Ecological evolution in northern Iberia (SW Europe) during the Late Pleistocene through isotopic analysis on ungulate teeth

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Abstract

During the Late Pleistocene, stadial and interstadial fluctuations affected vegetation, fauna, and human groups that were forced to cope with these pronounced climatic and environmental changes in time and space. These changes were especially abrupt during the Marine Isotopic Stage (MIS) 3. However, little is still known about the local and regional climatic conditions experienced by hominins in Europe. Here we reconstruct the climatic trends in northern Iberia considering the stable isotopic composition of ungulate skeletal tissues found in archaeological deposits dated between 80 to 15,000 cal BP. The carbon and oxygen isotopic composition preserved in the carbonate fraction of tooth enamel provides a reliable and high-resolution proxy of the food and water consumed by these animals, which is indirectly related to the local vegetation, environment, and climate, allowing us to estimate paleotemperatures and rainfall data. This study presents 44 bovine, equid, and cervid teeth from five archaeological sites in the Vasco-Cantabrian region (El Castilló, El Otero, Axior, Labeko Koba, Aitzbitarte III) and one in the Mediterranean area (Canyars), where human evidence is attested from the Mousterian to the Magdalenian. The carbon isotope values reflect animals feeding on C3 plants with a mix-feeder diet mainly developed in open environments. However, carbon isotope value ranges point to differentiated ecological niches for equids and bovines, especially during the Aurignacian in the Vasco-Cantabrian region. Temperature estimations based on oxygen isotopic compositions and rainfall obtained from carbon isotopic compositions indicate colder and more arid conditions than nowadays from the Late Mousterian to the Aurignacian. The contemporary Mediterranean site shows slightly lower temperatures related to an arid period when animals mainly graze in open landscapes. In the Vasco-Cantabrian region, during the MIS2, the Gravettian data reflect a landscape opening, whereas the Magdalenian point to warmer conditions but still arid.

Keywords: Middle and Upper Palaeolithic; Neanderthal; Homo sapiens, palaeoecology; geochemistry

1. Introduction

Understanding the local and regional climatic evolution during the Late Pleistocene in southern Europe is crucial for assessing the potential impact of climate on the adaptation and decline of Neanderthals, as well as the subsequent expansion and resilience of Anatomically Modern Humans during the Upper Paleolithic. During the Late Pleistocene, the climatic records demonstrate stadial and interstadial continuous fluctuations during the Marine Isotope Stage 3 (MIS 3, ca. 60-27 ka) and MIS 2 (ca. 27-11 ka). Human groups had to face those episodes, which affected different vegetation and fauna depending on the region.
Northern Iberia is a key study area due to the abundance of well-preserved archaeological caves and rock shelters where, in the last decade, an updated and multidisciplinary approach has been applied to disentangle how changing environmental conditions affected the subsistence dynamics of Middle and Upper Paleolithic hominins. Recent chronological, technological, and subsistence studies are revealing a more complex panorama than previously known.

The Vasco-Cantabrian region, located in northwestern Iberia, is subject to the influence of Atlantic climatic conditions and has been widely debated as a region that was significantly impacted by the glacial-interglacial oscillations during the MIS3 (Vidal-Cordasco et al., 2022). Modelling of traditional environmental proxies (small vertebrates and pollen) from archaeological-paleontological deposits show a progressive shift in the climatic conditions with decreasing temperatures and rainfall levels detected during the late Mousterian (Fernández-García et al., 2023). Ecological alterations have been observed in large mammals, such as niche partitioning between horses and cervids, a decrease in the available biomass for secondary consumers, and consequently decrease in herbivores carrying capacity (Jones et al., 2018; Vidal-Cordasco et al., 2022).

Cold and arid conditions are maintained during the Aurignacian and the Gravettian until the onset of MIS2. Afterwards, during the Last Glacial Maximum (LGM, 23-19 ka), the global climatic deterioration associated with this glacial phase results in colder and more arid conditions in the region, with a predominance of open landscapes. However, this region still provided resources for human exploitation survival acting as a refugia area with more humid conditions in comparison to the Mediterranean area (Cascalheira et al., 2021; García-Ibaibarriaga et al., 2019a; Lécuyer et al., 2021; Fernández-García et al., 2023, Fagoaga, 2014, Posth et al., 2023). By the end of the LGM, a climate amelioration and a moderate expansion of the deciduous forest are documented from the late Solutrean through the Magdalenian (Jones et al., 2021; García-Ibaibarriaga et al., 2019a).

In contrast, northeastern Iberia is influenced by the Mediterranean climate. During MIS 3, this period has frequently been described as characterised by colder temperatures, higher rainfall compared to the present, and less pronounced climatic fluctuations when compared to the Vasco-Cantabrian region (López-García et al., 2014; Fernández-García et al., 2020; Vidal-Cordasco et al., 2022). Small-vertebrate communities and archaeobotanical evidence indicate relatively stable climatic conditions, but also the persistence of open forests during the Middle to Upper Paleolithic transition as in northwestern Iberia (Allué et al., 2018; Ochando et al., 2021). However, certain records indicate specific climatic excursions, such as increased aridity and landscape opening during Heinrich Events 4 and 5 (e.g., Álvarez-Lao et al., 2017; Daura et al., 2013; López-García et al., 2022; Rufí et al., 2018). These multi-proxy studies have significantly expanded our understanding of the environment in Iberia. However, there is still limited availability of high-resolution proxies directly linked to human activity. In this study, we propose to investigate the ecology and environmental dynamics of past ungulates during the late Middle and Upper Paleolithic by measuring the carbon and oxygen isotopic composition (δ13C, δ18O) of bioapatite carbonates preserved in archaeological mammal teeth. Tooth enamel forms incrementally and does not biologically remodel, in contrast to other body tissues such as bone, which implies that the isotope values measured on them reflect the animal diet and water sources consumed during its mineralisation, around one to two years of animal life in our study species. The preserved carbon and oxygen isotope composition in the carbonate fraction of tooth enamel offers a high-resolution record of the dietary choices of the plants and water animals consume, which indirectly reflects the vegetation, environmental conditions, and climate. This allows us to estimate past temperatures, rainfall, and moisture levels on a sub-annual scale, returning isotopic information of the foraging areas where animals were feeding during tooth formation.

By analysing the stable isotopic composition of 44 ungulate teeth obtained from 15 archaeological levels associated with human occupation, including El Castillo, El Otero, Axlor, Labeko Koba, Alitzbitarte III in northwestern Iberia, and Terrasses de la Riera dels Canyars in northeastern Iberia, this study presents novel insights into local and regional environmental and climatic trends during the Late Pleistocene (Fig.1; Fig.2; Appendix A). Specifically, it focuses on the Middle to Upper Paleolithic transition in both areas and the post-
LGM period in the Vasco-Cantabrian region. The main objectives of this work are: 1) to assess how regional environmental conditions, including changes in moisture and vegetation cover, but also temperatures and rainfall are recorded in tooth enamel stable isotopic composition; 2) to approach animal diet and their ecological niches; 3) to obtain quantitative temperature data to compare with available proxies; 4) to characterise seasonal patterns of animals found in the archaeological sites by identifying winter and summer fluctuations. The chronological resolution in the study areas for this period allows us to correlate regional paleoenvironmental changes with global records.

Figure 1. Location of the archaeological sites included in this work. From west to east, in the autonomous community of Cantabria, El Castillo, and El Otero; in the Basque Country, Axlor and Aitzbitarte III; in Catalonia, Canyars.

Figure 2. The temporal position of the archaeological levels included in the study is shown to the occurrence of different techno-complexes in both northwestern and northeastern Iberia, as well as the d18O record from the NGRIP ice core (North Greenland Ice Core Project members, 2004; Rasmussen et al., 2014). Detailed chronological information is presented in Appendix B.

2. Archaeological sites and sampled material

This study selected a total of 44 ungulate teeth including 25 bovines (Bos primigenius, Bison priscus, Bos/Bison sp.), 14 equids (Equus sp. and Equus ferus), and 5 cervids (Cervus elaphus) found in five archaeological sites in the Vasco-Cantabrian region (El Castillo, El Otero, Axlor, Labeko Koba, Aitzbitarte...
III) and one in the Mediterranean area (Terrasses de la Riera dels Canyars, henceforth Canyars. These teeth were recovered from 15 archaeological levels attributed to the following technocomplexes: Mousterian (n=14), Transitional Aurignacian (n=10), Châtelperronian (n=2), Aurignacian (n=12), Gravettian (n=1) and Magdalenian (n=5) (Table 1 and 2; Appendix B). Archaeozoological studies of the archaeological sites are available (synthesis in Marín-Arroyo and Sanz-Royo, 2022; Daura et al., 2013) and most prove that faunal remains were accumulated by human acquisition during the different cultural phases. The isotopic results of equids from El Castillo were previously published by Jones et al. (2019) in combination with the bone collagen stable isotopes on ungulate prey capture at the site, as well as bioapatite phosphate analyses of bovines from Axlor (Pederzani et al., 2023).

3. Methods

3.1 Tooth sampling

All teeth included were sequentially sampled to reconstruct the complete δ¹⁸O and δ¹³C intratooth profiles based on enamel carbonate bioapatite. Intratooth sequential sampling was applied to the second and third molars and third and fourth premolars. Bovine and horse teeth sampled exceeded 3-4 cm of crown height to ensure that at least a one-year isotopic record of animal life was obtained (Hoppe et al., 2004; Britton et al., 2019). Samples were taken perpendicular to the growth axis on the tooth where the enamel was best preserved, avoiding, whenever possible taphonomic alterations such as cracks or postdepositional damages. Samples were performed in the labial face for the lower teeth and the lingual part for the upper ones. The outermost enamel surface was abraded to remove the superficial enamel, calculus, cementum, or concretions adhering to the surface to avoid contaminations. The sequential sampling consisted of straight strips (ca. 8 x 1.5 x 1 mm) covering the width of the selected lobe, approximately every 2-3 mm, from the crown to the Enamel-Root-Junction (ERJ). The sample depth covered around 75% of the enamel depth, and dentine inclusion was avoided. A low-revolution variable-speed manual drill was used, equipped with 1 mm diamond-coated drill bits of conical and cylindrical shape. About 10-15mg of enamel powder was collected in each subsample, generating 693 subsamples for IRMS measures (see complete intratooth profiles in Appendix C).

