained a spectacular result with a 0% amplification success rate in the case of the “old” fossil bones and a 100% amplification success rate in the case of the “fresh” fossil bones. This is ultimate proof of the detrimental effect of standard post-excavation treatments of fossil bones on the “survival” of amplifiable DNA.

In conclusion, even if amplification results from collection fossil bones can be obtained, fewer fossil bones will yield PCR results and less DNA will be retrieved. This effect will be the more pronounced the less DNA is preserved in the fossil, thus leading to a North-South gradient of suitability of fossils for palaeogenetic studies, for which fossils from permafrost areas and cold caves are more suitable than those from hot and dry climate zones. Our finding has major implications for palaeogenetic studies, which are a key to the study of extinct species and populations and can reveal the mechanisms leading to extinction. We propose that
recently excavated and untreated fossil material should be preferred to fossil material that has been washed, treated with chemicals and stored for a long time in regular museum collections. Furthermore, excavation, preparation and conservation protocols and storage conditions for fossil bones in collections should be revised if genetic information is to be preserved and retrieved. If, at a given archaeological or palaeontological site, palaeogenetic results are to retain their potential to answer archaeological, palaeontological and biological questions, a selected number of the fossils should not be subjected to any treatment but instead stored in the cold - at least in a cold room but preferably in a freezer and ideally in a cryobank - in small aliquots to avoid repeated freezing and thawing cycles of the same sample. This approach calls for a close collaboration between palaeogeneticists, palaeontologists, archaeologists, conservation managers and curators.

**Materials and Methods**

**Samples, fossil excavation and storage procedure.** The aurochs bone samples (here called “fossils”) used for this study originate from a palaeontological site in France (Pontvallain, La Sarthe). Moreover, the results of a study of 247 bovine and equine bones c. 600 to c. 50,000 years old and originating from France, Germany, Switzerland, Spain, Georgia, Armenia, Turkey, Syria, United Arab Emirates, Bahrein are discussed. The specimens examined included both bones excavated using standard archaeological and palaeontological field procedures, and bones excavated under strict protocols designed for bones destined for DNA analysis. (Table S1 summarizes provenance, contexts and treatment of the specimens.)

We analyzed samples of five rib shafts (diaphyses) from the skeleton of a c. 3,200 year-old aurochs buried in a crevice in Pontvallain (La Sarthe, France), two excavated in 1947 and three in 2004. The climatic conditions in this geographic region correspond to a moderate, oceanic climate type with an annual rainfall of 678 mm (between 45 and 70 mm per month)
and temperatures range from 4°C to 19°C throughout the year. The temperature in the crevice was between 5-10°C. The fossils excavated in 1947 were kept in a dry state in cardboard boxes and drawers in the museum of Le Mans. Environmental records kept since 1995 in the museum show that the mean temperature during the year varies between 15 and 25°C and the relative humidity between 40 and 60%. The storage conditions of the bone collection before 1995 were not controlled, but were rather those of basic, uninsulated stores typical for fossil bone collections in the past and where temperature possibly fluctuated between 0° and 40°C, and relative humidity between 20 and 90%. The fossil bones newly excavated in 2004 were subjected to strict protocols, that is, specimens were handled with gloves, were not washed, brushed or treated with consolidants or other chemicals and were immediately frozen at –20°C surrounded by their sediment. The circumstances of recovery and subsequent handling of this skeleton are discussed in detail in the Results and Discussion section.

Eight of the fossil bones from the Turkish Neolithic site of Telleilat-Mezraa were excavated in 2002 using this strict protocol. Eleven “old” fossil bones from this site had been excavated between 1992 and 2003. These old bones were excavated according to standard archaeological field procedures, and had been brushed in water, sundried and then stored in cardboard boxes, first in Turkey (with fluctuations in temperature from 0°C to 30°C) and then in Southern Germany. The climate in the area of Telleilat-Mezraa is characterized by an average temperature of 16.7°C (average high temperature of 23.9°C and average low temperature of 9.3°C), annual precipitation of 21.3 mm and mean humidity of 56.2% (weather station of Bireçik, Turkey).

Samples from the diaphysis of the unburied humerus from a Batina zebu that had died naturally and undergone putrefaction was collected from the surface in Muscat in North Oman in 1983. This specimen was already naturally putrefied and almost totally defleshed (remnants
of flesh were removed manually) and was stored in the bone collection of the University of Tübingen, Germany.

A fresh cow bone was obtained from a butcher and frozen at –20°C until analysis.

**Dating of the aurochs bones from Pontvallain.** $^{14}$C dating of one bone sample gave an uncalibrated radiocarbon age of 3204 +/- 56 years. Radiocarbon dating was performed on extracted pure collagen (C/N ratio of 2.8) by the Physical Institute of the University of Erlangen/Nürnberg, Germany.

