



Pliocene–Early Pleistocene climatic trends in the Italian Peninsula based on stable oxygen and carbon isotope compositions of rhinoceros and gomphothere tooth enamel



Péter Szabó ^{a, b, c, *}, László Kocsis ^d, Torsten Vennemann ^c, Luca Pandolfi ^e, János Kovács ^{a, f}, Edoardo Martinetto ^g, Attila Demény ^h

^a Environmental Analytical and Geoanalytical Research Group, Szentágothai Research Centre, University of Pécs, Pécs, Hungary

^b Doctoral School of Chemistry, University of Pécs, Pécs, Hungary

^c Institut des Dynamiques de la Surface Terrestre, Université de Lausanne, 1015 Lausanne, Switzerland

^d Geology Group, Faculty of Science, Universiti Brunei Darussalam - UBD, Brunei Darussalam

^e Department of Sciences, Section of Geology, University of Roma TRE, Roma, Italy

^f Department of Geology and Meteorology, University of Pécs, Pécs, Hungary

^g Department of Earth Sciences, University of Torino, Torino, Italy

^h Institute of Geological and Geochemical Research, Research Centre for Astronomy and Earth Sciences, Budapest, Hungary

ARTICLE INFO

Article history:

Received 20 July 2016

Accepted 6 November 2016

Keywords:

Stable isotopes

Enamel

Climate

Pliocene

Early Pleistocene

Italy

ABSTRACT

The Pliocene and Early Pleistocene (5.2–1 Ma) palaeoclimate for localities in Italy is evaluated using stable carbon and oxygen isotope compositions of tooth enamel of fossil specimens from Rhinocerotidae (*Stephanorhinus* sp.) and Gomphotheriidae (*Anancus* sp.) taxa. Carbon isotope composition was measured in the structural carbonate ($\delta^{13}\text{C}$), while oxygen isotope values were determined both in the structural carbonate ($\delta^{18}\text{O}_{\text{CO}_3}$) and the phosphate ($\delta^{18}\text{O}_{\text{PO}_4}$) of bioapatite.

The $\delta^{13}\text{C}_{\text{CO}_3}$ values indicate that the taxa were grazers-browsers of a pure C₃ vegetation. Low $\delta^{13}\text{C}_{\text{CO}_3}$ values for Central and North Italy indicate a humid climate with woodlands and forest cover in the Pliocene. For northern localities the $\delta^{13}\text{C}$ values increase between MN16a and MNQ16b biozones most likely linked to the Northern Hemisphere Glaciation at 2.7 Ma after the “Mid-Pliocene Warm Period”. For Central Italy the values have a wide range with a long term increasing trend in the Early Pleistocene, indicating more arid climate and/or more open vegetation.

Overall, the $\delta^{18}\text{O}_{\text{PO}_4}$ values in Central Italy change together with the $\delta^{13}\text{C}_{\text{CO}_3}$ values and are taken to reflect the warmer/wetter interglacials and cooler/more arid glacial phases. The $\delta^{18}\text{O}_{\text{PO}_4}$ values in North Italy are lower than those in Central Italy and show no clear temporal trend. One explanation for the low values especially in MN14–15 biozone is that these $\delta^{18}\text{O}_{\text{PO}_4}$ values do not reflect entirely the isotopic composition of local precipitation but river waters from the Alps with ¹⁸O-depleted isotopic compositions or a N-S directed rain-shadow effect on the precipitation. In general the new isotope data agree well with palaeoclimate reconstructions based on palynological and other proxies.

© 2016 Elsevier Ltd. All rights reserved.

1. Introduction

There is an increasing interest in understanding Pliocene and Early Pleistocene terrestrial climates and environments because it can help to predict future climate changes more accurately. Reconstructions of terrestrial palaeoenvironments, palaeoecology,

and palaeoclimates are commonly based on proxies like paleosols, fossil plants (including pollen), vertebrate occurrences, speleothems, travertines, tufas, as well as the geochemistry and isotope geochemistry of biogenic carbonates and phosphates such as mammal teeth and bones. (Sullivan and Krueger, 1981; Luz et al., 1984, 1990; Luz and Kolodny, 1985; Thorp and Van Der Merwe, 1987; Leethorp et al., 1989; Ayliffe et al., 1992, 1994; Bryant and Froelich, 1995; Bocherens et al., 1996; Fricke et al., 1998; Cerling and Harris, 1999; Kohn et al., 1999; Kohn and Cerling, 2002; Arpke and Karhu, 2006; Levin et al., 2006; Tütken et al., 2006;

* Corresponding author. Doctoral School of Chemistry, University of Pécs, Pécs, Hungary.

E-mail address: sz.piiit01@gmail.com (P. Szabó).

Martin et al., 2008; Kohn, 2010; Pellegrini et al., 2011; Kocsis et al., 2014; Pushkina et al., 2014; García-Alix, 2015; Kovács et al., 2015; Hartman et al., 2016; Metcalfe et al., 2016 and references therein).

In this study 51 Pliocene and Early Pleistocene tooth enamel samples of fossil rhinoceros ($n = 44$) and gomphothere ($n = 7$) were studied from Italy. Most of the samples are from Central and North Italy and a few samples from South Italy (Fig. 1). The age of the fossils covers the Early Pliocene to late Early Pleistocene, from about 5.2 to 1 Ma.

The carbon and oxygen isotope compositions of structural carbonate and the oxygen isotope composition of phosphate in enamel bioapatite were used to explore variations in past climate and the environment of the animals. The aims of this study are: 1) Reconstructing the isotope composition of environmental water ($\delta^{18}\text{O}_{\text{W}}$) and estimating the changes in mean annual surface air temperature (MAT) on the basis of $\delta^{18}\text{O}_{\text{PO}_4}$ values of fossil tooth enamel; 2) interpretation of the diet of the species; 3) detect changes in vegetation based on the enamel carbon isotope analyses.

These climatic and ecological parameters derived from the isotope results are compared with palaeobotanical proxies, palaeontological and palaeoecological information (Fauquette et al., 1999; Pontini and Bertini, 2000; Bertini, 2001, 2010; Bredenkamp et al., 2002; Ji et al., 2002; Fauquette and Bertini, 2003; Klotz et al., 2006; Palombo, 2007; Magri et al., 2010; Rook and Martínez-Navarro, 2010; Petronio et al., 2011; Combourieu-Nebout et al., 2015; Loftus et al., 2015; Rivals et al., 2015).

2. Background

2.1. Bioapatite and preservation of its isotopic compositions

Biogenic hydroxyapatite (i.e., bioapatite) is the main inorganic fraction of the skeletal tissues of mammals with up to 6 wt% of carbonate as structurally bound carbonate. The simplified formula

is $\text{Ca}_5(\text{PO}_4, \text{CO}_3)_3(\text{OH}, \text{CO}_3)$ (e.g., Kohn et al., 1999). There are two types of carbonate forms in bioapatite, structural and labile carbonate. Structural carbonate substitutes for PO_4^{3-} and OH^- , while the labile CO_3^{2-} component's structural identity is ambiguous as it is often considered adsorbed onto the surface. Oxygen is present in three different ions in bioapatite: phosphate, hydroxyl and carbonate ions. Because of the strong P–O bonds phosphate oxygen is considered more resistant to low temperature inorganic alteration processes, than carbonate oxygen (Kohn et al., 1999; Kohn and Cerling, 2002). However, during microbiological reactions even the phosphate oxygen isotope composition can be changed due to enzymatic catalysis between PO_4^{3-} and water (e.g., Blake et al., 1997; Zazzo et al., 2004a,b; Liang and Blake, 2007). Nevertheless, it was shown experimentally that when enamel (i.e., well-crystallized bioapatite) is subjected to bacterially mediated conditions, the oxygen isotopic composition of the phosphate group was not affected (e.g., Zazzo et al., 2004a,b). Moreover, enamel is the preferred tissue for isotopic investigations because it contains the highest proportion of apatite (96%), and structurally compact with little pore space and large phosphate crystallites (up to 1 μm long), arranged in a decussate texture (Kohn et al., 1999). As a result enamel is more resistant to diagenesis than other tissues in the same taphonomic context (Thorp and Van Der Merwe, 1987; Quade et al., 1992; Ayliffe et al., 1994; Wang and Cerling, 1994; Koch et al., 1997).

There is a widely used method to monitor diagenetic effects. Oxygen from the body fluids is in isotopic equilibrium with both the carbonate and phosphate ions of the inorganic tissue. Several studies have found a constant offset between the $\delta^{18}\text{O}_{\text{CO}_3}$ and $\delta^{18}\text{O}_{\text{PO}_4}$ values for non-altered mammal tooth enamel and the $\delta^{18}\text{O}_{\text{CO}_3} - \delta^{18}\text{O}_{\text{PO}_4}$ slopes are similar for different species (Bryant et al., 1996; Iacumin et al., 1996; Arppe and Karhu, 2006, 2010; Tütken et al., 2007; Pellegrini et al., 2011; Domingo et al., 2013). Because it seems quite improbable to find isotopically altered



Fig. 1. Location of paleontological sites cited in Table A1 (the map shows the present day locations.).

skeletal remains showing a good correlation and a similar offset to modern samples, it is plausible to use the parallel $\delta^{18}\text{O}_{\text{CO}_3} - \delta^{18}\text{O}_{\text{PO}_4}$ measurements as a monitor of diagenetic alteration (Iacumin et al., 1996). Most studies found an offset 8.4–9.1‰ for modern, unaltered samples (Bryant et al., 1996; Iacumin et al., 1996; Martin et al., 2008; Pellegrini et al., 2011). Nevertheless, Martin et al. (2008) found ~2‰ intra tooth differences in two upper molars (M1 and M2) of extant *Rhinoceros unicornis* and further variations can be present due to environmental, physiological and tooth growth process differences. The final recommendation is that if the $\delta^{18}\text{O}_{\text{CO}_3} - \delta^{18}\text{O}_{\text{PO}_4}$ difference is in the range of 7.2‰–10.6‰ the values of the samples can be considered as unaltered (Martin et al., 2008).