<table>
<thead>
<tr>
<th>Site</th>
<th>Level - Cultural period</th>
<th>Bovines</th>
<th>Horses</th>
<th>Red deer</th>
<th>Teeth</th>
<th>Subsamples</th>
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<tr>
<td>Axlor</td>
<td>VI - Mousterian</td>
<td>2</td>
<td>2</td>
<td>3</td>
<td>2</td>
<td>32</td>
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<tr>
<td></td>
<td>IV - Mousterian</td>
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<td>1</td>
<td>2</td>
<td>1</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>III - Mousterian</td>
<td>4</td>
<td>4</td>
<td>6</td>
<td>6</td>
<td>62</td>
</tr>
<tr>
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<td>21 A - Mousterian</td>
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<td>1</td>
<td>3</td>
<td>4</td>
<td>47</td>
</tr>
<tr>
<td></td>
<td>20 E - Mousterian</td>
<td>2</td>
<td>2</td>
<td>4</td>
<td>6</td>
<td>56</td>
</tr>
<tr>
<td></td>
<td>18C - Trans. Aurignacian</td>
<td>4</td>
<td>4</td>
<td>6</td>
<td>66</td>
<td></td>
</tr>
<tr>
<td></td>
<td>18B - Trans. Aurignacian</td>
<td>3</td>
<td>2</td>
<td>1</td>
<td>6</td>
<td>93</td>
</tr>
<tr>
<td>Labeko Koba</td>
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<td>1</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td></td>
<td>VII - ProtoAurignacian</td>
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<td>3</td>
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<tr>
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<td>VI - Aurignacian</td>
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<td>1</td>
<td>1</td>
<td>1</td>
<td>16</td>
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<tr>
<td></td>
<td>V - Aurignacian</td>
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<tr>
<td></td>
<td>IV - Aurignacian</td>
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<td>1</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>Canyars</td>
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<td>1</td>
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<tr>
<td>El Otero</td>
<td>IV - Magdalenian</td>
<td>2</td>
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<td>5</td>
<td>68</td>
<td></td>
</tr>
<tr>
<td>TOTAL</td>
<td></td>
<td>25</td>
<td>14</td>
<td>5</td>
<td>44</td>
<td>693</td>
</tr>
</tbody>
</table>

Table 1. Number of teeth sampled by species, archaeological sites and cultural periods.

3.2 Sample treatment and stable isotope mass spectrometry

Several authors have debated the necessity of chemical pre-treatments to remove organic matter and secondary carbonates from bioapatite carbonates before stable isotopic analysis. Some chemical treatments can introduce secondary carbonates, increase carbonate content, and alter the original isotopic
signal (Snoeck and Pellegrini, 2015; Pellegrini and Snoeck, 2016). The "side effects" of these pre-treatments can compromise the final isotopic signal measured. For this reason, in this work, most of the samples were not pretreated, except for the equid samples from Labeko Koba and Altzbitarte III, and the cervids and equids from El Otero that were sampled and pretreated in the context of the initial project. Pretreatment followed was established by Balasse et al. (2002), where around 7 mg of powdered enamel was prepared and pretreated with 3% of sodium hypochlorite (NaOCl) at room temperature for 24 h (0.1 ml/mg sample), and thoroughly rinsed with deionised water, before a reaction with 0.1M acetic acid for 4 h (0.1 ml/mg sample) (equivalent protocol in Jones et al., 2019). Samples were then thoroughly rinsed, frozen, and freeze-dried. NaOCl is one of the most common agents used for pretreating carbonates and works as a base that removes organic matter by oxidation. Although it is considered one of the most efficient agents for removing organic matter, it can induce the absorption of exogenous carbonates, such as atmospheric CO$_2$ and secondary carbonates (Snoeck and Pellegrini, 2015; Pellegrini and Snoeck, 2016). It is argued that using acetic acid after NaOCl pretreatment can remove exogenous carbonates absorbed during NaOCl application. However, it is unclear if all newly introduced carbonates are finally released and which effect they produce on the original isotopic composition. While variations in pretreatment methods exist among samples in this study, the lack of a universally accepted protocol necessitates careful consideration of any potential isotopic effects resulting from these differences. These samples were analysed in the Godwin Laboratory (Department of Earth Sciences, University of Cambridge). Enamel powder samples were reacted with 100% orthophosphoric acid for 2h at 70°C in individual vessels in an automated Gasbench interfaced with a Thermo Finnigan MAT253 isotope ratio mass spectrometer. Results were reported in reference to the international standard VPDB and calibrated using the NBS-19 standard (limestone, $\delta^{13}$C = +1.95‰ and $\delta^{18}$O = -2.2‰, Coplen, 2011) for which the precision is better than 0.08‰ for $\delta^{13}$C and 0.11‰ for $\delta^{18}$O.

For the non-pre-treated samples, carbon and oxygen stable isotopic ratios were measured using continuous flow-isotope ratio mass spectrometry, specifically a Europa Scientific 20-20 IRMS coupled to a chromatograph, at the Iso-Analytical laboratory in Cheshire, UK. The samples were weighed into clean extainer tubes after being flushed with 99.995% helium. Phosphoric acid was then added to the samples, and they were allowed to react overnight to ensure the complete conversion of carbonate to CO$_2$, following the method outlined by Coplen et al. (1983). The reference materials used for VPDB calibration and quality control of the analysis included: IA-R022 (calcium carbonate, $\delta^{13}$C = -28.63‰, $\delta^{18}$O = -22.69‰), NBS-18 (carbonatite, $\delta^{13}$C = -5.01‰, $\delta^{18}$O = -23.2‰), IA-R066 (chalk, $\delta^{13}$C = +2.33‰, $\delta^{18}$O = -1.52). The accepted values of the in-house standards IA-R022 and IA-R066 were obtained by calibrating against IAEA international reference materials, NBS-18 and NBS-19, and NBS-18 and IAEA-CO-1 (Carrara marble, $\delta^{13}$C = 2.5‰, and $\delta^{18}$O = -2.4‰), respectively. Additionally, in-house standards long-term measured were used: ILC1 (calcite, $\delta^{13}$C = 2.13, $\delta^{18}$O = -3.99‰), and Y-02 (calcite, $\delta^{13}$C = 1.48, $\delta^{18}$O = -9.59‰). The analytical precision of quality control standard replicates was better than 0.09‰ for $\delta^{13}$C and better than 0.12‰ for $\delta^{18}$O.

### 3.3 Carbon stable isotopic compositions as environmental tracers

To unravel animal diet and to compare the different species, in standardised terms, it is necessary to consider the fractionation factor ($\epsilon^*$) between $\delta^{13}$C obtained by the animal on its diet and $\delta^{13}$C recorded on enamel carbonates (Bocherens, 2003; Cerling and Harris, 1999). The $\epsilon^*$ estimated for large ruminant mammals results in an offset of around 14.1‰ between diet and dental enamel, which is commonly applied generally to medium-sized herbivores. However, it is well-known that this offset varies between species, considering animals’ different physiological parameters. Recently, a formal model to predict species-specific diet-consumer isotopic offsets has been proposed, which uses body mass (BM) and digestive physiology
as the main factors that regulating the $\varepsilon^*$ (Tejada-Lara et al., 2018). This model proposes the following prediction equations for ruminant or foregut fermenters (Equation 1: Eq.1) and hindgut fermenters (Eq. 2):

$$\varepsilon^* = 2.34 + 0.05 \text{ (BM)} \quad [r^2=0.78; \text{p-value}=0.008]$$

$$\varepsilon^* = 2.42 + 0.032 \text{ (BM)} \quad [r^2=0.74; \text{p-value}=0.003]$$

In this work, we compare species with different digestive physiology, ruminants for bovines and cervids, and non-ruminants for equids. The $\varepsilon^*$ value was adjusted to each animal to avoid bias from digestive physiology when comparing these species. The following fractionation factors have been used: 14.6‰ for *Bos taurus* (Passey et al., 2005), 13.7‰ for *Equus caballus* (Cerling and Harris, 1999), and 13.2‰ for *Cervus elaphus* (Merceron et al., 2021) following (Eq. 1) for ruminants with a mean body mass of 125 kg.

Carbon isotopic composition in body tissues is considered a combination of diet (understood as consumed food), environment openness (and associated exposure to light), and the amount of precipitation. Having the precipitation in mind, Lécuyer et al. (2021) proposed to estimate Mean Annual Precipitations (MAP) from $\delta^{13}C$ preserved in enamel carbonates, derived from diets based on C3 plants. After transforming $\delta^{13}C$ from enamel carbonate ($\delta^{13}C_{\text{carb}}$) to $\delta^{13}C$ of the diet ($\delta^{13}C_{\text{diet}}$) using the fractionation factors established above, this work suggested transforming this value to $\delta^{13}C$ from vegetation ($\delta^{13}C_{\text{veg}}$). The MAP estimation is based on least square regression developed by Rey et al. (2013) and based on Kohn (2010) dataset (Eq.4), which requires first to estimate the $\delta^{13}C_{\text{veg}}$ (Eq. 3). The $\delta^{13}C$ values of atmospheric CO$_2$ ($\delta^{13}C_{\text{atm}}$) are fixed in -7‰ (Lécuyer et al., 2021; Leuenberger et al., 1992; Schmitt et al., 2012).

$$\delta^{13}C_{\text{atm}} \text{ (VPDB)} = (\delta^{13}C_{\text{carb}} - \delta^{13}C_{\text{atm}}) / [1 + (\delta^{13}C_{\text{atm}} / 1000)]$$

$$\log_1 \text{MAP+300} = 0.092(\pm 0.004) \times \delta^{13}C_{\text{atm}} + 1.148(\pm 0.074)$$

The Lécuyer et al. (2021) equation incorporates the consideration of the pCO$_2$ effect on $\delta^{13}C_{\text{atm}}$ estimation, which is expected to result in an offset of +1‰ from current levels (considering that pCO$_2$ was 180 ppm during the LGM, which is lower than the 300ppm experienced during the post deglacial around 15 ka). If this correction was not applied MAP results will be underestimated by -150mm.

### 3.4 Oxygen stable isotope compositions as environmental tracers

Intratooth profiles are known to provide a time-averaged signal compared to input isotopic signal during enamel formation (Passey et al., 2005). This signal attenuation is caused both by time-averaging effects incurred through the extended nature of amelogenesis and tooth formation, and through the sampling strategy. During mineralisation, the maturation zone, which is time-averaged, often affects a large portion of the crown height and might affect the temporal resolution of the input signal of the sample taken. To obtain climatically informative seasonal information on the analysed teeth, the application of the inverse modelling method proposed by Passey et al. (2005) is, therefore, required. This method allows us to computationally estimate the time-averaging effects of sampling and tooth formation to obtain more accurately the original amplitude of the isotopic input signal, thus to summer and winter extremes (Appendix D). This method considers parameters based on the amelogenesis trends of each species and sampling geometry, which are critical for a meaningful interpretation of intratooth isotope profiles. To evaluate the data’s reproducibility and precision, the model also estimates the error derived from the uncertainty of the sampling and the mass spectrometer measurements. This method was initially developed for continuously growing teeth, taking into account a constant growth rate within a linear maturation model, which a progressive time-average increment as sampling advances along the teeth profile. The species studied in this research exhibit non-linear tooth enamel formation, particularly in later-forming molars (Bendrey et al., 2015; Zazzo et al., 2012; Passey and Cerling, 2002; Kohn, 2004; Blumenthal et al., 2014). Although the aforementioned model is not...
ideal, as it does not account for non-linear enamel formation and certain growth parameters for the species
included are unknown, it is the best estimation based on the current state of the field and remains widely
used (Pederzani et al., 2023, 2021a, b).

Stable oxygen isotopes from meteoric water (mainly derived from rainfall) have a strong relationship with
mean air temperatures in mid to high latitudes (Rozanski et al., 1992; Dansgaard, 1964) on a regional-to-
local scale. Obligate drinkers, such as bovines and horses, acquire this water and record its isotopic
composition in their teeth and bones with a fixed, but species-specific offset (Pederzani and Britton, 2019).