**Taphonomic analysis of the aurochs bones from Pontvallain.** The aurochs bones from Pontvallain were analyzed to identify any possible pre-burial treatment such as boiling, burning or digestion and any post-excavation (conservation) treatment with glue, resins, varnish, consolidants, or washing with alkaline, acidic or peroxide solutions, formol, alcohol or acetone as previously described (18). We carried out surface analysis of the fossil bones avoiding any such treatment and using both a binocular microscope (0.7x to 80x Leica MZ 7.5) and environmental scanning electron microscopy (QUANTA 200 Environmental Scanning Electron Microscope). The analysis of the fossils was thus possible without any additional preparation as described (18). The elemental composition of the samples and the identification of inclusions and mineralization was analyzed using wavelength dispersive X-ray and energy dispersive X-ray spectrometry (WDS and EDS, respectively).

**DNA extraction.** All pre-PCR work was carried out in a physically isolated work area in a part of the building (basement) where no other DNA work was done. The cleaning and powdering steps were performed in an area dedicated to work on fossil bones that was separate from the laboratory where DNA was extracted. For the fossil aurochs bones from Pontvallain excavated in 1947 and those excavated in 2004, the middle parts of the rib shafts were analyzed. The fossil bones were cut and the surface removed in a UV-irradiated glove box. They were then ground to a fine powder in a freezer mill (Freezer Mill 6750, Spex
Certiprep®). Further processing of the bone powder was performed in a laboratory dedicated to ancient DNA work (“fossil laboratory”) as previously described (12, 13 and supporting information). Blank extractions were carried out for each extraction series.

**DNA amplification.** PCR amplification, experiments with modern DNA and experiments with amplified and cloned DNA were carried out in three different laboratories that were not in the same part of the building as the “fossil laboratory”. To reduce the number of potential sources of error-prone sequences, we used the quantitative real-time PCR approach, UQPCR (13). Thus, for each fossil extract and PCR amplification we (i) quantified the target molecules present in a given fossil extract, (ii) diluted the fossil extract to abolish its inhibitory power as evaluated by the amplification of an external reference DNA, and (iii) destroyed with uracil-N-glycosylase (UNG) potential previous PCR and cloning products, thereby avoiding carry-over contamination. All PCR amplifications were performed in the Light Cycler® (Roche Applied Science) in individual glass capillaries using UQPCR as described (13 and supporting information). Reamplifications were never performed. Quantification of the target molecules in the extracts was performed as described (13).

A total of 29 PCRs (9 with primer pair BB1/2 and 20 with primer pair BB3/4) were performed on 10 independent extracts from samples of the shafts of two different ribs of the aurochs that had been excavated in 1947. Nine extracts from nine samples of the shafts of three different ribs of the aurochs that had been excavated in 2004 were amplified in 48 PCRs with primer pair BB3/4, and 12 PCR amplifications with primer pair BB1/2.

**Authentication of the ancient DNA sequence of the aurochs of Pontvallain.** (i) The nine DNA extractions from the rib samples of the aurochs from Pontvallain excavated in 2004 were amplified in 60 reactions using UQPCR (13) with two primer pairs from the hypervariable control region of bovine mitochondrial DNA and yielded a PCR product. Blank extractions, performed with each fossil extraction, never yielded any amplification products.
Negative controls, performed for each PCR amplification, were always negative. (ii) The initial quantity of target molecules and the quality of the results were assessed as described (12, 13) and was on average 761±715 molecules per gram of fossil powder from rib 1, 2,675±3164 molecules per gram of fossil powder from rib 2 and 4,926±9744 molecules per gram fossil powder from rib 3 of the aurochs from Pontvallain (based on the assumption that one cell contains 1,000 mitochondrial genomes). Sequences of amplification products obtained from a small number of starting molecules were compared with those starting from 100 authenticated mitochondrial molecules and found to be identical. (iii) For each fossil extract, the inhibitory effect was assessed on the basis of the decrease in the PCR efficiency and the amplification delay of modern genomic bovine DNA. The inhibiting extracts were diluted until the inhibition of the amplification reaction was abolished, i.e., in general 1:2. (iv) Prior to each PCR amplification, the products of previous PCR amplifications as well as cloned DNA were destroyed and deaminated cytosines were eliminated with UNG (13). (v) Sequencing was carried out directly on the PCR product itself and on several clones of the PCR products: 29 PCR products (20 BB3/4 and 9 BB1/2) and 34 clones (25 BB3/4 and 9 BB1/2) were sequenced. Identical *B. primigenius* sequences were obtained, except for one clone from the BB3/4 products, which showed a mutation (A -> G in position 16043). (vi) The PCR amplifications were repeated in a different laboratory (Genoscope, Evry, France) and identical sequences were obtained. (vii) A deer bone from the same excavation site yielded a *Cervus elaphus* sequence that showed two mutations when compared to the *C. elaphus* sequence that had previously been published (22). (viii) No contamination by cloned products was observed as assayed for by amplifying the PCR-positive extracts with primers hybridizing to the cloning vector on either side of the cloned fragment (13).