2.2. Stable carbon isotope composition and diet

Carbon isotope differences among vertebrates largely reflect the differences in the diet of the animals. Variations in $\delta^{13}\text{C}$ values of the plants reflect mostly the three different photosynthetic pathways used by plants. The C₃ pathway is the most common, occurring in all trees, most shrubs and herbs, and grasses in regions with a cool-temperate growing season. C₃ plants have a mean $\delta^{13}\text{C}$ value of about –27‰ with a range from –35 to –22‰. C₄ photosynthesis occurs in grasses from regions with a warm growing season, and in some sedges (Bruhl and Wilson, 2007) and dicots. C₄ plants have higher $\delta^{13}\text{C}$ values (mean ca. –13‰, range between –19 and –9‰). Crassulacean acid metabolism (CAM) is the least common pathway, occurring mostly in succulent plants, with a range between the C₃ and C₄ plants (O'Leary, 1988; Farquhar et al., 1989; Martinelli et al., 1991). Enamel $\delta^{13}\text{C}$ values of large non-ruminant herbivores have a bioapatite-plant fractionation offset of about 13.3‰ (Passey et al., 2005). In C₃ ecosystems, the variation in $\delta^{13}\text{C}$ values can provide further information about mean annual precipitation (MAP), humidity, the type of vegetation, habitat differences or niche partitioning between animals (Van der Merwe and Medina, 1991; Feranec and MacFadden, 2006; Kohn, 2010). Generally C₃ plants show a monotonic increase in $\delta^{13}\text{C}$ with decreasing MAP (Kohn, 2010). Effects of vegetation like the canopy effect complicate this relationship though. In forests the limited sunlight availability, high relative humidity and the low $\delta^{13}\text{C}_{\text{CO}_2}$ values from decaying litter of the soil organic matter lowers the $\delta^{13}\text{C}$ values (Van der Merwe and Medina, 1991; Bocherens et al., 1996; Cerling and Harris, 1999; Drucker et al., 2008). As the $\delta^{13}\text{C}$ values of herbivores reflect the values of the consumed plants, preferential consumption of some specific plants could imply that the $\delta^{13}\text{C}$ values ofapatite can not be taken to determine the average vegetation. Food preference of the animals is also not necessarily constant within a taxonomic group, and for a given taxon through space and time.

Because of the complex information encoded in the $\delta^{13}\text{C}$ values, several studies applied it to address a variety of palaeodietary, palaeoecological, and palaeoenvironmental problems (e.g. Leethorp et al., 1989; Quade et al., 1992; Bocherens et al., 1996; Cerling and Harris, 1999; Iacumin et al., 2000; Feranec, 2004; Metcalfe et al., 2009; Arpke et al., 2011; Montanari et al., 2013; Kocsis et al., 2014; Scherler et al., 2014; Zanazzi et al., 2015).

2.3. Stable oxygen isotope composition and environmental water

Oxygen isotope compositions of mammal teeth are linearly related to the body water of the animals (Longinelli, 1984). The isotope composition of body water depends on the total oxygen flux through the body, but water from drinking and food sources have the largest effect compared to other factors (Luz et al., 1984). In the case of elephants the ratio is approximately 2:1 of water ingested via drinking and that ingested by consumption of plant matter (Ayliffe et al., 1992), while for rhinoceroses this ratio is 4:1

(Clauss et al., 2005; Martin et al., 2008). Thus teeth of these two groups are potentially good indicators of the $\delta^{18}\text{O}$ values of local environmental waters (Martin et al., 2008). The drinking water of the animals may include different sources including streams, ponds, rivers, lakes, and leaves. Each of these reservoirs typically can have different $\delta^{18}\text{O}$ values relative to precipitation, due to variable mixing of temporally different precipitation and potential evaporative effects on the different reservoirs (e.g., Montanari et al., 2013). Teeth of large (>100 kg) obligate drinking mammals record an average oxygen isotope composition of ingested waters ($\delta^{18}\text{O}_{\text{w}}$) from different sources from a longer period. Thus the correlation between the average $\delta^{18}\text{O}$ values of teeth enamel and the local precipitation is usually strong (Ayliffe et al., 1992; Bryant and Froelich, 1995; Koch, 1998). However, there are exceptions for example glacial meltwaters, rivers from mountainous area, or lakes with high evaporation in arid areas can have a systematic $\delta^{18}\text{O}$ offset from the local precipitation. Migration of the animals can also complicate the relationship. If animals migrated in order to escape harsh winters or to exploit seasonally available resources, isotope composition of their enamel may not be representative for the area where the fossils are found (Hoppe and Koch, 2007). The $\delta^{18}\text{O}$ values can also offer information about the ecology of the different taxa if they lived under the same climatic regime. Browsing taxa that ingest a higher proportion of ¹⁸O-enriched water with their food tend to have higher $\delta^{18}\text{O}$ values compared to sympatric grazing taxa (Kohn, 1996; Tütken et al., 2013).

The oxygen isotope composition of local precipitation ($\delta^{18}\text{O}_{\text{ppt}}$) is controlled by several factors. The mean annual temperature (MAT), the different moisture source, and air mass trajectories, the latitude, altitude, continental and amount effects, all determine the isotopic compositions of precipitation. The MAT generally has the greatest effect on the $\delta^{18}\text{O}$ values of the precipitation ($\delta^{18}\text{O}_{\text{ppt}}$) at mid and high latitudes and in case of no extreme low or high precipitation or humidity (Dansgaard, 1964; Rozanski et al., 1993).

2.4. Pliocene-Pleistocene palaeoclimates and palaeoenvironments in Italy

According to different proxies and models, the climate in the Early Pliocene was in general warmer and wetter than today (2–3 °C higher global MAT with a decreased equator to pole gradient) (Dowsett et al., 1996, 2013; Hill et al., 2011; Salzmann et al., 2011; Haywood et al., 2013). In continental Europe thick forests were present with a temperate climate. Palynological data suggest a relatively stable, warm-temperate climate in Northern Italy with 15–17 °C MAT and a MAP above 1000 mm (Bertini, 2001, 2010). Nevertheless, the slight differences between megathermic elements and meso-microthermic ones suggest modest fluctuations of temperature (Bertini, 2001). The Climatic Amplitude Method (CAM) applied to other pollen records from Northern Italian sites gave MAT between 16 and 20 °C, with the Most Likely Value (MLV) around 17–19 °C; and MAP between 1100 and 1500–1600 mm, with the MLV around 1200–1300 mm (Fauquette et al., 1999; Fauquette and Bertini, 2003). In Sicily the climate was warmer and drier (or with equal humidity) than today (Fauquette et al., 1999, 2006; Combourieu-Nebout et al., 2015).

During the MN16a Neogene Mammalian biozone the climate in general became less stable compared to the Early Pliocene (Hilgen et al., 2012). In SW Europe the modern Mediterranean climatic regime was established and aridity was enhanced (Agustí et al., 2001; Meyers and Hinnov, 2010; Rook and Martínez-Navarro, 2010; De Schepper et al., 2013, 2014; Woodard et al., 2014). The Villanyian/Villafranchian mammal turnover occurred at this time in Europe and a renewal of archaic faunas to mammal assemblages with more modern characteristics indicative to less humid

environments under cooler climates documented also in Italy (Bredenkamp et al., 2002; Petronio et al., 2011). From 3.3 to 3.0 Ma the “Mid-Pliocene Warm Period” (MPWP) was characterized by 3 °C warmer global temperatures, 10–40 m higher sea-level and less continental ice sheets compared to present day, but even this stable, warm period was preceded by a short-lived, intense global glaciation (3.305–3.285 Ma) (De Schepper et al., 2014). Around 3.3 Ma (MN16a, Traversa FU) in Central Italy the faunas are characterized by the occurrence of new species and by the diffusion of animals also adapted to open habitats. The palaeobotanical data indicates a considerable homogeneity of the vegetation, at least in Northern Italy. Hence, besides the new species, animals strictly relying on the forests still survived. For example *Anancus arvernensis* as a forest living taxon survived from the Ruscianian.

At approximately 2.7 Ma the Northern Hemisphere Glaciation (NHG) occurred, followed by glacial-interglacial cycles of moderate amplitude (orbital periodicity of 41 ka) (Dowsett et al., 2013; Woodard et al., 2014; Salzmann et al., 2011; Haywood et al., 2013). The resulting increase in aridity and more intense seasonality caused a replacement of forests by tundra-like vegetation in Northern Europe (Bredenkamp et al., 2002) and a faunal renewal in Italy, where several forest-dwelling taxa, especially small carnivores and arboreal-scansorial taxa disappeared, whereas new large grazers, mixed feeders or even browsers appeared. This renewal, called the “*Equus*-elephant event” can be regarded as a true turnover phase (Palombo, 2007; Petronio et al., 2011). The irreversible transformation of the climate regime determines rapid variations of the vegetation community, according to a repetitive succession of vegetation types. This event also corresponds to a massive local disappearance of thermophile and humidity-requiring woody plant taxa (Martinetto et al., 2015), even if a few of them survived in Central Italy longer than in the rest of Europe (until the end of the Gelasian, ca. 1.8 Ma), due to a combination of locally high precipitation and high temperatures.

The entire interval between 2.5 and 1.0 Ma is characterized by relatively long interglacial periods, with rather thick forest cover, and short glacial periods, with open vegetation in Central and Southern Italy (Petronio et al., 2011 and references therein; Combouieu-Nebout et al., 2015). In the Mediterranean area the typical glacial-interglacial cycles are characterized by the succession of four main vegetation assemblages. Deciduous forest, subtropical/warm-temperate forest, altitudinal conifer forest and open vegetation. This succession supports four dominant climatic conditions: (1) a first increase of the temperature followed by (2) an increase of the humidity, then (3) a decrease of temperature without variations of humidity and (4) finally a strong decrease of the humidity, corresponding to a gradual transition from warm and humid conditions during interglacials to cold and dry conditions during glacials (Bertini, 2001, 2010). Pollen data indicate that the steppe-type of conditions, which are longer lasting and more frequent in Southern Italy, developed only during some extreme glacial peaks in Central Italy (Russo Ermolli et al., 2010; Pontini and Bertini, 2000; Petronio et al., 2011). In North Italy the glacial periods led to a progressive expansion of a cooler forest-type of vegetation cover (i.e., progressive increase of *Picea* and *Fagus*), but herbs including steppe taxa remained a minor component (Bertini, 2001, 2010).

Despite the many variations it can be concluded that interglacial phases can be characterized with high MAT and MAP, similar to that in the Zanclean, while during the glacial phases the MAT and MAP were close to modern values. This cyclicity was controlled by the 41 ka rhythm of the orbital periodicity continued in the entire period until 0.9 Ma (Bertini, 2000).

The Pliocene and Early Pleistocene sediments of Northern and Central Italy (often from the same successions that yielded the

studied fossil vertebrates) are rich in macrofossil plant assemblages (Forno et al., 2015; Irace et al., 2015; Martinetto, 2015; Martinetto et al., 2015) that have not yet been exploited for a thorough palaeoclimatic analysis, despite their potentially very accurate indications (Teodoridis et al., 2015). The aims of this study are to detect the temporal changes in the vegetation and in MAT and to compare the new isotope results with the palaeobotanical proxies and palaeontological information. The comparisons of the results obtained from the different approaches are summarized in Table 1.