Considering this two-step relationship, past climatic conditions can be estimated. However, most of the
temperature reconstructions based on δ18O have considered the δ18O from the phosphate fraction of
bioapatite enamel to build linear correlations between tooth enamel and drinking water δ18O and obtain
climatic information. For this reason, the δ18O from carbonates values obtained in this work (δ18O_carbonate) were
converted into δ18O from phosphates (δ18O_phosphate). To do so, first, to express in VSMOW notation, the δ18O
was corrected using the following correlation (Coplen et al., 1983; Brand et al., 2014):$$
\text{Eq.5} \; \delta^{18}\text{O}_{\text{carbonate}} (\text{VSMOW}) = 1.0309 \times \delta^{18}\text{O}_{\text{carbonate}} (\text{VPDB}) + 30.91
$$

Second, considering the relationship existing in tooth enamel between the carbonate and phosphate fraction
(Pederzani et al., 2011), from a compilation of the existent bibliography of modern animals measurements (Trayler and Kohn, 2017; Pellegrini et al., 2011; Bryant et al., 1996), Pederzani et
al. (2023) proposed the following correlation:
$$
\text{Eq.6} \; \delta^{18}\text{O}_{\text{phosphate}} (\text{VSMOW}) = 0.941 \times \delta^{18}\text{O}_{\text{carbonate}} (\text{VSMOW}) - 7.16
$$

Once the isotopic information is expressed in δ18O_phosphate (VSMOW), we can estimate the δ18O on meteoric
waters (δ18O_water). It is known that different physiological factors will condition how oxygen isotope composition
is fixed in each mammalian group. Thus, usually, the correlations are species-specific and developed
considering the particular physiology of each animal group. The correlation employed by this work relies on
recent data compilations (Pederzani et al., 2021b, 2023). In the case of horses (Eq. 7), it has been
considered the data combination of Blumenthal et al. (2019); Chillón et al. (1994); Bryant et al., 1994;
Delgado Huertas et al., 1995), whereas for bovines (Eq.8) the data from D'Angela and Longinelli (1990) and
Hoppe (2006) have been put together in Eq. 4. To estimate δ18O_water from red deer remains, we selected
D'Angela and Longinelli (1990) correlation (Eq. 9):

$$
\text{Eq.7} \; \delta^{18}\text{O}_{\text{water}} (\text{VSMOW}) = (\delta^{18}\text{O}_{\text{phosphate}} (\text{VSMOW}) - 22.14) / 0.87
$$

$$
\text{Eq.8} \; \delta^{18}\text{O}_{\text{water}} (\text{VSMOW}) = (\delta^{18}\text{O}_{\text{phosphate}} (\text{VSMOW}) - 22.13) / 0.85
$$

$$
\text{Eq.9} \; \delta^{18}\text{O}_{\text{water}} (\text{VSMOW}) = (\delta^{18}\text{O}_{\text{phosphate}} (\text{VSMOW}) - 24.39) / 0.91
$$

Finally, the mean annual temperatures (MAT) were calculated from δ18O_water, considering the linear regression
model relating δ18O_water and air temperatures from Iberia proposed by Fernández-García et al. (2019). This
correlation is based on monthly climatic records (monthly mean δ18O_water and monthly mean air temperatures)
from all Iberian stations from the Global Network of Isotopes in Precipitation, operated by the International

$$
\text{Eq.10} \; \text{MAT} (\degree\text{C}) = 2.38(\pm 0.10) \times \delta^{18}\text{O}_{\text{water}} + 28.19(\pm 0.58) \quad [r^{2}= 0.65; \; n=304; \; p\text{-value} >0.0001]
$$

Following Pederzani et al. (2021b, 2021a), MAT was deduced from the δ18O mean value between summer
and winter in each tooth before modeling to reduce associated error and maximise number of usable data
records. Summer and winter estimations were extracted from the obtained δ18O values after inverse
modeling application, to identify seasonal variation. Due to the uncertainties incurred from converting stable
isotope measurements to palaeotemperature, the final estimations in this work should be considered
exploratory and as a method of standardisation to make results comparable with different sites, data from
different species, and other non-isotopic palaeoclimatic records. In these estimations, the associated error
from converting $\delta^{18}$O$_{\text{pp}}$ to MAT is enlarged by the uncertainty derived from the transformation of $\delta^{18}$O$_{\text{pp}}$
(VPDB) to $\delta^{18}$O$_{\text{pp}}$ (VSMOW) (see Pryor et al., 2014; Skrzypek et al., 2016 for further discussion).

### 3.4 Present-day isotopic and climatic data

Present-day climatic conditions surrounding each site have been considered, allowing an inter-site
comparison, essential for our study but also a regional to a global perspective. Considering current MATs
and MAPs, estimated climatic data is expressed in relative terms as MAT and MAP anomalies. Present-day
summer and winter temperatures were also considered. Present-day temperatures and precipitation values
were obtained from the WorldClim Dataset v2 (Fick and Hijmans, 2017) (Appendix B). This dataset includes
the average of bioclimatic variables between 1970-2000 in a set of raster files with a spatial resolution every
2.5 minutes. The exact location of the selected archeo-palaeontological sites was used, using geographical
coordinates in the projection on modern climatic maps with QGIS software.

Present-day $\delta^{18}$Omw values from the analysed sites’ areas were obtained using the Online Isotopes in
Precipitation Calculator (OIPC Version 3.1 (4/2017); Bowen, 2022) based on datasets collected by the
Global Network for Isotopes in Precipitation from the IAEA/WMO (Appendix B).

### Table 2. Mean, minimum and maximum (Min), standard deviation (SD) of $\delta^{18}$O and $\delta^{18}$H values per
archaeological site and level organised by cultural periods. n = number of introchot subsamples measured. In tooth type: position (U, upper; L, lower); laterality (R, right; L, left); tooth (M, molar; P, premolar).

<table>
<thead>
<tr>
<th>Site</th>
<th>Level</th>
<th>Culture</th>
<th>Species</th>
<th>Tooth type</th>
<th>Code</th>
<th>$\delta^{18}$O VPDB (%)</th>
<th>$\delta^{18}$H VPDB (%)</th>
<th>n</th>
<th>min</th>
<th>max</th>
<th>SD</th>
<th>Range</th>
<th>MAT</th>
<th>MAP</th>
<th>Range</th>
<th>MAT</th>
<th>MAP</th>
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*Note: The table continues with similar entries.*
4. Results

In northwestern Iberia, specifically in the Vasco-Cantabrian region, the mean $\delta^{13}C_{\text{carb}}$ values range from -8.9‰ to -13‰, with a mean value of -11‰ (SD = 1.2‰) (Table 2; Table 3). Considering species' different fractionation factors, the $\delta^{13}C_{\text{carb}}$ were transformed in $\delta^{13}C_{\text{diet}}$, resulting in mean values that extend from -23.5‰ to -27‰ (Fig. 4). It must be considered that average values may reflect slightly different periods or be affected by seasonal bias because different teeth encompass different periods, but it has been verified in our teeth that the variations are limited when the seasonal information of the sequential sampling is incorporated (±0.2; Appendix B). The carbon isotopic composition varies between species. The bovines have generally higher mean $\delta^{13}C_{\text{carb}}$ (from -8.9‰ to -12.4‰) than the horses (from -11.3‰ to -12.6‰), whereas the red deer samples fall within the horses' range (from -11.3‰ to -13‰). Average values of $\delta^{18}O_{\text{carb}}$ in all Vasco-Cantabrian individuals extend between -3.3‰ and -7.2‰ (mean = -5.5‰; SD = 0.8‰). When transformed to $\delta^{18}O$ expected from meteoric waters ($\delta^{18}O_{\text{mw}}$), with species-adapted correlations, the $\delta^{18}O_{\text{mw}}$ values range from -3.9‰ to -9.9‰. Less clear patterns in $\delta^{18}O_{\text{carb}}$ are observed between bovines and horses, with mean values of -5.7‰ and -5.2‰, respectively. In the Mediterranean area, the site of Canyars, both species have relatively high $\delta^{18}O_{\text{carb}}$ values that fall inside the range of variation observed in the Cantabria region, between -3.8‰ and -5.5‰ in bovines and between -4.4‰ and -4.8‰ in case of horses.

<table>
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<th>Mediterranean region (NE Iberia)</th>
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<td>$\delta^{18}O_{\text{mw}}$ VSMOW (‰)</td>
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<td>Max</td>
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Table 3. Mean $\delta^{13}C$ from enamel carbonate ($\delta^{13}C_{\text{Carb}}$) and diet ($\delta^{13}C_{\text{diet}}$), and $\delta^{18}O$ from enamel carbonate ($\delta^{18}O_{\text{carb}}$) and meteoric waters ($\delta^{18}O_{\text{mw}}$), by species on the Vasco-Cantabrian and Mediterranean areas. Max: maximum value; Min: minimum value; SD: standard deviation.

4.1 Axlor

A total of seven bovine teeth were included from levels III (n = 4), IV (n = 1), and VI (n = 2) of Axlor cave. The mean $\delta^{13}C_{\text{carb}}$ range from -8.9‰ to -9.9‰ ($\delta^{13}C_{\text{diet}}$ = -23.5‰ to -24.5‰), whereas mean $\delta^{18}O_{\text{carb}}$ values are between -4.8‰ and -6.2‰ ($\delta^{18}O_{\text{mw}}$ = -5.7‰ and -7.3‰), indicating a range of variation around 1‰ and 1.4‰, respectively (Fig. 3; 4). Considering isotopic compositions by levels, mean $\delta^{13}C_{\text{carb}}$ decreases from level III to level IV, whereas mean $\delta^{18}O_{\text{carb}}$ remains stable through the sequence (Table 2; Appendix B). A range between 0.3‰ and 0.5‰ is observed in $\delta^{13}C_{\text{carb}}$ variation within tooth profiles. Individuals show clear $\delta^{18}O$ sinusoidal profiles, with peaks and troughs, and intratooth ranges from 2.1‰ to 3.4‰. The $\delta^{18}O_{\text{carb}}$ after inverse modelling intratooth profiles range from -5‰ to -6.5‰ (Appendix C; D). Mean Annual Temperatures (MATs), estimated from mean $\delta^{18}O_{\text{carb}}$ values, with seasonal control, oscillated between 10.1ºC and 14ºC (MATAs = +1.8/-2.1ºC) (Table 4). From sinusoidal profiles, summer temperatures were extracted from peaks, which are estimated to extend from 15.4ºC to 23.7ºC, and winter temperatures from troughs provided values ranging from -7ºC to 10.8ºC. Mean Annual Precipitation (MAPs), extracted from $\delta^{18}C_{\text{carb}}$, extend...
between 204mm and 326mm (MAPAs = -721/-843mm). Based on these estimations, a non-clear climatic trend is observed through these levels.

**Figure 3.** Distribution of mean carbon (δ¹³C<sub>carb</sub>) and oxygen (δ¹⁸O<sub>carb</sub>) isotopic values from enamel carbonate for species and archaeological site.