**Calculation of the depurination rate of DNA.** We used the classical Arrhenius formula to determine the reaction rate (*k*):
where \( R \) is the universal gas constant, \( T \) is the absolute temperature, and \( A \) is the Arrhenius constant, which relates to the geometric requirements of the reaction and must be determined experimentally. The activation energy \( E_a \) for the depurination reaction is 130 kJ/mol (19) and \( A \) can be estimated from data in Lindahl and Nyberg (19) to be \( 2.46 \times 10^{11} \, \text{s}^{-1} \) at pH 7.4. The depurination rate \( k \) at 37°C and pH 7.4 is \( 3 \times 10^{-11} \, \text{s}^{-1} \) (20) and therefore can be calculated to be \( 4 \times 10^{-12} \, \text{s}^{-1} \) at 25°C, \( 2.5 \times 10^{-13} \, \text{s}^{-1} \) at 10°C, and \( 9 \times 10^{-14} \, \text{s}^{-1} \) at 5°C. Assuming that one depurination event in a DNA target molecule within the region to be amplified is sufficient to prevent amplification, the inactivation rate of one strand of a 150 nucleotide-long DNA fragment at 10°C and pH 7.4 would therefore be \( 1.2 \times 10^{-3} \, \text{yr}^{-1} \). Using these parameters and the first order decay formula \( [A]/[A_0] = e^{-kt} \) (see Results and Discussion) it can be estimated that 90% of the DNA molecules would be inactivated in 1,900 years.

We are grateful to the archaeologists who provided access to their sites and to the archaeozoologists who provided the faunal remains (see supporting material). We thank Sophie Penet at the Genoscope, Evry, France, for the reproduction of the PCR amplifications, Olivier Ploux and Stéphane Mann for help with the synthesis of PTB, Gordon Turner-Walker for critical reading of a previous version of the manuscript, Antonia Kropfinger for corrections of the English language and several anonymous reviewers for helpful comments.

We thank Jean-Laurent Monnier et Serge Cassen, UMR 6566, Civilisations atlantiques et Archéosciences, Université Rennes 1, France, for allowing MP to obtain a fellowship of the French Ministry of Culture. RS was supported by an Erasmus fellowship and VBC by a grant from the Spanish Ministry for Education, Culture and Sports (Ap-2001-2090). This work was financially supported by the French Centre National de la Recherche Scientifique (CNRS), the Spanish Ministerio de Ciencia y Tecnologia, within the Project “Taphonomic processes:
repercussion on palaeoecological, palaeoenvironmental and biomolecular interpretations”, BTE2003-01552”, the Institut de Biodiversité (IFB) within the framework “Biodiversité et changement global” and the French Centre National d’Etudes Spatiales (CNES).
References


Legends to Figures:

Figure 1: The hip of the fossil *B. primigenius* specimen from Pontvallain showing a bone fragment excavated in 1947 and a bone fragment excavated in 2004 perfectly fitting at the line of breakage (arrow).

Figure 2: Scanning electron microphotographs from cross sections of two different fossil bone fragments from the aurochs’ ribs recovered from Pontvallain in 1947 (A, B) and in 2004 (C, D). A and C: magnification 40 x, B and D: magnification 400 x.

Table 1: Bovine DNA sequences of the mitochondrial D loop: European consensus sequence (23); African consensus sequence (24); Indian consensus sequence (24); British Pleistocene aurochs sequences (D740, D812, CHWF, NORF, CPC98, TP65) (15); PVL04, sequence obtained from the aurochs of Pontvallain (this study).

<table>
<thead>
<tr>
<th>Sequence</th>
<th>European</th>
<th>African</th>
<th>Indian</th>
<th>D740</th>
<th>D812</th>
<th>CHWF</th>
<th>NORF</th>
<th>CPC98</th>
<th>TP65</th>
<th>PVL04</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C C T G</td>
<td>T</td>
<td>A</td>
<td>C C</td>
<td>C C</td>
<td>C C</td>
<td>C C</td>
<td>C C</td>
<td>C C</td>
<td>C</td>
</tr>
<tr>
<td></td>
<td>G C T A</td>
<td>G T</td>
<td>C T G</td>
<td>G C</td>
<td>G C</td>
<td>G C</td>
<td>G C</td>
<td>G C</td>
<td>G C</td>
<td>G C</td>
</tr>
<tr>
<td></td>
<td>T G T T</td>
<td>T T T T</td>
<td>T T T T</td>
<td>T T T T</td>
<td>T T T T</td>
<td>T T T T</td>
<td>T T T T</td>
<td>T T T T</td>
<td>T T T T</td>
<td>T T T T</td>
</tr>
</tbody>
</table>

The sequences of the 3,200 year-old *Bos primigenius* and *Cervus elaphus* sequences will be deposited pending acceptance of the manuscript.
Fig. 1. The hip of the fossil *B. primigenius* specimen from Pontvallain showing a bone fragment excavated in 1947 and one excavated in 2004, perfectly fitting at the line of breakage (arrow)
Fig. 2 Scanning electron microphotographs from cross-sections of two different fossil bone fragments from the aurochs ribs recovered from Pontvallain in 1947 (A and B) and in 2004 (C and D) (A and C, x40; B and D, x400)