3. Materials and methods

3.1. Fossil materials

Fig. 1 and Table A1 give detailed information about the localities, their relative age relationships and the types of investigated fossils. Altogether 51 fossils from 21 localities were collected from the time period of about 5.2 to 1 Ma. For some localities only one sample whereas for others up to 7 samples were collected. The fossils belong to four different species, three of them are rhinocerotid: *Stephanorhinus etruscus* (n = 21), “*Dihoplus*” *megarhinus* (n = 9), *Stephanorhinus jeanvireti* (n = 6), and one proboscidean: *Anancus arvernensis* (n = 7) (Fig. 1, Table A1). These include rhinoceros samples that cannot be determined at specific level (n = 8). Wherever possible, teeth of adult individuals were sampled. Each fossil probably represents a different individual. Samples were acquired from the following institutions: DST, Department of Earth Science, Sapienza University of Roma (Roma); MSNAF, Natural History Museum, Fisiocritici Academy, Siena (Siena); MGC, Museum of Geology G. Capellini, (Bologna), MGPT, Museum of Geology and Paleontology, University of Turin (Turin), MHMB, Natural History Museum Basel (Basel).

Many of the investigated sites and fossils are chronologically well calibrated using magnetostratigraphy, cyclostratigraphy and biochronology. For the interpretation of the results the samples were grouped in the Neogene Mammal biozones. The numbers of samples from different time periods are different in Central and in North Italy. The oldest samples are from MN14 in Central and from MN14–15 in North Italy. The precise stratigraphic and geographic position of the “*Dihoplus*” *megarhinus* specimens from Dusino San Paolo is uncertain; the MN14–15 interval is based on the well-known record of the taxa.

In Central Italy MNQ18, MNQ19 and MNQ19–20 biozones are treated separately, while in the case of North Italy due to the fewer and less well-dated samples from these time periods, the MN18–19–20 biozones are drawn together (n = 3). In North Italy the MN16 biozone can be separated to MN16a (Traversa FU, 5 samples) and MNQ16b (Montopoli FU, 4 samples) biozones. However recent analyses on plant macrofossils assemblages from the same area where MNQ16b mammal assemblages have been reported, in NW Italy (Irace et al., 2015), noted the absence of several thermophilous plants (HUTEA of Martinetto et al., 2015) already within the latest Pliocene. This suggests a change of vegetation already in the late Pliocene, and not only at the beginning of the Pleistocene. Since vegetation changes often coincide with mammal turnover events, the actual assignment to the MN16a/MNQ16b mammal zones should be further investigated. In fact the MNQ16b mammals are documented by 19th century findings of uncertain stratigraphic position that could fall within the latest Pliocene as well as within the earliest Pleistocene.

3.2. Sampling

After cleaning the surface mechanically, about 10 mg of bulk enamel was sampled from each fossil using a Dremel diamond-

Table 1

Calculated MAT changes based on $\delta^{18}\text{O}_{\text{PO}_4}$ and vegetation estimates based on the $\delta^{13}\text{C}_{\text{CO}_3}$ isotope results compared with palaeobotanical proxies, (1) Bertini, 2001; (2) Bertini, 2010; (3) Fauquette et al., 1999; (4) Combourieu-Nebout et al., 2015; (5) Fauquette and Bertini, 2003; (6) Magri et al., 2010; (7) Klotz et al., 2006; (8) Fauquette et al., 2006; MLV: most likely value, NHG: Northern Hemisphere Glaciation.

	Age (Ma)	~5.2–3.5	~3.3–2.8	~2.8–2.5	~2.5–1.9	~1.9–1.7	~1.7–1.3	~1.3–1	Recent
	MN zone	MN14–15	MN16a	MN16b	MN17	MNQ18	MNQ19	MNQ19–20	
Central Italy	MAT changes ($^{\circ}\text{C}$) indications (this study)		2–3.6 $^{\circ}\text{C}$ decrease of MAT from MN14			2–3.6 $^{\circ}\text{C}$ decrease of MAT from MNQ18 to MNQ19–20			
	estimated vegetation (this study)	woodland-mesic grassland	woodland-mesic grassland			woodland-mesic grassland, more open vegetation/less humid/less precipitation than in the former biozones	woodland-mesic grassland, more open vegetation/less humid/less precipitation than in the former biozones		woodland-mesic grassland, more open vegetation/less humid/less precipitation than in the former biozones
North Italy	estimated vegetation (this study)	woodland-mesic grassland	woodland-mesic grassland	woodland-mesic grassland, more open vegetation/less humid/less precipitation than in MN6a		Interglacials: higher MAT and MAP than present day. Glacials: similar MAT and MAP to present day (2), vegetation openings at 2.5Ma and from 1.9 to 1.7Ma (MNQ18) (1)	Interglacials: higher MAT and MAP than present day. Glacials: similar MAT and MAP to present day (2), vegetation openings at 2.5Ma and from 1.9 to 1.7Ma (MNQ18) (1)		MAT 14–16 $^{\circ}\text{C}$, MAP 760–970 mm (4), 730 mm (6)
	palaeobotanical indications					woodland-mesic grassland, (one sample from Leffe indicate open woodland-xeric grassland)			
South Italy	estimated vegetation (this study)					Interglacials: similar MAP and MAT to Early Pliocene, Glacials: similar MAT to present day, and 500–600 mm higher MAP than present, (5) (Stirone site), alternations of different types of forests with no open vegetation (2)	Interglacials: similar MAP and MAT to Early Pliocene, Glacials: similar MAT to present day, and 500–600 mm higher MAP than present, (5) (Stirone site), alternations of different types of forests with no open vegetation (2)	MAT, 13–15 $^{\circ}\text{C}$, MAP 930–1100 mm (4)	
	palaeobotanical indications	MAT 16–20 $^{\circ}\text{C}$ (MLV 17–19 $^{\circ}\text{C}$) (1; 2), 12–20 $^{\circ}\text{C}$ (3), 1–4 $^{\circ}\text{C}$ higher than present day (8), MAP 1100–1600 mm (MLV 1200–1300 mm) (1,2), 1100–1400 mm (3), 400–700 mm higher than present day (8)							
						woodland-mesic grassland	open vegetation and forest alternations	MAT, 15–18 $^{\circ}\text{C}$, MAP 440–830 mm (4)	
							(2)		

studded drill. Where it was possible, enamel was sampled along a vertical line over the whole enamel length from the crown to the root to get a representative mean sample of the period of enamel formation. Due to sample limitations, in some cases enamel from tooth fragments was collected. Because the complex process of enamel formation and mineralization, which can be 2–3 years in the case of rhinos and proboscideans, sampling enamel from a tooth fragment can still represent an average isotope record of a longer period (Tafforeau et al., 2007; Metcalfe and Longstaffe, 2012). Where possible, the teeth from adult individuals were sampled, because physiological effects such as weaning/nursing can modify the $\delta^{13}\text{C}$ and $\delta^{18}\text{O}$ values of the bioapatite (Martin et al., 2008; Metcalfe et al., 2010). Where possible, several samples were collected from each site to avoid the effects of environmental factors and individual differences. According to Pryor et al. (2014) typical uncertainty in temperature inferred from a single sample can be at least $\pm 4^\circ\text{C}$, while analyses of multiple samples can reduce the uncertainty to about 1–2 $^\circ\text{C}$.

3.3. Pre-treatment and measurements

Sample powder was pre-cleaned according to the method given in Koch et al. (1997). NaOCl was used to remove soluble organic material and acetic acid-Ca-acetate buffer to remove exogenous carbonates (Koch et al., 1997; Kocsis, 2011). The pre-treatment methods have been recently reviewed by Pellegrini and Snoeck (2016). The pre-treatment procedure suggested is supported by the authors, but it is also clear that some unpredictable contamination cannot be excluded. This may influence the finer crystalline material (e.g., bone, dentine) and could also be the cause for the slightly lower NBS-120c oxygen isotope compositions measured parallel with the teeth.

After pre-cleaning, about 2 mg of sample was used for the carbonate isotopic measurements, while for the $\delta^{18}\text{O}$ analyses, another 2 mg was dissolved and the phosphate ions were precipitated as silver phosphate, according to methods adapted after Dettman et al. (2001) and Kocsis (2011).

The carbon and oxygen isotopic compositions of structural carbonate were measured on a Finnigan MAT Delta Plus XL mass spectrometer equipped with a GASBENCH-II preparation unit. The samples were reacted with 99% orthophosphoric acid and the produced CO_2 was introduced to the mass spectrometer with He-carrier gas following procedures similar to those described in Spötl and Vennemann (2003).

Carrara Marble in-house standards ($\delta^{18}\text{O} = -1.70\text{\textperthousand}$, VPDB; $\delta^{13}\text{C} = 2.05\text{\textperthousand}$, VPDB) were run in the same sequence with the samples and used for correcting the data. The reproducibility of the in-house standard is better than $0.1\text{\textperthousand}$ (1σ) for both oxygen and carbon isotopic compositions. NBS-120c reference material (Florida phosphate rock) was treated and analysed in parallel with the samples and gave values of $\delta^{13}\text{C}$ of $-6.3 \pm 0.1\text{\textperthousand}$ ($n = 6$) and $\delta^{18}\text{O}$ of $-2.3 \pm 0.2\text{\textperthousand}$ ($n = 6$) on VPDB scale. These values are identical to those of the long-term average values of NBS-120c in the laboratory.

For the phosphate $\delta^{18}\text{O}$ values the silver-phosphate was analysed via reduction with graphite in a TC/EA (high-temperature conversion elemental analyser) coupled to a Finnigan MAT Delta Plus XL mass spectrometer according to the values and method given in Vennemann et al. (2002). The results were corrected to in-house Ag_3PO_4 phosphate standards (LK-2L: $12.1\text{\textperthousand}$ and LK-3L: $17.9\text{\textperthousand}$) that showed standard deviations (1σ) better than $\pm 0.3\text{\textperthousand}$ during the measurements. As internal standard the NBS-120c was prepared and run together with the samples in order to test sample preparation. The analyses gave an average $\delta^{18}\text{O}$ value of $21.2 \pm 0.3\text{\textperthousand}$, ($n = 8$), which is slightly lower than reported by others

and also lower compared to the NBS-120c that was measured for the calibration of TU-1 and TU-2 ($21.7 \pm 0.3\text{\textperthousand}$, e.g., Vennemann et al., 2002; Halas et al., 2011). Because there is no internationally accepted $\delta^{18}\text{O}$ value for this material and as it is a sedimentary rock, the mammal teeth data were not corrected further. The carbon isotope compositions are expressed relative to VPDB standard (Vienna Pee Dee Belemnite), while the oxygen either to VPDB or VSMOW (Vienna Standard Mean Ocean Water).

4. Results

The results of the carbon ($\delta^{13}\text{C}$) and oxygen ($\delta^{18}\text{O}_{\text{CO}_3}$ and $\delta^{18}\text{O}_{\text{PO}_4}$) isotope measurements along with the calculated $\delta^{13}\text{C}$ values of vegetation, $\delta^{18}\text{O}$ values of environmental water ($\delta^{18}\text{O}_w$) (see equations in Table 2) are presented in Table A1. The raw data, along with a step-by-step set of calculations will allow a possible future verification and recalculation of the results (recommendation of Skrzypek et al., 2016). Fig. 2 illustrates the $\delta^{13}\text{C}$ and $\delta^{18}\text{O}_{\text{PO}_4}$ values of the samples clustered in time bins with vegetation estimates and relative MAT scales.