**Figure 4.** Biplot crossing δ¹³C from diet and δ¹⁸O from meteoric waters by species and sites.
4.3 Labeko Koba

This work includes teeth of bovines (n = 4), horses (n = 4), and red deer (n = 1) from levels related to Châtelperronian (IXb), ProtoAurignacian (VII), and Aurignacian (VI, V, and IV). Significant differentiation in mean δ¹³C_carb between bovines and horses is observed, with higher values between -9.3‰ and -10.4‰ in bovines (δ¹³C_carb = -23.8‰ to -25.9‰) than equids, whose values extend from -11.6‰ to -12‰ (δ¹³C_carb = -25.2‰ to -25.8‰) (Fig. 3). These horses’ values are within the ranges observed from this species in the region. Red deer have similar δ¹³C_carb values to those of horses (δ¹³C_carb = -12.3‰; δ¹³C_carb = -25.5‰). Mean δ¹³C_carb values are similar between species from -4.7‰ to -7.2‰ (δ¹³O_carb = -6.1‰ to -8.5‰). However, bovines have a very high variation within mean δ¹³O values (2.1‰), also reflected in the intratooth profiles. These δ¹³O values are lower than in other Vasco-Cantabrian sites, especially for two individuals in level VII and V (Table 3). Differences in δ¹³C values between bovines and horses result in isotopic niche differentiation between both species (Fig. 4). The red deer niche is placed within the horses' niche. The evolution of this niche over time cannot be evaluated by levels due to the limited sample. Considering the isotopic compositions by levels (Fig. 5), both bovines and horses experienced a slight increase in mean δ¹³C from levels IXb to IV, that is, from Châtelperronian to Aurignacian. Mean δ¹⁸O values of bovines decrease from VII to V, whereas in the case of horses increase from Xb to VI then to decrease from VI to IV.

Variability of δ¹³C_carb values in intratooth profiles is slightly higher (0.1-0.7‰), especially in bovines (0.3-0.9‰), with more oscillating profiles than generally flat profiles observed in horses and red deer (Appendix C; D). Intratooth profiles ranges of δ¹³C_carb are also larger within bovines (2-4‰) than in horses (1-2‰). Inverse modelled individual δ¹³O_carb ranges oscillated between 5-8‰ and 2-4‰, respectively. Sinusoidal curves are observed both in horses and bovines, but bovine profiles are noisier. The red deer has a large δ¹³O_carb range (6.3‰) from summer peak to an incomplete winter thought. We detect an inverse relation between δ¹³C and δ¹³O in some points of these individual profiles. MATs oscillated between 7°C and 13.7°C (MATas = -5.6/+1.1°C), with summer temperatures from 15.3°C to 25°C and winter temperatures from 0°C to 9.1°C. MAPs extend between 248mm and 521mm, which are notable drier conditions than nowadays (MAPas = -798/-525mm) (Table 4). Lower rainfall levels and higher seasonal amplitudes are recorded along the sequence, especially in samples from the ProtoAurignacian level VII. Relevant differences are noticed between MAPs estimated from bovines and equids, the first providing more arid conditions.

4.4 Aitzbitarte III

A single bovine individual was analysed from level V, related to Gravettian. It has a high mean δ¹³C_carb (-9.2‰) considering the observed range in bovines from the Vasco-Cantabrian region, whereas the δ¹³O_carb mean value (-5.5‰) is inside the common δ¹³O variation observed (Fig. 3). The mean δ¹³C_carb value of -23.8‰ is comparable with Canyars and some individuals from Axtior, but different from Labeko Koba and El Castillo individuals. The individual δ¹³C fluctuation is small (0.3‰) (Appendix C; D). These teeth show not quite sinusoidal profile shape in δ¹³O_carb, with an intratooth range of around 2.2‰. Climatic information is extracted but may be considered cautiously due to the profile shape and the limited sample size. From the inverse modelled mean δ¹³O_carb value (-5.4‰), we estimate a MAT of 14.5°C (MATa = +1.1°C) with a summer temperature of 17.5°C and winter temperature of 2.1°C. The MAP estimation reached 235mm (-1127mm to nowadays) (Table 4).

4.5 El Otero

Two equids and three cervids are included from level IV from El Otero, related to the Magdalenian phase. The mean δ¹³C_carb values are close, between -11.3‰ and -11.4‰ for red deer (δ¹³C_carb = -24.4‰ and -24.6‰) and -11.3‰ and -11.6‰ for horse (δ¹³C_carb = -25‰; -25.3‰) (Fig. 3). These δ¹³C values for both species are relatively high concerning other studied samples, especially for cervids (around +1.2‰). Both species have higher δ¹³O_carb values concerning the common range of variation observed in the Vasco-
Cantabria region, between -3.9‰ and -5‰ for horses and between -4.4‰ and -5.1‰ for red deer. When values are transformed to δ^{13}C_{diet} and δ^{18}O_{mw}, equids and cervids isotopic niches are separated (Fig. 4). All individuals show low-amplitude δ^{13}C_{carb} intratooth profiles (<0.3‰), but especially equids with an intratooth variation around 0.1‰ (Appendix C; D). Equids and cervids show δ^{18}O_{carb} sinusoidal profiles, with intratooth ranges between 1.4‰ and 2.4‰. Climatic estimations are proposed only for equids, providing MATs estimations from 13.4ºC to 16.7ºC (MAT_{As} = -0.3ºC/+3ºC) and MAP between 400mm and 456mm (MAP_{As} = -755/-699mm) (Table 4). A high-temperature seasonality can be seen, with summer temperatures between 19.4ºC and 22.5ºC and winter temperatures from 0.7ºC to 7.2ºC.

Figure 5. Evolution of δ^{13}C in diet (δ^{13}C_{diet}) and δ^{18}O in meteoric waters (δ^{18}O_{mw}) by archaeological levels in a diachronic order. From right to left: all species, including cervids, bovines and horses. Colors correspond to different chrono-cultures.

4.6 Canyars

From the archaeological layer I at Canyars corresponding to the Aurignacian, this work includes bovines (n = 2) and equids (n = 3) teeth. The mean δ^{13}C_{carb} values for bovines are between -9‰ to -9.3‰ (δ^{13}C_{diet} = -23.6‰ and -23.8‰), and for horses between -10‰ and -10.7‰ (δ^{13}C_{diet} = -23.7‰ and -24.4‰) (Fig. 3). In this site, the δ^{13}C_{carb} values for horses are notably higher than samples in the Vasco-Cantabrian region (around +1.2‰) (Table 3). Both species have relatively high δ^{18}O_{carb} values, but they fall inside the range of variation observed in the Vasco-Cantabrian region, between -3.6‰ and -5.5‰ in bovines and between -4.4‰ and -4.8‰ in horses. Different responses are seen in mean δ^{18}O values between the two bovines,
with one high mean value; but with close δ¹³C mean values. Bovine and equid isotopic niches overlap (Fig. 4).

All individuals show flat δ¹³C, intratooth profiles (<0.3‰ variation). Some individuals analysed do not show δ¹⁸O sinusoidal profiles, with intratooth profiles moderately flat and ranging from 1.1‰ to 1.6‰. We detect an inverse relationship between δ¹³C and δ¹⁸O in some points of bovine individual isotopic profiles. MATs oscillated between 12.5ºC and 14.8ºC (MATAs = -0.3ºC/-2.6ºC), with summer temperatures from 15.2ºC to 25.2ºC and winter temperatures from 7.3ºC to 11.4ºC (Table 4). MAPs extend between 211mm and 316mm (MAPAs = -431/-326mm). No substantial differences are noticed in the estimations based on bovines and equids because mean δ¹³C diet values differed relatively little.

<table>
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<th>Sample Level</th>
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<th>Estimated</th>
<th>Relative</th>
<th>Estimated</th>
<th>Relative</th>
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<th>Relative</th>
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Table 4. Summary of paleoclimatic estimations, based on δ¹³C for temperatures (Mean Annual Temperatures, MAT; summer; winter) and in δ¹⁸O for precipitation (Mean Annual Precipitations, MAP). Only teeth with validated seasonal curves are included in the summer and winter temperature estimation after modelling teeth peaks and trough profiles. For some profiles with an unclear seasonal shape, MATs were deducted from the original average of teeth without a seasonal profile (values marked in red). Details on teeth selection are presented in Appendix B.

5. Discussion

5.1 Diet and ecological niches: carbon ratios

Carbon isotopic ratios are valuable indicators for discerning past animal diets based on the ecosystems where the animals most frequently foraged. Considering species trends in the studied sites, bovines have generally higher mean δ¹³C values (from -8.9‰ to -12.4‰) than horses (from -11.3‰ to -12.6‰), whereas the red deer fall within the horses’ range (from -11.3‰ to -13‰). In Cenaras, in the Mediterranean area,
bovines also show higher mean δ¹³C values (-9‰ to -9.3‰) compared to horses (-10‰ to -10.7‰). These
differentiated isotopic ranges for equids and bovines can be potentially linked to feeding behaviour, but
these species are expected to present different basal δ¹³C considering their physiology and diet. Bovines,
being ruminants, have been suggested in previous studies to exhibit higher δ¹³C values due to increased
methane production (Cerling and Harris, 1999; Tejada-Lara et al., 2018). Therefore, transforming δ¹³C databy
δ¹³Cdet values is crucial to mitigate the species-specific impact, particularly when comparing ruminants and
non-ruminants. Bovines report δ¹³Cdet values between -23.5‰ and -27.5‰ and horses between -25‰ and
-26‰. These carbon compositions are typical of animals feeding on C3 plants (commonly accepted range
between -34‰ and -23‰), as can be expected from high-latitude ecosystems during the Pleistocene
(Cerling and Harris, 1999; Bocherens, 2003; Drucker, 2022).

Environmental factors such as light exposure, water stress, temperature fluctuations, salinity, and
atmospheric CO₂ changes can influence variations in δ¹³C values in a diet primarily based on C3 plants
(Kohn, 2010; Bocherens, 2003). Typically, δ¹³Cdet values below -27‰ (δ¹³Cdet = -13‰) are associated with
animals feeding on C3 vegetation found in closed forested environments, whereas δ¹³Cdet values between
-27‰ and -23‰ are linked to C3 open landscapes, which could include grasslands and steppe areas
(Bocherens, 2003). The relatively high δ¹³C observed here points to animals predominantly feeding in open
environments. The canopy effect, characterised by a depletion in ¹³C isotopes due to dense tree cover,
seems unlikely among the analysed samples since none of the individuals reported δ¹³Cdet below the
common cut-off of -27‰ (van der Merwe, 1991; Kohn, 2010; Drucker et al., 2008). Therefore, in general
terms, open mosaic landscapes, ranging from light forests to meadows and grasslands, can be inferred for
northwestern Iberia. Given the generally higher δ¹³Cdet values reported by bovines, it is likely that they were
foraging in more open environments than horses and can be considered predominantly grazers. Particularly,
bovines from El Castillo exhibit distinct feeding behavior compared to other Vasco-Cantabrian sites, as
evidenced by their lower δ¹³Cdet values, indicating a potential preference for browsing and feeding in closer
environments, possibly in lightly forested areas. Both extinct aurochs (Bos primigenius) and steppe bison
(Bison priscus) are usually classified as grass-dominant mix-feeders during the Pleistocene, although it
should be noted that modern European bison (Bison bonasus) could include browsing in their diet (Rivals
et al., 2022). For aurochs, a browse-dominated mixed feeding behaviour is also frequently described.

The δ¹³Cdet range in equids indicates feeding in open environments as well, suggesting a general mix-
feeding pattern for the Vasco-Cantabrian region. However, individuals from the Mediterranean area are
likely grazing in more open environments, as evidenced by their notably higher δ¹³Cdet values compared to
the Vasco-Cantabrian region (+1-2‰). It is important to evaluate if other factors are contributing to lower
δ¹³C values in horses. In the case of equid samples from the Vasco-Cantabrian region, it should be
considered that they have been pretreated with a combination of NaClO and acetic acid, which could
potentially affect the isotopic values. Samples after organic removal pretreatment can potentially show either
higher or lower δ¹³C values and higher δ¹⁸O values based on previous experiments (Pellegreni and Snoeck,
2016; Snoeck and Pellegreni, 2015), with δ¹³C values generally varying below 0.3‰. Based on the
observation that horses in the Vasco-Cantabrian region present lower δ¹³C values compared to bovines but
similar mean δ¹⁸O value ranges, the influence of the pre-treatment on our samples is deemed to be limited.
Furthermore, the high variability in δ¹⁸O values at El Castillo and Labeko Koba does not correlate with a
significant variation in δ¹³C values. Based on dental wear and stable isotopes analysis, Middle and Late
Pleistocene horses (Equus ferus) were primarily grazers, although some rare cases have been reported as
mixed feeders or browsers, such as at Igué des Rameaux amont or Schöningen (Kuiters et al., 2015; Rivals
et al., 2009, 2015; Uzunidis, 2020). Horse populations from northern and eastern Europe were found to be
browsers or mixed feeders, while those from the Mediterranean region tend to be grazers (Rivals et al.,
2022).
Finally, the few cervids included in this study exhibit δ\(^{13}\)C\(_{\text{diet}}\) values that frequently overlap with those of horses, indicating a mixed feeding behaviour that varies from more closed environments in El Castillo to more open habitats in El Otero. During the Pleistocene, the red deer (Cervus elaphus) exhibit a flexible, mixed-feeding behavior, consuming leaves, shrubs, forbs, grass, and sedges, similar to their present-day counterparts (Rivals et al., 2022; Merceron et al., 2021). This species inhabits diverse habitats ranging from steppes to closed temperate forests.

### 5.2 Seasonality, mobility and water acquisition: oxygen ratios and intratooth profiles

Average values of δ\(^{18}\)O in Vasco-Cantabrian individuals extend between -3.3‰ and -7.2‰ (Table 3). Even if no clear species patterns in δ\(^{18}\)O are observed, in general, bovines present slightly lower δ\(^{18}\)O values from -4.8‰ to -7.2‰ than other species; horses have a large variation from -3.3‰ to -6.6‰ and red deer from -4.4‰ to -6.8‰. In Canyars, both species have relatively high δ\(^{18}\)O values that fall inside the variation range observed in the Vasco-Cantabrian region, between -3.6‰ and -5.5‰ in bovines and between -4.4‰ and -4.8‰ in horses. Each species show different δ\(^{18}\)O intratooth ranges, with bovines between 1‰ and 3‰, equids mostly around 1.5‰, and cervids presenting the higher ranges, from 1‰ to 6‰ (Table 3; Appendix C). After applying inverse modelling to correct the dampening effect (Passey et al., 2005), the majority of teeth increase the δ\(^{18}\)O intratooth range, between 3‰ and 8‰ for bovines and 2‰ and 7‰ for horses (Appendix D). Most bovines from Axlor and Labeko Koba and equids from El Castillo and El Otero exhibit well-defined sinusoidal profiles in their δ\(^{18}\)O intratooth individual values, indicating potential seasonal fluctuations between δ\(^{18}\)O values of environmental summer and winter meteoric waters, although not all samples follow this pattern consistently. Certain intratooth profiles, particularly those from bovines in El Castillo and Canyars, exhibit sharp profiles with narrow ranges. This phenomenon was previously reported in the region in preliminary studies conducted at the sites of El Castillo (Jones et al., 2019) and in the Magdalenian levels of El Mirón cave (Geiling, 2020).

Non-sinusoidal profiles observed in the data can be attributed to various factors, including issues related to sample techniques and preservation and the inherent variability in the original isotopic signal. Factors related to sampling and methods can be connected to 1) the sampling process (e.g. too deep or too distant sampling grooves); 2) the imprecision of the mass spectrometer measurements; 3) uncontrolled effects of samples pretreatments; 4) diagenetic alterations affecting the carbonate fraction. However, it must be noted that technical reasons, whether related to sampling or pretreatment, do not appear to impact the obtained results significantly. First, this study reproduces the same intratooth sampling methods that previously yielded reliable results in similar research (e.g., Pederzani et al., 2023, 2021a). Second, non-significant alterations in intratooth profiles of treated horse samples (El Castillo, Labeko Koba, Otero) are noticed in comparison to untreated bovid samples (Appendix C). Some bovid samples are equally showing these non-sinusoidal profiles. In sites where both species are analysed, no correlation is observed between δ\(^{18}\)O and δ\(^{13}\)C. In tooth enamel, diagenetic alterations are generally less pronounced than bone due to its larger mineral content. However, carbonates within tooth enamel can be more susceptible to diagenesis and recrystallisation compared to the phosphate fraction, which contains a larger reservoir of oxygen and stronger oxygen bonds (Zazzo et al., 2004; Chenery et al., 2012; Bryant et al., 1996). The carbonate content in our samples, ranging from 3.9‰ to 8.9‰, is similar to the proportion found in modern tooth enamel, suggesting no immediate indication of diagenetic alteration. Diagenesis can also be evaluated by comparing the isotopic values of the carbonate and phosphate fractions in a sample, as there is a predictable difference between them. However, phosphate fraction measurements were still unavailable in our study. Additionally, in the case of diagenetic alteration, we would expect specimens from the same archaeological levels to be affected similarly, which is not the case.

Based on these arguments, it is suggested that the non-sinusoidal δ\(^{18}\)O signal observed in some individuals is likely attributed to the preservation of the original isotopic signature from water input. Several factors can
explain why some teeth do not reflect a clear seasonal fluctuation, which could be related to animals' mobility or the isotopic composition of the water sources. The main factors considered in our study are 1) the high mobility of the animals analysed among ecosystems with different isotopic baselines due to large migrations; 2) the inland-coastal or short altitudinal movements through the region, which lead to the acquisition of water from sources with different isotopic signal; and 3) the acquisition of water from sources with no clear seasonal signal, such as large bodies of water, rivers, groundwaters, or meltwaters. Analyses of nitrogen and sulphur stable isotopes on unglue bone collagen from Axlor, El Castillo and Labeko Koba (Jones et al., 2019, 2018; Pederzani et al., 2023) have already revealed large variation ranges linked to the existence of several microenvironments just in a few kilometres within the Vasco-Cantabria region. Long migrations and long hunting distances cannot solely explain these diversified values. In our study, the minimal δ¹³C intratooth variation within individuals (<1‰) indicates limited seasonal changes in their feeding behavior that influenced the carbon isotopic composition (Appendix C). Therefore, considering the Vasco-Cantabrian diverse orography with perpendicular valleys that connect the Cantabrian Cordillera with the Atlantic Ocean through rivers over short distances (30-50 km), the availability in the past of a wide range of water sources in small areas seems highly likely. Furthermore, rivers in the region frequently contain meltwater from snow during the winter-spring months, and water springs are also common.

5.3 Regional trends and ecological niches

This study provides valuable insights despite the limited sample size at each archaeological level. It establishes a baseline of isotopic values for northern Iberia, allowing for the evaluation of regional trends. In the northwest, in the Vasco-Cantabrian region, the δ¹³C values obtained oscillated between -8.9‰ and -13‰ and between -3.3‰ and -7.2‰ in the case of δ¹⁸O values. These values are within the range expected, considering previous regional studies in ungulates (Lécuyer et al., 2021; Pederzani et al., 2023; Jones et al., 2019; Carvalho et al., 2022). Although oxygen variability trends are less precise, the main factor distinguishing the observed changes over time is the variation of carbon isotopic composition among species and regions. The combination of mean δ¹³C and δ¹⁸O values (Fig. 4; 5) accentuates disparities in ecological niche overlap between horses and bovines, whereas cervids and horses frequently exhibit shared ecological niches. The dissimilarities between bovines and horses could be attributed to shifts in feeding behavior, which may be accompanied by ecological and environmental changes, either independently or in parallel.

Upon evaluating the entire dataset by sites, the consistently lower δ¹³C values in horses compared to bovids throughout time suggest both animals inhabited open landscapes, with bovines exhibiting a grazer preference while horses show a mix-feeding diet. Only in the Middle-to-Upper Paleolithic transition 18B and 18C levels of El Castillo, an exception is observed with lower δ¹³C values in bovines, linked to a higher browser input due to a higher habitat in closer environments, such as open forests, similar to those inhabited by the horses. This generates a niche overlapping between horses and bovines, most likely reflecting stable conditions that could support both species in similar ecosystems. Contrarily, in the Châtelperronian and early Aurignacian levels from Labeko Koba, a clear differentiation between horses and bovines is observed, mainly in δ¹³C values, highlighting the occupation of different landscapes by both species. This niche fractionation between species could result from resource competition derived from an unstable climatic period, where species needed to specialise to adapt to the changing conditions. Notable changes are also observed in the δ¹⁸O values from Labeko Koba compared to the older El Castillo and Axlor sites, with bovines exhibiting a higher fluctuation range and the lowest values in the region. These trends are consistent with values observed on bone collagen from previous studies in the studied sites. During the Middle-to-Upper Paleolithic transition in the region, by comparing horses and red deer, a decrease in mean δ¹³C (from -20‰ to -21‰) and δ¹⁸O values (from 6‰ to 2.5‰) was observed in contrast to stable red deer mean δ¹³C (Fernández-García et al., 2023; Jones et al., 2018, 2019). This decrease was already interpreted as niche
fractionation, derived from an opening landscape, that drove equids into low-quality pastures compared to cervids. Pollen evidence in the region suggests a prevalence of steppe vegetation and low tree cover for the Châtelperronian and Aurignacian (Iriarte-Chiapusso, 2000).

In the same period, at the Mediterranean site of Canyars, higher mean δ¹³C
values are observed in both species (between -23.6‰ and -24.4‰), indicating a preference for more open landscapes by bovines and equids.

The indication of open areas could be linked to the arid climatic conditions associated with the Heinrich Event 4, which coincides with the formation of the archaeological level at Canyars. This predominance of open areas coincides with the presence of typical steppe herbivore species, such as Equus hydruntinus and Coelodonta antiquitatis, the microfauna and pollen taxa, and use-wear analysis on ungulate remains identified at the site (Daura et al., 2013; López-Garcia et al., 2022; Rivals et al., 2017). Aridity is a plausible explanation for the higher niche partitioning observed in Labeko Koba and the higher δ¹³C values found in Canyars for both species in Aurignacian levels. The δ¹³C results of bovines from Aitzbitarte III during the Gravettian are consistent with the trend observed in Labeko Koba, and previous studies have already suggested this time to be notably arid and cold. Finally, in the Magdalenian level of El Otero, higher δ¹³C values resemble those observed in Canyars. However, this time, carbon values are related to niche partitioning between horses and red deer. In contrast, higher δ¹⁸O values might indicate warmer conditions but are still associated with open landscapes in the Vasco-Cantabrian area.