Most of the collected samples for this study have a $\delta^{18}\text{O}_{\text{CO}_3} - \delta^{18}\text{O}_{\text{PO}_4}$ offset between $7.2\text{\textperthousand}$ and $10.6\text{\textperthousand}$ (see in Table A1), which is typical for samples that have not been subjected to diagenesis. Only two enamel samples have a lower carbonate-phosphate difference than $7.2\text{\textperthousand}$, that is 6.8 and $6.9\text{\textperthousand}$, which could indicate some degree of diagenetic alteration, but these samples have similar phosphate $\delta^{18}\text{O}$ and carbonate $\delta^{13}\text{C}$ values as other samples from the sites, so using these results does not change the site averages. It is possible that in these samples the $\delta^{18}\text{O}_{\text{CO}_3}$ values were slightly affected by late diagenetic processes (e.g., exchange with pore fluids).

To detect differences between species, MN biozones and areas, both parametric (F-tests for variances and two-tailed homo- or heteroscedastic t-tests for central values) and non-parametric (Levene tests for variances and two-tailed Mann-Whitney tests for central values) statistical tests were made. One way ANOVA and non-parametric Kruskal-Wallis tests were performed to compare variances of more than two groups. Statistical significance is based on $p < 0.05$.

4.1. Carbon isotope results

The $\delta^{13}\text{C}$ values for the whole fauna range from $-15.9\text{\textperthousand}$ to $-9.2\text{\textperthousand}$ (VPDB). The range of values is from -15.9 to $-10.8\text{\textperthousand}$ for Central Italy and from -15.1 to $-9.2\text{\textperthousand}$ for North Italy. The three samples from South Italy from MNQ19 and MNQ20 vary between -12.1 and $-11.5\text{\textperthousand}$ (VPDB).

Two tailed homoscedastic t-test ($p = 0.92$) and Mann-Whitney test ($p = 0.75$) indicate that there is no significant difference if all the values from Central and North Italy are compared.

The average values for the different time periods in Central and in North Italy are represented in Fig. 2. There are significant differences between some time periods. In the case of Central Italy t-tests indicate a change between MNQ18 and MNQ19 ($p = 0.044$) and between MNQ19 and MNQ19-20 ($p = 0.014$). However, according to the Mann-Whitney test these differences are not significant ($p = 0.125$ and $p = 0.092$). The average $\delta^{13}\text{C}$ values decrease by 1\textperthousand from MNQ18 to MNQ19 and increase by $2.3\text{\textperthousand}$ from MNQ19 to MNQ19-20.

In the case of North Italy both t-test ($p = 0.002$) and Mann-Whitney test ($p = 0.02$) support changes between MN16a and MNQ16b. The change in average values is $2.1\text{\textperthousand}$ in this transition.

The four species have average $\delta^{13}\text{C}$ values between $-14\text{\textperthousand}$ and $-13\text{\textperthousand}$ with no difference between them (ANOVA test, $p = 0.501$, Kruskal Wallis test, $p = 0.21$). The average values are $-13.0 \pm 1.1\text{\textperthousand}$ ($n = 21$) for *S. etruscus*; $-13.9 \pm 0.9\text{\textperthousand}$ ($n = 9$) for

Table 2The equations used for the calculations of water $\delta^{18}\text{O}$ values from those measured for the phosphates.

Number of equation	Relationship	Equation	type of regression	R^2 , slope	References
1	$\delta^{13}\text{C}_{\text{diet, meq}}$; $\delta^{13}\text{C}_{\text{leaf}}$	$\delta^{13}\text{C}_{\text{diet, meq}} = \delta^{13}\text{C}_{\text{leaf}} + (\delta^{13}\text{C}_{\text{modern atm CO}_2} - \delta^{13}\text{C}_{\text{ancient atm CO}_2})$			Domingo et al. (2013)
2	$\delta^{13}\text{C}_{\text{diet, meq}}$; MAP	$\delta^{13}\text{C}_{\text{diet, meq}} (\text{‰, VPDB}) = -10.29 + 1.9 \times 10^{-4} \text{ altitude (m)} - 5.61 \log_{10}(\text{MAP} + 300, \text{mm/yr}) - 0.0124 \text{ Abs (latitude, o)}$			Kohn (2010)
3	$\delta^{18}\text{O}_{\text{w}}$; $\delta^{18}\text{O}_{\text{PO}_4}$	$\delta^{18}\text{O}_{\text{w}} = (\delta^{18}\text{O}_{\text{PO}_4} - 23) / 0.90$	inverted forward regression	slope = 1.11	Kohn and Cerling (2002)
4	$\delta^{18}\text{O}_{\text{w}}$; $\delta^{18}\text{O}_{\text{PO}_4}$	$\delta^{18}\text{O}_{\text{w}} = (\delta^{18}\text{O}_{\text{PO}_4} - 26.8 + 8.9 \text{ h}) / 0.76$; (where h is the relative humidity)	inverted forward regression	slope = 1.32	Kohn (1996)
5	$\delta^{18}\text{O}_{\text{w}}$; $\delta^{18}\text{O}_{\text{PO}_4}$	$\delta^{18}\text{O}_{\text{w}} = 1.1128 (\pm 0.0029) \delta^{18}\text{O}_{\text{PO}_4} - 26.4414 (\pm 0.0508)$	transposed regression	$R^2 = 0.998$; slope = 1.11	Amiot et al. (2004)
6	$\delta^{18}\text{O}_{\text{w}}$; $\delta^{18}\text{O}_{\text{PO}_4}$	$\delta^{18}\text{O}_{\text{w}} = (\delta^{18}\text{O}_{\text{PO}_4} - 23.3 (\pm 0.7)) / 0.94 (\pm 0.10)$	inverted forward regression	$R^2 = 0.85$; slope = 1.06	Ayliffe et al. (1992)
7	MAT; $\delta^{18}\text{O}_{\text{w}}$; for Europe	$\text{MAT} = (\delta^{18}\text{O}_{\text{w}} + 13.74) / 0.53$	inverted forward regression	$R^2 = 0.6$, slope = 1.89	Pryor et al. (2014)
8	MAT; $\delta^{18}\text{O}_{\text{w}}$; for Europe	$\text{MAT} = 1.4118\text{O}_{\text{w}} + 21.64$	transposed regression	$R^2 = 0.49$, slope = 1.41	Skrzypek et al. (2016)
9	MAT; $\delta^{18}\text{O}_{\text{w}}$; for Italy (excluded the islands)	$\text{MAT} = 1.56\delta^{18}\text{O}_{\text{w}} + 23.31$	transposed regression	$R^2 = 0.3$, slope = 1.56	data of GNIP stations from Skrzypek et al. (2016)
10	MAT; $\delta^{18}\text{O}_{\text{w}}$; global equation	$\text{MAT} = (\delta^{18}\text{O}_{\text{w}} + 13.6) / 0.69$	inverted forward regression	slope = 1.45	Dansgaard (1964)
11	MAT; $\delta^{18}\text{O}_{\text{w}}$; global equation	$\text{MAT} = (\delta^{18}\text{O}_{\text{w}} + 14.18 \pm 0.52) / (0.49 \pm 0.03)$	inverted forward regression	$R^2 = 0.81$; slope = 2.04	Amiot et al. (2004)

"D." *megarhinus*; $-13.6 \pm 1.3\text{‰}$, ($n = 6$) for *S. jeanvireti*; and $-13.9 \pm 1.7\text{‰}$ ($n = 7$) for *A. arvernensis*.

4.2. $\delta^{18}\text{O}_{\text{PO}_4}$ values

The $\delta^{18}\text{O}_{\text{PO}_4}$ values for the whole fauna have a range from 12.0 to 18.7‰ (V-SMOW). Regarding the regions, the values vary from 14.1‰ to 18.4‰ in Central Italy and from 12.0‰ to 15.7‰ in North Italy. The three samples from South Italy from MNQ19 and MNQ20 have values between 16.6‰ and 18.7‰ . Comparing the average $\delta^{18}\text{O}_{\text{PO}_4}$ values from Central and North Italy the difference is significant (t -test, $p = 5.5 \times 10^{-6}$, Mann-Whitney test, $p = 2.1 \times 10^{-5}$). In Central Italy the averages are higher than values of North Italy in all of the time intervals. The differences are $1\text{--}3\text{‰}$ with the highest difference in the MN14–15 biozone. There are also differences in values between Central and South Italy in MNQ19–20. Samples from South Italy have about 3‰ higher average $\delta^{18}\text{O}_{\text{PO}_4}$ values than samples from Central Italy.

The average values for the different time periods in Central and in North Italy are represented in Fig. 2. In Central Italy the average $\delta^{18}\text{O}_{\text{PO}_4}$ values decrease significantly from MN14 to MN16 (t -test, $p = 0.004$, Mann-Whitney test, $p = 0.006$). In North Italy there are some changes in the average values between the different biozones but these changes are not significant.

The different species have the following average values: $15.5 \pm 1.5\text{‰}$ ($n = 21$) for *S. etruscus*; $15.9 \pm 1.9\text{‰}$ ($n = 9$) for "D." *megarhinus*; $15.7 \pm 0.7\text{‰}$ ($n = 7$) for *A. arvernensis* and $14.7 \pm 0.9\text{‰}$ ($n = 6$) for *S. jeanvireti*. The three species have similar average values, while *S. jeanvireti* has an average value about 1‰ lower. Differences among the species are not significant.

5. Discussion

5.1. Palaeoenvironment, and diet of the taxa based on $\delta^{13}\text{C}$ values

In this study all the $\delta^{13}\text{C}$ values indicate that these taxa were

browsers or grazers in a pure C_3 ecosystem. It is compatible with earlier suggestions that C_4 grasses were absent in Europe during the Pliocene to Late Pleistocene (Kürschner, 2010). Nonetheless, it must be mentioned that Iraze et al. (2015) reported the presence of C_4 sedges in NW Italy already in the latest Pliocene. The ecology of the nearest living relatives (e.g. *Cyperus glomeratus*) suggests that the distribution of these sedges was mainly limited to the disturbed environments along the rivers, and there is no evidence permitting to reconstruct their frequency in the vegetation and their possible importance as food for vertebrates (see Mishra et al., 2015).