### 5.4 Late Pleistocene climatic evolution in Northern Iberia

Carbon and oxygen isotopes were used to estimate quantitative parameters related to past temperatures and precipitation. In the case of oxygen isotopic compositions, an evaluation of environmental water composition can be addressed before approaching temperature estimations. When transformed to δ¹⁸O

Using species-adapted correlations, the summer δ¹⁸O values obtained from the modelled teeth range from -1‰ to -4.9‰, while the winter values range from -4.2‰ to -10.6‰. These values agree with the current δ¹⁸O range of values recorded by the IAEA station (IAEA/ WMO, 2022) in Santander (from -3.5‰ in summer to -6.6‰ in winter) and in Barcelona (from -2.2‰ in summer to -6.3‰ in winter) and the OIPC (Bowen, 2022) estimations for studied locations (from -1‰ to -9‰) (Appendix B). As observed in the present, Canyars exhibit higher mean annual δ¹⁸O values of -5.5‰, which are close to the current δ¹⁸O estimated for this location (-5.4‰). This suggests that the baseline δ¹⁸O differences between Canyars and the other sites can be primarily attributed to the Mediterranean influence rather than the Atlantic, assuming equivalent air circulation patterns in the past as experienced in the present and considering IAEA stations.

Considering this work’s climatic reconstruction, temperatures are generally colder and precipitation levels are notably lower than nowadays (Table 4; Appendix B). From 80,000 to 57,000 cal BP, in the Mousterian levels of the Axlor site, temperatures are generally colder than today, but there is no clear trend observed throughout the sequence, with older levels showing higher differences between summer and winter temperatures. Rainfall estimations in these levels exhibit an unusual arid pattern, possibly affected by bovines mainly feeding in open areas. Indeed, even if the species’ impact has been corrected through δ¹³C diet estimation (Tejada-Lara et al., 2018), this study observed that the influence of basal feeding behavior on rainfall estimations previously advised by Lécuyer et al. (2021) should be considered. In this case, it is not possible to isolate the effect of diet from environmental interference, but previous studies have highlighted stable climatic conditions for the site (Pederzani et al., 2023).

During the Middle-to-Upper Paleolithic transition and early Aurignacian occupations, a shift in the niche configuration of species is observed, indicating climatic perturbations. There is a decreasing trend in temperatures from the Transitional Aurignacian levels in El Castillo (18C and 18B; ca. 47,000 cal BP) to the Châtelperronian (Xinf; 42,100 cal BP) and Early Aurignacian (VII-V; from 41,136 to 38,570 cal BP) levels in Labeko Koba. Lower mean annual and winter temperatures are particularly notable in El Castillo, while...
Labeko Koba exhibits high seasonal amplitude, especially in level VII. Additionally, there is a slight decrease in rainfall and increased fluctuations from the Transitional Aurignacian levels in El Castillo (18B-18C) to the Aurignacian levels in Labeko Koba (VII-V). Previous studies in the northern Iberian region underlined an environmental and ecological shift after GS13/HE5, from 48,000 to 44,000 cal BP, based on a progressive trend to colder temperatures, aridity increase, and open environmental conditions, matching with the late Neanderthal occupations, followed by a population hiatus before the arrival of Anatomically Modern Humans (Jiménez-Sánchez et al., 2013; Ruiz-Fernández et al., 2022). Moreover, previous isotopic analyses in the region pointed to some ecological alterations considering perturbations observed in the δ13C and δ18O of bone collagen (Jones et al., 2019, 2018).

When comparing the environmental reconstruction of the Aurignacian period in the Vasco-Cantabrian region (levels V-IV from Labeko Koba) and the Mediterranean region (Layer I from Canyars), which are synchronous to HE4 (39,000 BP), the study reveals notably lower rainfall levels for the Mediterranean, consistent with the feeding behavior described for animals mainly feeding in open areas. These drier conditions align with the unique climatic expectations for this period and support previous findings suggesting aridity and the predominance of open landscapes (Rivals et al., 2017; Daura et al., 2013). The temperature data indicates that Canyars experienced colder conditions, especially during the winter season, compared to the present. However, in comparison to Labeko Koba, Canyars experienced warmer conditions. As explained earlier, the Mediterranean basin had consistently higher temperatures, even during colder periods. In line with this, previous studies conducted at the site have also highlighted the persistence of Mediterranean open forests in the surroundings (Rivals et al., 2017; López-García et al., 2013).

Finally, the sites Aitzbitarte III (26,692 cal BP) and El Otero (19,303 cal BP) provided valuable climatic insights into the Vasco-Cantabrian region during the Upper Paleolithic, specifically during the Gravettian and Magdalenian periods, respectively. Considering previous research in the region, the climatic trend reported for the Aurignacian, characterised by colder and more arid conditions, was expected to continue or even intensify during the Gravettian period (Fernández-García et al., 2023; Garcia-Ibarrarriaga et al., 2019b; Lécuyer et al., 2021). Both sites exhibit lower precipitation levels compared to the present, indicating significant aridity, supported by the animals predominantly feeding in open landscapes. El Otero’s higher mean annual temperatures recorded in the Magdalenian horses are consistent with a climatic amelioration following the Last Glacial Maximum (Jones et al., 2021).

5. Conclusions

This study provides a comprehensive analysis of the temporal evolution of the environment and climatic conditions in northern Iberia, spanning from the late Middle Paleolithic to the late Upper Paleolithic, this is from the GS21 to the GS2, ranging from 80,000 to 15,000 cal BP. In the Vasco-Cantabrian region, the results reveal a heterogeneous landscape characterised by an open mosaic, ranging from light forest to meadows and grasslands. This landscape reconstruction is primarily influenced by the feeding locations of the studied animals and, consequently, related to the ecosystems where hominins captured them. Despite shifts in niche configuration observed between equids and bovines, both species typically forage in open areas, with bovines showing a higher preference for grazing. Only in El Castillo, during the late Mousterian and the Transitional Aurignacian levels, bovines show unusually low δ13C related to higher browsing and overlapping with horse isotopic niche. This might indicate a slightly closed mosaic landscape that could sustain both species. In contrast, only horses from Canyars exhibit a preference for grazing behavior.

Stable climatic conditions are described for Mousterian Axlor and El Castillo levels from 80,000 to 50,000 BP. However, some elements indicate some environmental perturbations initiated during the Transitional...
Aurignacian levels from El Castillo, around 48,000 BP and after HE5/GS13. After GS12 (44,200-43,300 BP), horses and bovines are potentially occupying different ecological niches during the Châteleperronian and early Aurignacian levels from Labeko Koba, pointing to a species' environmental specialisation, which can be a consequence of competition for food resources during an unstable ecological period. The climatic estimations indicate a temperature shift during this period, with a slight decrease in temperatures and evidence of fluctuations in rainfall from the Transitional Aurignacian levels of El Castillo to the early Aurignacian levels of Labeko Koba. Previous environmental studies on the region have underlined ecological stress and aridity increase around 48,000 cal BP, which could sustain this biological impoverishment. When comparing the environmental conditions during the Aurignacian period in the Mediterranean (Canyars) and the Vasco-Cantabrian region (Labeko Koba), the Mediterranean area had higher baseline temperatures but also experienced higher aridity. Animals continued to feed on open landscapes during the Gravettian and Magdalenian levels in the Vasco-Cantabrian region, represented by Altzbitarte III and El Otero sites. However, there is evidence of a temperature recovery after the LGM at the El Otero site.

For the first time, a regional approach is obtained by measuring δ¹³C and δ¹⁸O in enamel carbonates from ungulates teeth for the late Middle and Upper Paleolithic in northern Iberia. Stable isotope composition of oxygen and carbon from ungulate teeth has provided valuable insights into the diet and foraging areas of bovines, equids, and cervids. These results, derived from this extensive sampling in the Vasco-Cantabrian, establish the basis of stable isotopic studies on teeth enamel, which were slightly explored in the region. Despite the uncertainties inherent in this work, derived from using the carbonate enamel fraction for paleoclimatic estimations, both δ¹⁸O and δ¹³C contributed to the regional climatic characterisation, including the estimation of temperatures and precipitations, as well as the seasonality range between summer and winter. The potential influence of pretreatment effects and uncontrolled diagenetic alterations on the enamel carbonate fraction has been assessed. However, further investigation using new techniques like phosphate analysis and FTIR are needed to gain more insights into sample preservation. Ongoing sulphur, hydrogen and strontium studies will provide additional information on the animal mobility patterns consumed for hominins and, therefore, landscape exploitation through this transition between late Neanderthal and early modern human groups. A more comprehensive characterisation of the baseline oxygen values in the region would also enhance the environmental interpretation of the existing data.

Appendices

Appendices A, C and D are presented after bibliography. Raw data is found in Appendix B available at https://github.com/ERC-Subsilence/Ungulate_enamel-carbonate

Code availability

R code used to perform plots and models in this manuscript can be accessed at GitHub (https://github.com/ERC-Subsilence/Ungulate_enamel-carbonate).

Data availability

The available datasets used for this article are provided in the supplementary materials (Appendix A-D).

Author contribution

responsible for the excavations in Canyars and contribute to the discussion. All the authors revised and commented on the manuscript.

Competing interests

The contact author has declared that none of the authors has any competing interests

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References


Fagoaga, A.: Aproximación paleoecológica y paisajística durante el MIS3 a partir del estudio de los micromamíferos del yacimiento de El Salt (Alcoi, Alicante), University of Burgos, 34 pp., 2014.


Appendix A. Sites description

A1. Vasco-Cantabrian sites

Axlor (Dima, Vizcaya, País Vasco)

Axlor is a rock-shelter located in Dima (43.2706; -1.8905), with a continuous Middle Paleolithic sequence from the MIS5 to the MIS3 (DeMuro et al., 2023; Pederzani et al., 2023; Marín-Arroyo et al., 2018). It is placed on the southwestern slope of the Dima valley, with an elevation of approximately 320 m above sea level (a.s.l.), at 33 km straight from the present-day coastline, next to one of the lowest mountain passes linking the Cantabrian basins and the Alavese Plateau. The site was discovered in 1932 and initial excavations were performed by Barandiarán (1967-1974). J. M. Barandiarán undertook the excavations between 1967 and 1974, identifying eight Mousterian levels (I-VIII) (Barandiarán, 1980). From 2000 to 2008, new excavations by González-Urquijo, Ibáñez-Estévez and Ríos-Garaizar were achieved and, since 2019, these are ongoing by González-Urquijo and Lazuén. Due to the lack of chronology during Barandiarán excavations, among other aspects, work was focused on obtaining a detailed stratigraphy on the new excavation areas to correlate it with Barandiarán’s levels (González-Urquijo & Ibáñez-Estévez, 2021; González Urquijo et al., 2005). The new stratigraphic sequence is roughly equivalent to the previous one, but with additional levels, not previously identified or excavated by Barandiarán. Some of these levels were deposited before Level VIII (Gómez-Olivencia et al., 2018; 2020). The Middle Paleolithic sequence extends from layers VIII to III (or from N to B-C). Levallois production is predominant in the lower levels (VI to VIII), while Quina Mousterian technocomplex does in the upper ones (from III to V) (Ríos-Garaizar, 2012, 2017). New chronological data by radiocarbon and OSL methods confirm that a sequence Axlor levels VI, VIII, and VIII probably accumulated during MIS5d–a (109–82 ka), while levels D to B probably were formed during the period encompassing the start of MIS 4 (71–57 ka) through to the beginning or middle of MIS 3 (57–29 ka) (Demuro et al., 2023) and upper Level III to 46,200 ±3,000 BP, which calibrates between 45,510 cal BP and the end of the calibration curve at > 55,000 cal BP (see Pederzani et al., 2023: Fig. 1).