Several calculations were made for the interpretation of the isotope results. The tested equations are summarized in Table 2. From the $\delta^{13}\text{C}$ values the modern equivalent diet composition are calculated and for this, the changes of the isotopic composition of atmospheric CO_2 through time have to be taken into account. Based on isotopic data from marine foraminifera, the reconstructed $\delta^{13}\text{C}$ value of the atmospheric CO_2 for the Pliocene is about -6.3‰ , ($\delta^{13}\text{C}_{\text{ancient atm CO}_2}$, Pliocene), for the Early Pleistocene it is about -6.5‰ ($\delta^{13}\text{C}_{\text{ancient atm CO}_2}$, Pleistocene), while the modern value was referenced to -8‰ ($\delta^{13}\text{C}_{\text{modern atm}}$, year 2000, Kohn, 2010), respectively (Tipple et al., 2010; Domingo et al., 2013). The modern equivalent diet composition ($\delta^{13}\text{C}_{\text{diet, meq}}$) can be calculated with the equation of Domingo et al. (2013) (Table 2.; Equation (1)). Based on the expected $\delta^{13}\text{C}_{\text{diet, meq}}$ cut-off values for different habitats by Domingo et al. (2013) our data indicate woodland to mesic C_3 grassland (-30‰ to -25‰) as a major flora type in Italy during the Pliocene. Some samples have lower values suggesting the existence of closed-canopy forest but site averages are always higher than -30‰ . There is only one sample that is slightly out of the above ranges (Leffe -23.7‰), where probably local conditions were more enhanced. The value itself would fit the category of open woodland-xeric C_3 grassland (-25‰ to -22‰) (Domingo et al., 2013).

The differences between $\delta^{13}\text{C}$ values could be the result of a complex combination of local climatic control on the vegetation, habitat and dietary differences of the species. Measuring samples

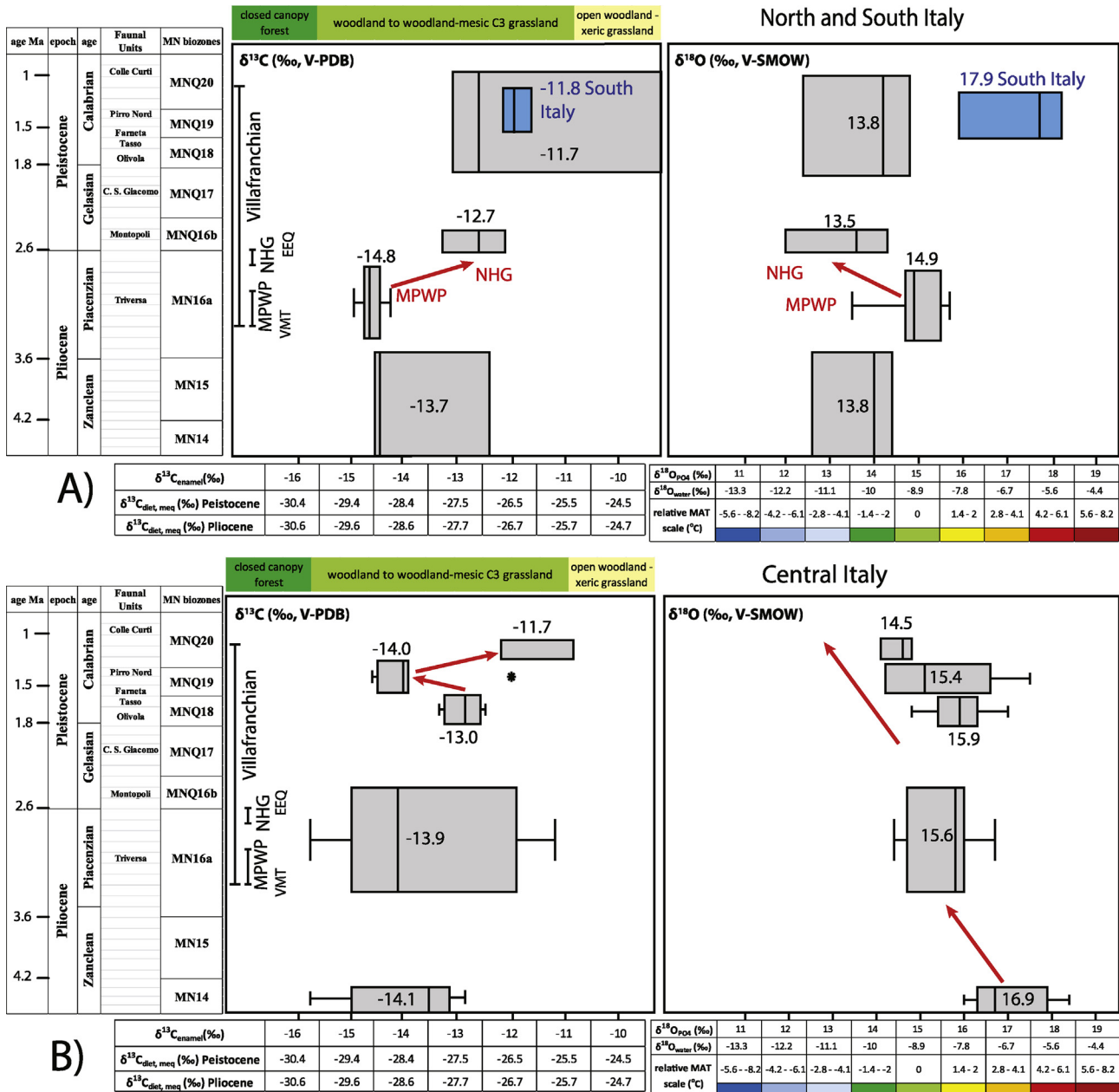


Fig. 2. Average $\delta^{13}\text{C}$ (‰, VPDB) and $\delta^{18}\text{O}_{\text{PO4}}$ (‰, VSMOW) values obtained from the teeth with calculated environmental water values, relative MAT scales and vegetation estimation. A) North Italy B) Central Italy, Vegetation information is from Domingo et al. (2013), environmental water values were calculated by applying Equation (3). (after Kohn and Cerling, 2002), relative MAT scales were calculated using the minimum and maximum slopes of the equations from Table 2. Chronology according to Hilgen et al. (2012). The onset of the Quaternary according to the chronology confirmed in 2009 by the International Union of Geological Sciences. NHG = Northern Hemisphere Glaciation, MPWP = "Mid-Pliocene Warm Period", VMT = Villafranchian mammal turnover, EEQ = Elephant-Equus event, the * sign represent an outlier in Central Italy in MNQ19.

from different taxa from the same location or the same species from different locations would allow a better separation of these different controls. This separation is not always possible due to sample limitations. For example, in this study all of the *S. jeanvireti* samples are from the MN16 biozone and five of the six samples are from North Italy, so different $\delta^{13}\text{C}$ values for this species could indicate dietary differences compared to the other species and/or different climatic signal. Although the results must be interpreted with caution, some implications of these differences can be considered.

Palaeontological observations of *A. arvernensis* suggest that it was a browser taxon that lived in a forest or woodland (Kahlke et al., 2011; Rivals et al., 2015). Morphologically, the bunodont

molars generally imply a soft diet of leaves, fruits and twigs. Their feet were adapted to walk on soft soil, also suggesting that it was an inhabitant of moist woodlands (Ji et al., 2002; Rivals et al., 2015). The dentition changes in the fossil record imply a trend in its diet from soft forest food towards the incorporation of grasses, requiring a grinding component to mastication (Rivals et al., 2015). In agreement with the palaeontological observations, the -13.9‰ average $\delta^{13}\text{C}$ values suggests that *A. arvernensis* lived in woodlands. *S. jeanvireti* seems characteristic for a humid, forest-dominated environment with relatively open areas where gramineous plants could be found (Guérin, 1972, 1980; Lacombat and Mörs, 2008). *S. etruscus* has brachydont dentition and slender, subcursorially structured limbs (Guérin, 1980; Mazza, 1988) suggesting that this

species probably inhabited environments with variable forest cover similar to those in which black rhinos live today, i.e. open scrub woodlands and the margins of small woods and feeding mainly on leaves, shrubs and twigs (Guérin, 1980; Mazza, 1988). At Leffe, palynological and faunal records of a lacustrine basin succession from approximately 1.8 to 1.1 Ma indicate several environmental changes. The rhinoceros species (*S. etruscus* or *S. hundsheimiensis*) lived in warm-temperate dense mixed or conifer forest, to open xerophytic communities and steppe with tree birch and with sparse woodland patches (Ravazzi et al., 2009).

Overall, the palaeontological and environmental observations and the -13‰ to -13.9‰ $\delta^{13}\text{C}$ values are in agreement and suggest that all the investigated taxa could live in forested and partly open environments and were browsers, feeding leaves, twigs shrubs and also grass if these were present in the area. This could also imply that changes in measured $\delta^{13}\text{C}$ values likely the effect of vegetation or climatic changes than change in food preference of the animals.

5.2. Environmental changes based on $\delta^{13}\text{C}$ data

Variation in the measured $\delta^{13}\text{C}$ values in time and space depend on several factors, including vegetation, precipitation and humidity and it is not always possible to separate these effects. For example low $\delta^{13}\text{C}$ values could represent high precipitation and can indicate close canopy forest, or both. However, temporal trends in $\delta^{13}\text{C}$ seem to reflect the environmental and climatic changes in Italy from MN14 to MNQ20. A comparison of the $\delta^{13}\text{C}$ results and the indications of palaeobotanical proxies can be seen in Table 1.

The low $\delta^{13}\text{C}$ values in the Pliocene could represent high humidity, high precipitation values and/or more closed woodlands in North and in Central Italy. The values remain the same in Central Italy, where all the samples from MN16 biozone were averaged. In North Italy, where the samples have a better age control, this biozone can be dissected; a sharp increase in $\delta^{13}\text{C}$ values between the late Pliocene and the early Pleistocene (MN16a – MN16b) is noted. The 1.9‰ increase in $\delta^{13}\text{C}$ values (Fig. 2) could indicate vegetation changes from more closed woodlands to more open areas and/or a large decrease in precipitation or in humidity. If only the drop in MAP had an effect on the $\delta^{13}\text{C}$ values then this drop would be $1250 \pm 630 \text{ mm/year}$ according to the equation of Kohn (2010), (Table 2, Equation (2), a latitude of 45° and an altitude of 150 m was substituted in the equation). The low $\delta^{13}\text{C}$ values in MN16a probably reflect warm and humid climate of the "Mid-Pliocene Warm Period", while the high values in MN16b are most likely linked to the Northern Hemisphere Glaciation. This result can confirm the considerable environmental changes also observed in palynological records with an increase of a cooler type forest (*Picea* and *Fagus*) in North Italy (Stirone section) and a vegetation opening in Central Italy around 2.6 Ma (Bertini, 2001, 2010).

Samples with younger ages are more common from the Central Italian area. Higher $\delta^{13}\text{C}$ for the MNQ18 could indicate drier climate, less precipitation and/or more open vegetation compared to previous biozones. This is in good agreement with pollen records, showing decreased humidity at that time. After 2.6 Ma, the next indication of the vegetation change may be noted for MNQ18 from 1.9 to 1.7 Ma in Central Italy (Bertini, 2001, 2010). During MNQ19 (1.7–1.3 Ma) the $\delta^{13}\text{C}$ values decrease, which could represent wetter interglacial periods and a more closed vegetation in the area. The youngest samples from MNQ19–20 (~1.2 Ma) from Madonna della Strada site show a more marked increase in $\delta^{13}\text{C}$ values. Floral and faunal data suggest environments still characteristic of an interglacial between 1.3 and 1.1 Ma at this site (Magri et al., 2010). The recorded forest phase showed a clear development, starting with an expansion of conifers, progressively replaced by a mixed oak forest, and then showing an increase in Mediterranean

vegetation. An appreciable diffusion of open vegetation is also recorded, as indicated by pollen of grasses and other herbs. The finds of *Mammuthus meridionalis* and "Arvernoceros" or *Eucladoceros giulii* also suggest the presence of open environments. Overall the climate of the site could be similar to present day climate, which is a semi-continental climate, with a transitional Mediterranean-temperate character with about 700 mm precipitation (Magri et al., 2010).

From North Italy only three samples are presented from the younger biozones. These samples from MNQ18–19–20 have higher average $\delta^{13}\text{C}$ values than samples from the Pliocene. The sample from Leffe has the highest $\delta^{13}\text{C}$ value in the database (-9.2‰) indicating open woodland-xeric grassland. As stated by Ravazzi et al. (2009), the observed variable forest cover at the Leffe record documents the occurrence of the rhinoceros (*S. etruscus* or *S. hundsheimiensis*) in cold-temperate, partially open environments, which is the coldest extreme of Early Pleistocene climate cycles registered in the Biogenic Unit of the Leffe Formation. According to this and the high $\delta^{13}\text{C}$ value, the sampled rhinoceros are presumably from the younger part of the site and could live in a more open environment than the other sampled taxa. However, as only one sample was analysed, the results are merely a suggestion as seasonal effects can also be present and there could be individual differences between the animals.

The average $\delta^{13}\text{C}$ values of the three samples from South Italy from MNQ19–20 ($-13.8 \pm 0.3\text{‰}$) are similar to the values in Central and Northern Italy at that time and indicate partly open vegetation.

Overall the temporal changes in $\delta^{13}\text{C}$ values in Italy are in good agreement with the pollen data indicating that the changes in $\delta^{13}\text{C}$ reflect vegetation changes and/or changes in precipitation or in humidity.

5.3. Palaeotemperature changes derived from the $\delta^{18}\text{O}_{\text{PO}_4}$ data

Estimating MAT from $\delta^{18}\text{O}_{\text{PO}_4}$ records require two data conversion steps that are based on well-demonstrated correlations. The first is the correlation between the $\delta^{18}\text{O}_{\text{PO}_4}$ and the $\delta^{18}\text{O}$ values of consumed water (environmental water – $\delta^{18}\text{O}_w$), and the second is the correlation between this $\delta^{18}\text{O}_w$ and the MAT of the site. Several uncertainties and assumptions are inherent to both conversion steps. Different types of equations exist to calculate drinking water of the animal from the $\delta^{18}\text{O}_{\text{PO}_4}$ data. General equations based on very large dataset often show that the $\delta^{18}\text{O}_{\text{PO}_4}$ values of different species can have very similar offsets from the drinking water. The species-specific equations, however, can be more reliable for the given species with the given physiology.

There are more uncertainties in the second step regarding MAT. An assumption is needed for the calculations, such that as the $\delta^{18}\text{O}$ of drinking water represents the $\delta^{18}\text{O}$ of the local mean precipitation, which is not always the case (see effects of evaporation or meltwaters in background section and below in the discussion) and that the estimate of past MAT can be based on present day correlation between MAT and the isotope composition of precipitation. There are local, regional and global equations for the datasets of one IAEA meteorological station or for several stations worldwide. The advantage of using global equations is that they represent an average mid-latitude climate regime and that they are not so sensitive to changes in the local climates with time. Using site-specific precipitation-temperature relationships can, however, give more precise estimates for a local area, notably if it is a mountainous area where local climate variability may be important.

Amongst other influencing factors (different moisture source and air mass trajectories, latitude, altitude, continental and amount effects) $\delta^{18}\text{O}$ of the local precipitation correlates with the MAT. Due to this complex relationship, the correlation between the $\delta^{18}\text{O}$ and

the MAT is changing in time and space, and the correlation may be weak for non-continental or monsoon-influenced stations. In these cases the type of the regression method and the type of the dataset (monthly or annual average precipitation data) becomes an important factor and could influence the palaeoclimatic interpretations (Pryor et al., 2014; Skrzypek et al., 2016). There is currently a debate regarding the regression type to be used for palaeotemperature reconstructions (see Pryor et al., 2014 vs. Skrzypek et al., 2016). Given the above uncertainties, the relative changes in MAT were calculated in this study, instead of absolute MAT values. Results of several equations were compared to observe the possible range of MAT changes linked to the changes in $\delta^{18}\text{O}_{\text{PO}_4}$. The results were calculated using the equations with the highest and the lowest slopes in both conversion steps. The equations used are summarized in Table 2.

In the first step ($\delta^{18}\text{O}_{\text{PO}_4}$ to $\delta^{18}\text{O}_{\text{W}}$), different general equations (Kohn, 1996; Kohn and Cerling, 2002; Amiot et al., 2004) and one species-specific equation (Ayliffe et al., 1992; for proboscidea) were used. In the second step ($\delta^{18}\text{O}_{\text{W}}$ to MAT) several regional and global equations were compared: two equations for Europe (Pryor et al., 2014; Skrzypek et al., 2016), and two global equations (Dansgaard, 1964; Amiot et al., 2004). Additionally a new equation for Italy was produced based on the dataset of the Italian GNIP stations following the work of Skrzypek et al. (2016). The R^2 of this regression is low, but the slope of the equation is very similar to that of the regional and global equations. Equations with similar slopes can be obtained for North and Central Italy as well.

The spatial distribution of the isotopic compositions of precipitation in present day Italy is quite complex. Air masses from different origins have different $\delta^{18}\text{O}$ values and the Apennines and the Alps also have an influence. Longinelli and Selmo (2003) found that the relationship between the isotopic composition of precipitation and the mean monthly temperature values are, on average, very poor. They found no latitudinal gradient along the Tyrrhenian coast from Sicily to the Italian–French border, despite the considerable range of latitude. However, there are 4–5‰ spatial differences in the isotopic composition of precipitations and about 6–7 °C differences in MAT, as a result of the complex topography and climate of Italy, there is not a simple south–north gradient.

The calculated MAT changes between MN14 and MNQ20 in Central Italy compared to other proxies are shown in Table 1 and illustrated in Fig. 2. The 1.4‰ decrease in $\delta^{18}\text{O}_{\text{PO}_4}$ values from MN14 to MN16 can represent a 1.5–1.8‰ change in $\delta^{18}\text{O}_{\text{W}}$ and a 2–3.6 °C decrease in MAT. The cooling trend from Early- to Late Pliocene – to Early Pleistocene is in good agreement with indications of other proxies. Another decrease from MNQ18 to MNQ19–20 is not significant, but with the average shift of −1.4‰ in $\delta^{18}\text{O}_{\text{PO}_4}$ could imply a similar 2–3.6 °C decrease in MAT. Because of the climate in MN19–20 (1.3–1.1 Ma) the Madonna della Strada site was described as an interglacial with thermophilous taxa (Magri et al., 2010), it can be assumed that during the glacial peaks in Early Pleistocene, temperature changes with higher amplitude could have occurred.

In arid climates high evaporation of water reservoirs could raise the $\delta^{18}\text{O}$ values of the animals' drinking water and thus the $\delta^{18}\text{O}_{\text{PO}_4}$ values. If the humidity changed with time, calculations from the $\delta^{18}\text{O}_{\text{PO}_4}$ values would lead to over- or underestimation of MAT changes. In this study for most of the time intervals the $\delta^{18}\text{O}_{\text{PO}_4}$ and $\delta^{13}\text{C}$ values appear to co-vary, likely reflecting the warmer/wetter and cooler/more arid climates. However, some exceptions, for example the MNQ19 zone in Central Italy, could be more humid and slightly cooler than the previous mammal zone, which could imply that samples from that biozone are from a different phase of a glacial-interglacial cycle. Low $\delta^{13}\text{C}$ values in most of the time periods are in agreement with palaeobotanical proxies, indicating humid climates with dense woodlands. In most of these conditions

it is unlikely that strong evaporation could have led to unusual $\delta^{18}\text{O}$ enrichment in water and also phosphate. Some influence is only conceivable in the case of the relatively cooler and more arid time periods such as MNQ18 in Central and MNQ18–19–20 in North or in South Italy but in these cases the $\delta^{13}\text{C}$ values still indicate woodlands so this effect should not be strong.

For Central Italy the decreasing temperatures from Early Pliocene to Late Pliocene – Early Pleistocene followed by fluctuations but a long-term cooling trend towards the glacial-interglacial cycles in Early Pleistocene are supported by the $\delta^{18}\text{O}_{\text{PO}_4}$ in the results and are in agreement with other proxies.

In North Italy there are no large temporal changes in MAT. Whereas, the small increase in $\delta^{18}\text{O}_{\text{PO}_4}$ values between MN14–MN15 and MN16a could be linked to the MPWP, while the following decrease for the MNQ16b can refer to the NHG global cooling event. After this the values stayed relatively constant for the younger time periods. The average $\delta^{18}\text{O}_{\text{PO}_4}$ values are all lower than the values in Central Italy. The measured low $\delta^{18}\text{O}_{\text{PO}_4}$ values could suggest colder MAT, especially during MN14–15, but would contradict other reported proxy data (Fauquette et al., 1999, 2006, Bertini, 2001, 2010). In contrast, while samples from the end of the Zanclean may well have lower $\delta^{18}\text{O}_{\text{PO}_4}$ compared to values from the warmest part of the MPPW, the values being similar to MNQ18–19–20 are considered unlikely.

One explanation for the low $\delta^{18}\text{O}_{\text{PO}_4}$ values could be the influence of glacial meltwaters and river waters draining from the Alps. This would have lower isotopic composition compared to the local precipitation given an altitude effect on the catchment and/or a rain-shadow effect with air mass transport dominated from the north. Therefore, drinking from water source influenced by meltwater could have also lowered the $\delta^{18}\text{O}_{\text{PO}_4}$ values in the teeth. Isotope measurements of waters from the upper part of the Po river support this assumption for a present-day situation. The average composition of the river water is characterized by $\delta^{18}\text{O}$ of −12.5‰, while the $\delta^{18}\text{O}$ values of the precipitation are −7 to −9‰ in the same area (e.g., Marchina et al., 2015; Longinelli and Selmo, 2003). As in Central Italy trends in $\delta^{18}\text{O}_{\text{PO}_4}$ values are in agreement with other proxies, these local source-water effects, are considered more likely compared to a regional-scale cooler climate. If it is the case, the $\delta^{18}\text{O}$ values in North Italy could not be used for MAT calculations unless appropriate equations can be deduced. Nonetheless, samples from more locations and broader time segments could corroborate these hypotheses.