The archaeozoological study indicate an anthropic origin of the faunal assemblage with scarce carnivore activity documented (Altuna, 1989; Castaños, 2005; Gómez-Olivia et al., 2018). In lower layers, the most abundant taxa are Cervus elaphus (VIII) and Capra pyrenaica (VII), while in upper layers III-V. Cervus elaphus is substituted by Bos primigenious/Bison priscus and Equus sp. The material included in this work comes from the faunal collection of the Barandiarán excavation currently curated at the Bizkaia Museum of Archaeology (Bilbao) where teeth were sampled and the stable isotope analyses on enamel phosphate were included in Pederzani et al. (2023).

El Castillo (Puente Viesgo, Cantabria)

El Castillo is cave located in Puente Viesgo (43.2924; -3.9656), with an elevation of approximately 195 m a.s.l., at 17 km straight from the present-day coastline. The cave belongs to the karstic system that was formed in the Monte Castillo, which dominates the Pas valley. The site was discovered in 1903 by H. Alcalde del Río. H. Obermaier carried out the first excavation seasons between 1910 and 1914, when many of the archeological remains were recovered, mainly from the hall of the cave. These interventions were done under the supervision of the “Institut de Paleontologie Humaine” (IPH) and of Prince Albert I of Monaco. From 1980 to 2011; V. Cabrera and F. Bermudo de Quiros undertook new excavations focusing on the cave entrance, on the Middle to Upper Paleolithic transitional levels, mainly 16, 18 and 20 (Cabrera-Valdes, 1984). The site has yielded an important stratigraphic sequence, composed by 26 sedimentological units.
related to different anthropic occupational units, often separated by archeologically sterile units:


Unit 21 is mostly sterile (Cabrera Valdés, 1984; Martín-Perea et al., 2023) and it was dated by ESR yielding a mean date of 69,000 ± 9,200 years BP (Rink et al., 1997). However, Martín-Perea et al. (2023) suggested some dating uncertainty arising from the interpretation of the initial stratigraphic nomenclature. They suggest that the ESR dates provided for level 21 by Rink et al. (1997) were erroneously attributed to this unit and it might correspond to 20E indicating that below that subunit the chronology is older than 70,000 years BP (Martín-Perea et al., 2023). The Mousterian Unit 20 cave is divided into several subunits (Martín-Perea et al., 2023). In Unit 20, a cave roof collapse took place transforming the cave system into an open rock shelter. This unit contains abundant archaeological and paleontological remains. Lithic industry consistent in sidescrapers, denticulates, notches and cleavers, the majority on quartzite and presents both unifacial, bifacial discoid debitage and Levallois debitage. Unit 20E was attributed to Quina Mousterian by Sánchez-Fernández and Bernaldo De Quiros (2009) and contains a Neanderthal tooth remain (Garralda, 2005).

Considering the geochronological uncertainties for dates on 20E related with Rink et al. (1997), we have decided to solely rely on ESR date of 47,000 ± 9400 BP provided by Liberta et al. (2010) for this level. Unit 20C presents clear evidence of the Mousterian lithic industry and radiocarbon dates of 48,700±3,400 uncal BP (OxA-22204) and 49,400±3,700 uncal BP (OxA-22205) (Wood et al., 2018) and mean ESR date of 42,700 ±9800 BP (Liberta et al., 2010). Level 19 is archaeologically sterile and separates Unit 20 from Unit 18 (Wood et al., 2018).

Unit 18 is divided into three parts: 18A (archaeologically sterile), 18B, and 18C. Levels 18B and 18C were classified as Transitional Aurignacian, representing a gradual transformation from the Mousterian to the Aurignacian, which is unique to El Castillo cave (Cabrera et al., 2001; Maillo and Bernaldo de Qiúros, 2010; Wood et al., 2018). The dates and the cultural attribution of these levels have been the subject of much debate (e.g. Zilhão and D’Errico, 2003; Wood et al., 2018). According to Wood et al. (2018), the last dates of these levels range between 42,000±1,500 uncal BP (OxA-22203) and 46,000±2,400 uncal BP (OxA-21973), which is much earlier than the start of the Aurignacian period in the Cantabrian region (Marín-Arroyo et al., 2018; Vidal-Cordasco et al., 2023). The lithic assemblage of Unit 18 appears to be dominated by Discoid/Levallois technology (Bernaldo de Qiúros and Maillo-Fernández, 2009) but with a high percentage of "Upper Paleolithic" pieces. Additionally, punctual bone industry, as well as pieces with incisions and engravings, were discovered in Unit 18 (Cabrera-Valdés et al., 2001). Three deciduous tooth crowns attributed to Neanderthals were found in Unit 18B (Garralda et al., 2022). Above, Unit 17 is sterile but contains scarce lithic and faunal materials, while Level 16 was attributed to the Proto-Aurignacian, with dates of 38,600±1,000 uncal BP (OxA-22200) (Wood et al., 2018).

According to Luret et al. (2020), there was a shift in hunting practices between the Late Mousterian (unit 20) and the Transitional Aurignacian (unit 18). During the Late Mousterian, hunting strategies were less specialized, and the species hunted included red deer, horses, and bovines. However, in Unit 18, a specialization in red deer hunting is observed. However, the explanation of this shift has been proposed as a response to a cultural choice or induced by climatic changes. The material included in this work comes from the faunal collection recovered during the Cabrera-Valdés and Bernaldo de Qiúros excavations curated at Museo de Prehistoria y Arqueología de Cantabria (MUPAC, Santander).

**Labeko Koba (Arrastre, Guipúzcoa, País Vasco)**

Labeko Koba is a cave located in the Kurtzetxiki Hill (43.0619; -2.4833), at 246 m a.s.l. and 29 km straight from the present-day Atlantic coast. In 1987 and 1988, due to the construction of the Arrasate ring road, the...
site was discovered and a savage excavation was carried out (Arrizabalaga, 2000a). Unfortunately, the site was destroyed after that. The stratigraphic sequence identified nine different levels. The lower Level IX was attributed to the Châtelperronian, based on the presence of three Châtelperron points. Although there is a lack of human remains in few Cantabrian Châtelperronian sites, recent research has suggested that this techno-complex was produced by Neanderthals (Maroto et al., 2012; Rios-Garaizar et al., 2022). Level VII marks the beginning of the Aurignacian sequence, likely Proto-Aurignacian, with a lithic assemblage dominated by Dufour bladelets (Arrizabalaga, 2000a). Levels VI, V, and IV contain lithic assemblages that suggested an Early Aurignacian attribution (Arrizabalaga, 2000b; Arrizabalaga et al., 2009). This site is significant because it is one of the few sites with Châtelperronian assemblages and with both Proto-Aurignacian and Early Aurignacian separated (Arrizabalaga et al., 2009).

Initial radiocarbon dates were inconsistent with the stratigraphy of the site and much more recent than expected for the Early Upper Paleolithic (Arrizabalaga, 2000a). This incoherence was determined to be affected by taphonomic alterations (Wood et al., 2014). Later radiocarbon dates undertaken with an ultrafiltration pre-treatment provided a new regional framework for the regional Early Upper Paleolithic (Wood et al., 2014). The Châtelperronian layer is dated to 38,100±900 uncal BP (OxA-22562) and 37,400±800 uncal BP (OxA-22560). The Proto-Aurignacian levels cover a period from 36,850±800 uncal BP (OxA-21766) to 35,250±650 uncal BP (OxA-21793). The three Early Aurignacian levels are dated to 35,100±600 uncal BP (OxA-21778) for level VI, ~ 34,000 uncal BP (OxA-21767 and OxA-21779) for level V, and ~ 33,000 BP (OxA-21768 and OxA-21780) for level IV (Arrizabalaga et al., 2009).

Taphonomic studies indicate an alternation in the use of the cave between carnivores and humans, the latter ones during short occupation periods (Villaluenda et al., 2012; Rios-Garaizar et al., 2012; Arrizabalaga et al., 2010). Labeko Koba is considered to have functioned as a natural trap where carnivores, mainly hyenas, were accessed to animal carcasses. At least in the base of Labeko Koba IX, carnivore activity was higher, and they would have consumed the same prey as humans (Villaluengua et al., 2012). The presence of humans is linked to strategic use as a campsite associated with a small assemblage of lithic artifacts. The most consumed species by Châtelperronian groups were red deer, followed by the consumption of large bovids, equids, and woolly rhinoceros. During the Aurignacian period, there was some stability in human occupations, although still alternated with carnivore occupations (Arrizabalaga et al., 2010). Cold-adapted fauna such as reindeer and woolly rhinoceros were identified in association with the Châtelperronian. Reindeer were still present during the Aurignacian levels, as well as the woolly mammoth and arctic fox. The original sampling of the studied teeth by this work was performed in the San Sebastian Heritage Collection headquarters where the Guipuzcoa archaeological materials were deposited at that time.

Aitzbitarte III (Rentería, Guipúzcoa, País Vasco)

Aitzbitarte III is an archaeological site located within a karstic system comprising of nine caves in Rentería (43.270; -1.8905). The cave is situated 220 m.a.s.l. and is 10 km away from the present-day coastline. Initial archaeological interventions were carried out at the end of the 19th century by P.M. de Soraluce (Altuna, 2011). Recent excavations were initially conducted in the deep zone inside the cave between 1986 and 1993, and later focused on the cave entrance between 1994 and 2002, by J. Altuna, K. Mariezkurrena, and J. Ríos-Garaizar (Altuna et al., 2011; 2017). While the cave’s entrance area contains a sequence comprising Mousterian, Evolved Aurignacian, and Gravettian layers (Altuna et al., 2011; 2013), the stratigraphy in the inner cave presents 8 levels: level VIII (some tools with Mousterian features), VII (sterile), VIb, VIa and V (Middle Gravettian technocomplex with abundance of Noailles burins), IV-II (disturbed archaeological levels) and I (surface) (Altuna et al., 2017). Levels V have dates of 24,910 uncal BP (I-15208) and 23,230 uncal BP (Ua-2243); whereas level VI extends...
from 23,830 ± 345 uncal BP (Ua-2628) and 25,380 ± 430 uncal BP (Ua-2244) (Altuna, 1992; Altuna et al., 2017), with a possible outlier dated at 21,130 uncal BP (Ua-1917).

The Gravettian occupation in the inner part of the cave was originally thought to be more recent than the ones in the cave entrance. However, it was difficult to correlate the two excavation areas due to different sedimentation rates. The rich human occupations took place during a singular cold phase in the Middle Gravettian with a specialized paleoeconomy focused on the hunting of Bos primigenius and Bison priscus (85% in level VI and 68% in level V), which is unusual in the Cantabrian region mostly focused on red deer and ibex. Other ungulates present are Cervus elaphus and Rupicapra rupicapra, and to a lesser extent Capra pyrenaica, Capreolus capreolus, Rangifer tarandus, and Equus ferus (Altuna et al., 2017; Altuna & Mariezkurrena, 2020). There is a scarce representation of carnivores. The tooth studied was sampled at the Gordalúa Center for Heritage Collections of the Provincial Council of Gipuzkoa.