Palynological proxies indicate that the Zanclean MAT and precipitation were higher than the Piacenzian, and the climate in North Italy was comparable to that of the other sites of the north-western Mediterranean region (Table 1.). From the Piacenzian climatic and vegetation characteristics of the middle European region have been suggested (Fauquette and Bertini, 2003). Mediterranean plants are poorly presented in the pollen records and totally lack in plant macrofossil records (Martinetto, 2015 and unpublished data), while the persistence of forests indicates constant humidity. Stable isotope measurements from the Carpathian Basin also support similar results to those for North Italy in the given time period (e.g., Kovács et al., 2015). For a present-day situation, an approximate 1–2‰ difference in $\delta^{18}\text{O}$ values in precipitation and about 1–3 °C differences in MAT are given between areas near Torino and Central Italy (e.g., Longinelli and Selmo, 2003; Combourieu-Nebout et al., 2015). It can be concluded that about 1–2‰ lower $\delta^{18}\text{O}_{\text{pp}}\text{pt}$ values in North Italy can be realistic compared to Central Italy but larger differences may have an additional effect indicated, such as surface waters from the Alpine catchments.

In South Italy the three samples from MNQ19–20 have higher $\delta^{18}\text{O}_{\text{PO}_4}$ values than samples from Central Italy from this period. Two of the three samples are from the Pirro Nord site, with very

high values. This site is close to the Adriatic shore, closer than the other investigated sites. Recent annual average $\delta^{18}\text{O}_{\text{pp}}\text{pt}$ values in that area are about $-5\text{\textperthousand}$, higher than the -6 to $-9\text{\textperthousand}$ for other areas further from the sea, so the measured difference could be the result of the shore-inland gradient in $\delta^{18}\text{O}_{\text{pp}}\text{pt}$ values rather than a south-north temperature gradient at that time (Longinelli and Selmo, 2003).

6. Conclusions

Stable isotope analyses of 51 enamel samples from fossil megaherbivores from North, Central and South Italy (from 5.2 to about 1 Ma) provides information about the Pliocene and Early Pleistocene climate in Italy. Most of the samples have a $\delta^{18}\text{O}_{\text{CO}_3} - \delta^{18}\text{O}_{\text{PO}_4}$ offsets between 7.2 and 10.6%, which is compatible with a range typical for non-altered samples. All of the $\delta^{13}\text{C}$ values indicate that the investigated taxa lived in C_3 ecosystem. The $\delta^{13}\text{C}$ values support a humid climate with woodlands during the Early Pliocene in North and in Central Italy. The Northern Hemisphere Glaciation at 2.7 Ma is indicated as a sharp increase in $\delta^{13}\text{C}$ values between MN16a and MNQ16b in North Italy. After MNQ16b the fluctuations probably represent the moderate glacial-interglacial cycles with a long term increasing trend in $\delta^{13}\text{C}$ values. Higher values during MNQ18 and MNQ19–20 are in good agreement with pollen data indicating more open vegetation. In Central Italy both stable isotope systems change in tandem indicating warmer and more humid or cooler and more arid climates, respectively. The $\delta^{18}\text{O}$ values support the warmest climate during the Early Pliocene followed by cooling with fluctuations toward the end of the Early Pleistocene. In North Italy higher $\delta^{18}\text{O}_{\text{PO}_4}$ values in MN16a are apparent – probably linked to the MPWP – but no other trends can be observed. Attributing the low $\delta^{18}\text{O}_{\text{PO}_4}$ values in North Italy solely to MAT changes would result in low MAT values during the Early Pliocene, in contrast to temperature changes inferred from other proxies (Fauquette et al., 1999, 2006, Bertini, 2001, 2010). A possible explanation could be that the drinking water was influenced by an Alpine catchment with typically lower mean $\delta^{18}\text{O}$ values because of altitude and/or air mass origin differences for the northern parts of Italy. With the exception of the low $\delta^{18}\text{O}$ values in North Italy, the Pliocene vegetation and temperature changes fit well within the trends based on palynological records and changes in mammal fossil assemblages.

Acknowledgements

The research has been generously funded by the Swiss SCIEX program Nr. 13.083. L.P. thanks the European Commission's Research Infrastructure Action, EU-SYNTHESYS project AT-TAF-2550, DE-TAF-3049, GB-TAF-2825, HU-TAF-3593, ES-TAF-2997; part of this research received support from the SYNTHESYS Project <http://www.synthesys.info/> which is financed by European Community Research Infrastructure Action under the FP7 "Capacities" Program. The authors are grateful to Carlo Sarti (Museum of Geology G. Capellini, Bologna, Italy), Daniele Ormezzano (Museum of Geology and Paleontology, University of Turin, Turin, Italy), Ferruccio Farsi and Roberto Mazzei (Natural History Museum, Fisiocritici Academy, Siena, Italy), Carmelo Petronio (Department of Earth Science, Sapienza, University of Rome, Italy) and Loic Costeur (Natural History Museum Basel, Basel, Switzerland) for supplying the fossil materials.

Appendix A. Supplementary data

Supplementary data related to this article can be found at <http://dx.doi.org/10.1016/j.quascirev.2016.11.003>.

References

- Agustí, J., Cabrera, L., Garcés, M., Krijgsman, W., Oms, O., Parés, J.M., 2001. A calibrated mammal scale for the Neogene of Western Europe. State of the art. *Earth-Sci. Rev.* 52, 247–260. [http://dx.doi.org/10.1016/S0012-8252\(00\)00025-8](http://dx.doi.org/10.1016/S0012-8252(00)00025-8).
- Amiot, R., Lécuyer, C., Buffetaut, E., Fluteau, F., Legendre, S., Martineau, F., 2004. Latitudinal temperature gradient during the cretaceous upper campanian–Middle Maastrichtian: $\delta^{18}\text{O}$ record of continental vertebrates. *Earth Planet. Sci. Lett.* 226, 255–272. <http://dx.doi.org/10.1016/j.epsl.2004.07.015>.
- Arppe, L., Karhu, J.A., 2006. Implications for the Late Pleistocene climate in Finland and adjacent areas from the isotopic composition of mammoth skeletal remains. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 231, 322–330. <http://dx.doi.org/10.1016/j.palaeo.2005.08.007>.
- Arppe, L., Karhu, J.A., 2010. Oxygen isotope values of precipitation and the thermal climate in Europe during the middle to late Weichselian ice age. *Quat. Sci. Rev.* 29, 1263–1275. <http://dx.doi.org/10.1016/j.quascirev.2010.02.013>.
- Arppe, L., Aaris-Sørensen, K., Daugnora, L., Lougas, L., Wojtal, P., Zupins, I., 2011. The palaeoenvironmental delta C-13 record in European woolly mammoth tooth enamel. *Quat. Int.* 245, 285–290. <http://dx.doi.org/10.1016/j.quaint.2010.10.018>.
- Ayliffe, L.K., Lister, A.M., Chivas, A.R., 1992. The preservation of glacial-interglacial climatic signatures in the oxygen isotopes of elephant skeletal phosphate. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 99, 179–191. [http://dx.doi.org/10.1016/0031-0182\(92\)90014-v](http://dx.doi.org/10.1016/0031-0182(92)90014-v).
- Ayliffe, L.K., Chivas, A.R., Leakey, M.G., 1994. The retention of primary oxygen isotope compositions of fossil elephant skeletal phosphate. *Geochim. Cosmochim. Acta* 58, 5291–5298. [http://dx.doi.org/10.1016/0016-7037\(94\)90312-3](http://dx.doi.org/10.1016/0016-7037(94)90312-3).
- Bertini, A., 2000. Pollen record from Colle Curti and Cesi: early and middle Pleistocene mammal sites in the Umbro-Marchean Apennine Mountains (central Italy). *J. Quat. Sci.* 15, 825. [http://dx.doi.org/10.1002/1099-1417\(200012\)15:8<825::aid-jqs561>3.0.co;2-6](http://dx.doi.org/10.1002/1099-1417(200012)15:8<825::aid-jqs561>3.0.co;2-6).
- Bertini, A., 2001. Pliocene climatic cycles and altitudinal forest development from 2.7 Ma in the Northern Apennines (Italy): evidence from the pollen record of the Stirone section (~ 5.1 to ~ 2.2 Ma). *Geobios* 34, 253–265. [http://dx.doi.org/10.1016/S0016-6995\(01\)80074-7](http://dx.doi.org/10.1016/S0016-6995(01)80074-7).
- Bertini, A., 2010. Pliocene to Pleistocene palynoflora and vegetation in Italy: State of the art. *Quat. Int.* 225, 5–24. <http://dx.doi.org/10.1016/j.quaint.2010.04.025>.
- Blake, R.E., O'Neil, J.R., Garcia, G.A., 1997. Oxygen isotope systematics of biologically mediated reactions of phosphate. I. Microbial degradation of organophosphorus compounds. *Geochim. Cosmochim. Acta* 61, 4411–4422. [http://dx.doi.org/10.1016/s0016-7037\(97\)00272-x](http://dx.doi.org/10.1016/s0016-7037(97)00272-x).
- Bocherens, H., Pacaud, G., Lazarev, P.A., Mariotti, A., 1996. Stable isotope abundances (C-13, N-15) in collagen and soft tissues from Pleistocene mammals from Yakutia: implications for the palaeobiology of the Mammoth Steppe. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 126, 31–44. [http://dx.doi.org/10.1016/s0031-0182\(96\)00068-5](http://dx.doi.org/10.1016/s0031-0182(96)00068-5).
- Bredenkamp, G.J., Spada, F., Kazmierczak, E., 2002. On the origin of northern and southern hemisphere grasslands. *Plant Ecol.* 163, 209–229. <http://dx.doi.org/10.1023/a:1020957807971>.
- Bruhl, J.J., Wilson, K.L., 2007. Towards a comprehensive survey of C3 and C4 photosynthetic pathways in cyperaceae. *Aliso* 23, 99–148. <http://scholarship.claremont.edu/aliso/vol23/iss1/11>.
- Bryant, J.D., Froelich, P.N., 1995. A model of oxygen isotope fractionation in body water of large mammals. *Geochim. Cosmochim. Acta* 59, 4523–4537. [http://dx.doi.org/10.1016/0016-7037\(95\)00250-4](http://dx.doi.org/10.1016/0016-7037(95)00250-4).
- Bryant, J.D., Koch, P.L., Froelich, P.N., Showers, W.J., Genna, B.J., 1996. Oxygen isotope partitioning between phosphate and carbonate in mammalian apatite. *Geochim. Cosmochim. Acta* 60, 5145–5148. [http://dx.doi.org/10.1016/S0016-7037\(96\)00308-0](http://dx.doi.org/10.1016/S0016-7037(96)00308-0).
- Cerling, T.E., Harris, J.M., 1999. Carbon isotope fractionation between diet and bioapatite in ungulate mammals and implications for ecological and paleoecological studies. *Oecologia* 120, 347–363. <http://dx.doi.org/10.1007/s004420050868>.
- Clauss, M., Polster, C., Kienzle, E., Wiesner, H., Baumgartner, K., von Houwald, F., Streich, W.J., Dierenfeld, E., 2005. Energy and mineral nutrition and water intake in the captive Indian rhinoceros (*Rhinoceros unicornis*). *Zoo. Biol.* 24, 1–14. <http://dx.doi.org/10.1002/zoo.20032>.
- Combouret-Nebout, N., Bertini, A., Russo-Ermolli, E., Peyron, O., Klotz, S., Montade, V., Fauquette, S., Allen, J., Fusco, F., Goring, S., Huntley, B., Joannin, S., Lebreton, V., Magri, D., Martinetto, E., Orain, R., Sadori, L., 2015. Climate changes in the central Mediterranean and Italian vegetation dynamics since the Pliocene. *Rev. Paleobot. Palynol.* 218, 127–147. <http://dx.doi.org/10.1016/j.jrpaleobio.2015.03.001>.
- Dansgaard, W., 1964. Stable isotopes in precipitation. *Tellus* 16, 436–468. <http://dx.doi.org/10.1111/j.2153-3490.1964.tb00181.x>.
- De Schepper, S., Groeneveld, J., Naafs, B.D.A., Van Renterghem, C., Hennissen, J., Head, M.J., Louwye, S., Fabian, K., 2013. Northern Hemisphere Glaciation during the Globally Warm Early Late Pliocene. *Plos One* 8 (12), e81508. <http://dx.doi.org/10.1371/journal.pone.0081508>.
- De Schepper, S., Gibbard, P.L., Salzmann, U., Ehlers, J., 2014. A global synthesis of the marine and terrestrial evidence for glaciation during the Pliocene Epoch. *Earth-Sci. Rev.* 135, 83–102. <http://dx.doi.org/10.1016/j.earscirev.2014.04.003>.
- Dettman, D.L., Kohn, M.J., Quade, J., Ryerson, F.J., Ojha, T.P., Hamidullah, S., 2001. Seasonal stable isotope evidence for a strong Asian monsoon throughout the past 10.7 m.y. *Geology* 29, 31–34. [http://dx.doi.org/10.1130/0091-7613\(2001\)029<0031:SSIEFS>1.0.CO;2](http://dx.doi.org/10.1130/0091-7613(2001)029<0031:SSIEFS>1.0.CO;2).