**El Otero (Secedura, Voto, Cantabria)**

El Otero cave is located in Secadura (Voto) (43.3565; -3.5360), at 129 m.s.a.l and 12 km straight from the present-day coastline. Near the Matienzo valley in a coastal plain environment covered by meadows and gentle hills. The discovery was made in 1908 by Lorenzo Sierra. The site was excavated in 1963 by J. Gonzalez Echegaray and M. A. Garcia Guinea, in two different sectors (Sala I and Sala II) with an equivalent stratigraphic sequence (González Echegaray, 1966). A total of nine levels were identified in Sala I, from level IX to level I. Levels IX and VIII were originally related to the "Aurignacian-Mousterian, based on lithics assemblages with a combination of both technocomplex features. The overlying levels VI-IV were separated by a speleothem crust (level VII) and were initially related to Aurignacian, due to the presence of end-scrapers, bone points, blades, or burins on truncation (Freeman, 1964; Rios-Garaizar, 2013). Also, perforated deer, ibex, and fox teeth were found in levels V and IV. This site lacked chronological dating methods, until a selection of material from levels VI, V and IV revealed a difference chrono-cultural attribution (Marín-Arroyo et al., 2018). Radiocarbon results yielded younger dates for such a cultural attribution and show significant stratigraphic inconsistency. Level VI gave a result of 12,415 ± 55 (OxA-32585), two dates in Level V are 12,340 ± 55 (OxA-32509) and 10,585 ± 50 (OxA-32510) and a date in Level IV is 15,990 ± 80 (OxA-32508). All these results fall into the range of the Late Upper Paleolithic (Magdalenian-Azilian initially identified in levels III-I), eliminating attribution of these levels to the Aurignacian, despite the presence of apparently characteristic artefacts.

Red deer dominate the assemblage, except for level IV where horses are more abundant. Wild boar, roe deer, and ibex are also present, but large bovids are relatively rare (González Echegaray, 1966). Level IV is the richest and most anthropic level, with evidence of butchering in red deer (captured in winter and early summer) and chamois (in autumn) The formation of this level involved humans and carnivores, and although certain data may suggest an anthropic predominance, the limited sample analyzed taphonomically and the pre-selection of preserved pieces do not allow for a definitive conclusion (Yravedra & Gómez-Castaneda, 2010). The material included in this work is curated at the Museo de Prehistoria y Arqueología de Cantabria (MUPAC, Santander).

**A2. Mediterranean sites**

**Terrasses de la Riera dels Canyars (Gavà, Barcelona, Cataluña)**

Terrasses de la Riera dels Canyars (henceforth, Canyars) is an open-air site located near Gavà (Barcelona) (41.2961; 1.9797), at 28 m.s.a.l and 3 km straight from the present-day coastline. The site lies on a fluvial terrace at the confluence of Riera dels Canyars, a torrential stream between Garraf Massif, Llobregat delta...
and Riera de Can Llong (Daura et al., 2013). Archaeo-paleontological remains were discovered during quarry activities in 2005 and were complete excavated on 2007 by the Grupo de Recerca del Quaternari (Daura and Sanz, 2006; Daura et al., 2013). This intervention determined nine lithological units. The paleontological and archaeological remains come exclusively from one unit, the middle luthitic unit (MLU), and specifically from layer I. The MLU is composed of coarse sandy clays and gravels, filling a paleochannel network named lower detrital unit (LDU) (Daura et al., 2013). Five radiocarbon dates were obtained on charcoals from layer I, which yield statistically consistent ages from 33,800 ±350 uncal BP to 34,900 ±340 uncal BP, which results in mean age of 39,600 cal BP (from 37,405 to 40,916 cal BP) (Daura et al., 2013).

The layer I of the site has yielded a rich faunal assemblage, consisting of over 5,000 remains. Among the herbivores, the most common species found are Equus ferus, Bos primigenius, Equus hydruntinus, and Cervus elaphus (Daura et al., 2013; Sanz-Royo et al., 2020). Capra sp. and Sus scrofa are also present, although in lower frequencies. The carnivores found at the site are also noteworthy, with Crocuta crocuta and Lynx partitus being the most frequent. Presence of cold-adapted fauna associated to stepped environments is recorded, such as cf. Mammutthus sp., Coelodonta antiquitatis, and Equus hydruntinus. Small mammal analysis, pollen, and use-wear analysis have provided further evidence that a steppe-dominated landscape surrounded the Canyars site, supporting a correlation with the Heinrich Event 4, in coherence with the chronology obtained for the layer (López-García et al. 2013; 2023; Rivals et al., 2017).

However, the presence of woodland is also attested by forest taxa within charcoal and pollen assemblages (Daura et al., 2013).

Taphonomic study is ongoing. But several evidences point that hyenas have played an important role in the accumulation of the faunal assemblage (Daura et al., 2013; Jimenez et al. 2019). However, sporadic human presence is documented by few human modifications found in faunal remains (cutmarks and fire alterations). Although the paucity of the lithic assemblage in the site, it shows a clear attribution to Upper Palaeolithic technocomplex, most likely the Early Aurignacian (Daura et al., 2013). Recently, it was documented a perforated bone fragment, which has been identified as a perforated board for leather production (Doyon et al., 2023). All teeth included in this work were sampled in Laboratori de la Guixera (Ajuntament de Casteldefels) where the material is stored.

References


Appendix C. Intratooth curve plots

Original curves derived from enamel intratooth sampling on enamel carbonate. Provided by sites. In blue, oxygen stable isotope composition ($\delta^{18}O$), and, in brown, carbon stable isotope composition ($\delta^{13}C$). In the x-axis, the distance from Enamel Root Junction (ERJ). Notice that the y-axis can experience some variations between sites.

**Figure C1.** Intratooth plots of oxygen ($\delta^{18}O$) and carbon ($\delta^{13}C$) isotope composition from teeth from Axlor, considering distance from enamel root junction (ERC).
Figure C2. Intratooth plots of oxygen (δ¹⁸O) and carbon (δ¹³C) isotope composition from teeth from El Castillo, considering the sample's distance from the enamel root junction (ERC).
Figure C3. Intratooth plots of oxygen ($\delta^{18}$O) and carbon ($\delta^{13}$C) isotope composition from teeth from El Castillo, considering the sample's distance from the enamel root junction (ERC).
Figure C4. Intratooth plots of oxygen ($\delta^{18}$O) and carbon ($\delta^{13}$C) isotope composition from teeth from El Castillo, considering the sample’s distance from the enamel root junction (ERC).

Figure C5. Intratooth plots of oxygen ($\delta^{18}$O) and carbon ($\delta^{13}$C) isotope composition from teeth from Labeko Koba, considering the sample’s distance from the enamel root junction (ERC).
Figure C6. Intratooth plots of oxygen ($\delta^{18}O$) and carbon ($\delta^{13}C$) isotope composition from teeth from Labeko Koba, considering the sample's distance from the enamel root junction (ERC).

Figure C7. Intratooth plots of oxygen ($\delta^{18}O$) and carbon ($\delta^{13}C$) isotope composition from teeth from Aitzbitarte III, considering the sample's distance from the enamel root junction (ERC).
Figure C8. Intratooth plots of oxygen ($\delta^{18}O$) and carbon ($\delta^{13}C$) isotope composition from teeth from El Otero, considering the sample's distance from the enamel root junction (ERC).
Figure C9. Intratooth plots of oxygen ($\delta^{18}O$) and carbon ($\delta^{13}C$) isotope composition from teeth from Canyars considering the sample's distance from the enamel root junction (ERC).
Appendix D. Inverse Modelling: Methodological Details and Models

The intratooth δ¹⁸O profiles presented in this study were obtained through the application of inverse modelling, using an adapted version of the code published in reference (Passey et al., 2005). This modeling approach allowed for the correction of the damping effect and the reconstruction of the original δ¹⁸O input time series. The model utilizes different species-specific parameters related to enamel formation, which vary between bovines and equids. These parameters have been established based on previous studies (Bendrey et al., 2015; Blumenthal et al., 2014; Kohn, 2004; Passey and Cerling, 2002; Zazzo et al., 2012). For *Bos/Bison* sp., the initial mineral content of enamel is fixed at 25%, the enamel appositional length is set at 1.5 mm, and the maturation length is 25 mm. For *Equus* sp., the initial mineral content of enamel is fixed at 22%, the enamel appositional length is set at 6 mm, and the maturation length is 28 mm.

In addition, the model requires other variables related to sampling geometry, as well as error estimates derived from mass spectrometer measurements. The distance between samples varies for each tooth, but as a general trend, the sampling depth on the tooth enamel surface in the samples of this study represents approximately 70% of the total enamel depth. The standard deviation of the measurements obtained from the mass spectrometer was typically set at 0.12%, taking into account the uncertainty associated with the standards. Finally, the models require a damping factor that determines the cumulative damping along the isotopic profile by adjusting the measured error (Emeas) to the prediction error (Epred). In the teeth analysed in this study, the damping factor ranged from 0.001 to 0.1. The most likely model solutions were selected, and summer and winter values were extracted from the δ¹⁸O profiles, considering the original peaks and troughs identified in the unmodelled δ¹⁸O profile. This approach was adopted to prevent the introduction of artificial peaks that the model may produce, particularly in teeth without a distinct sinusoidal shape.
Figure D1. Inverse models for oxygen isotope composition ($\delta^{18}O$) from teeth from Axlor, considering distance from enamel root junction. The blue line and points correspond to original data and grey line the most likely model solution, with the 95% confidence interval shown in shaded areas.
Figure D2. Inverse models for oxygen isotope composition ($\delta^{18}O$) from teeth from El Castillo, considering distance from enamel root junction. The blue line and points correspond to original data and grey line the most likely model solution, with the 95% confidence interval shown in shaded areas.
Figure D3. Inverse models for oxygen isotope composition (δ¹⁸O) from teeth from El Castillo, considering distance from enamel root junction. The blue line and points correspond to original data and grey line the most likely model solution, with the 95% confidence interval shown in shaded areas.
Figure D4. Inverse models for oxygen isotope composition (δ¹⁸O) from teeth from Labeko Koba, considering distance from enamel root junction. The blue line and points correspond to original data and grey line the most likely model solution, with the 95% confidence interval shown in shaded areas.
Figure D5. Inverse models for oxygen isotope composition ($\delta^{18}$O) from teeth from Canyars considering distance from enamel root junction. The blue line and points correspond to original data and grey line the most likely model solution, with the 95% confidence interval shown in shaded areas.

Figure D6. Inverse models for oxygen isotope composition ($\delta^{18}$O) from teeth from El Otero, considering distance from enamel root junction. The blue line and points correspond to original data and grey line the most likely model solution, with the 95% confidence interval shown in shaded areas.
Figure D7. Inverse models for oxygen isotope composition (δ18O) from teeth from Aitzbitarte III, considering distance from enamel root junction. The blue line and points correspond to original data and grey line the most likely model solution, with the 95% confidence interval shown in shaded areas.

References Appendix D