- 029<0031:ssiefa>2.0.co;2.
- Domingo, L., Koch, P.L., Hernandez Fernandez, M., Fox, D.L., Domingo, M.S., Teresa Alberdi, M., 2013. Late Neogene and early quaternary Paleoenvironmental and Paleoclimatic conditions in southwestern Europe: isotopic analyses on Mammalian Taxa. *Plos One* 8, e63739. <http://dx.doi.org/10.1371/journal.pone.0063739>.
- Dowsett, H., Barron, J., Poore, R., 1996. Middle Pliocene sea surface temperatures: a global reconstruction. *Mar. Micropaleontol.* 27, 13–25. [http://dx.doi.org/10.1016/0377-8398\(95\)00050-X](http://dx.doi.org/10.1016/0377-8398(95)00050-X).
- Dowsett, H.J., Foley, K.M., Stoll, D.K., Chandler, M.A., Sohl, L.E., Bentsen, M., Otto-Btiesner, B.L., Bragg, F.J., Chan, W.L., Contoux, C., Dolan, A.M., Haywood, A.M., Jonas, J.A., Jost, A., Kamae, Y., Lohmann, G., Lunt, D.J., Nisancioglu, K.H., Abe-Ouchi, A., Ramstein, G., Riesselman, C.R., Robinson, M.M., Rosenbloom, N.A., Salzmann, U., Stepanek, C., Strother, S.L., Ueda, H., Yan, Q., Zhang, Z.S., 2013. Sea surface temperature of the mid-Piacenzian ocean: a data-model comparison. *Sci. Rep.* 3 <http://dx.doi.org/10.1038/srep02013> article number 2013.
- Drucker, D.G., Bridault, A., Hobson, K.A., Szuma, E., Bocherens, H., 2008. Can carbon-13 in large herbivores reflect the canopy effect in temperate and boreal ecosystems? Evidence from modern and ancient ungulates. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 266, 69–82. <http://dx.doi.org/10.1016/j.palaeo.2008.03.020>.
- Farquhar, G.D., Ehleringer, J.R., Hubick, K.T., 1989. Carbon isotope discrimination and photosynthesis. *Annu. Rev. Plant. Phys.* 40, 503–537. <http://dx.doi.org/10.1146/annurev.pp.40.060189.002443>.
- Fauquette, S., Bertini, A., 2003. Quantification of the northern Italy Pliocene climate from pollen data: evidence for a very peculiar climate pattern. *Boreas* 32, 361–369. <http://dx.doi.org/10.1080/03009480310002235>.
- Fauquette, S., Suc, J.P., Guiot, J., Diniz, F., Feddi, N., Zheng, Z., Bessais, E., Drivaliari, A., 1999. Climate and biomes in the west Mediterranean area during the Pliocene. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 152, 15–36. [http://dx.doi.org/10.1016/s0031-0182\(99\)00031-0](http://dx.doi.org/10.1016/s0031-0182(99)00031-0).
- Fauquette, S., Suc, J.P., Bertini, A., Popescu, S.M., Warny, S., Taoufiq, N.B., Villa, M.J.P., Chikhi, H., Feddi, N., Subally, D., Clauzon, G., Ferrier, J., 2006. How much did climate force the Messinian salinity crisis? Quantified climatic conditions from pollen records in the Mediterranean region. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 238, 281–301. <http://dx.doi.org/10.1016/j.palaeo.2006.03.029>.
- Feranec, R.S., 2004. Geographic variation in the diet of hypodont herbivores from the Rancholabrean of Florida. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 207, 359–369. <http://dx.doi.org/10.1016/j.palaeo.2003.09.031>.
- Feranec, R.S., MacFadden, B.J., 2006. Isotopic discrimination of resource partitioning among ungulates in C3-dominated communities from the Miocene of Florida and California. *Paleobiology* 32, 191–205. <http://dx.doi.org/10.1666/05006.1>.
- Forno, M.G., Gattiglio, M., Comina, C., Barbero, D., Bertini, A., Doglione, A., Gianotti, F., Irace, A., Martinetto, M., Mottura, A., Sala, B., 2015. Stratigraphic and tectonic notes on the Villafranca d'Asti succession in the type-area and Castelnovo Don Bosco sector (Asti reliefs, Piedmont). *A. M. Q.* 28, 5–27.
- Fricke, H.C., Clyde, W.C., O'Neil, J.R., 1998. Intra-tooth variations in delta O-18 (PO4) of mammalian tooth enamel as a record of seasonal variations in continental climate variables. *Geochim. Cosmochim. Acta* 62, 1839–1850. [http://dx.doi.org/10.1016/0016-7037\(98\)00114-8](http://dx.doi.org/10.1016/0016-7037(98)00114-8).
- Garcia-Alix, A., 2015. A multiproxy approach for the reconstruction of ancient continental environments. The case of the Mio–Pliocene deposits of the Granada Basin (southern Iberian Peninsula). *Glob. Planet. Change* 131, 1–10. <http://dx.doi.org/10.1016/j.gloplacha.2015.04.005>.
- Guérin, C., 1972. Une nouvelle espèce de Rhinocéros (Mammalia, Perissodactyla) à Viallette (Haute-Loire, France) et dans d'autres gisements du Villafranchien Inférieur Européen: *Dicerorhinus jeanvireti* n. sp. *Doc. Lab. Géol. Fac. Sci. Lyon* 49, 53–161.
- Guérin, C., 1980. Les rhinocéros (Mammalia, Perissodactyla) du Miocene terminal au Pléistocène supérieur en Europe occidentale. Comparaison avec les espèces actuelles. *Doc. Lab. Géol. Fac. Sci. Lyon* 79, 1–1183.
- Halas, S., Skryzpek, G., Meier-Augenstein, W., Pelc, A., Kemp, H.F., 2011. Inter-laboratory calibration of new silver orthophosphate comparison materials for the stable oxygen isotope analysis of phosphates. *Rapid Commun. Mass Spectrom.* 25, 579–584. <http://dx.doi.org/10.1002/rcm.4892>.
- Hartman, G., Bar-Yosef, O., Brittingham, A., Grosman, L., Munro, N.D., 2016. Hunted gazelles evidence cooling, but not drying, during the Younger Dryas in the southern Levant. *Proc. Natl. Acad. Sci.* 113, 3997–4002. <http://dx.doi.org/10.1073/pnas.1519862113>.
- Haywood, A.M., Hill, D.J., Dolan, A.M., Otto-Btiesner, B.L., Bragg, F., Chan, W.L., Chandler, M.A., Contoux, C., Dowsett, H.J., Jost, A., Kamae, Y., Lohmann, G., Lunt, D.J., Abe-Ouchi, A., Pickering, S.J., Ramstein, G., Rosenbloom, N.A., Salzmann, U., Sohl, L., Stepanek, C., Ueda, H., Yan, Q., Zhang, Z., 2013. Large-scale features of Pliocene climate: results from the Pliocene model intercomparison project. *Clim. Past* 9, 191–209. <http://dx.doi.org/10.5194/cp-9-191-2013>.
- Hilgen, F.J., Lourens, L.J., Van Dam, J.A., Beu, A.G., Boyes, A.F., Cooper, R.A., Krijgsman, W., Ogg, J.G., Piller, W.E., Wilson, D.S., 2012. The Neogene period. In: Gradstein, F.M., Schmitz, J.G.O.D., Ogg, G.M. (Eds.), *The Geologic Time Scale*. Elsevier, Boston, pp. 923–978.
- Hill, D.J., Csank, A.Z., Dolan, A.M., Lunt, D.J., 2011. Pliocene climate variability: northern Annular Mode in models and tree-ring data. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 309, 118–127. <http://dx.doi.org/10.1016/j.palaeo.2011.04.003>.
- Hoppe, K.A., Koch, P.L., 2007. Reconstructing the migration patterns of late Pleistocene mammals from northern Florida, USA. *Quat. Res.* 68, 347–352. <http://dx.doi.org/10.1016/j.yqres.2007.08.001>.
- Iacumin, P., Bocherens, H., Mariotti, A., Longinelli, A., 1996. Oxygen isotope analyses of co-existing carbonate and phosphate in biogenic apatite: a way to monitor diagenetic alteration of bone phosphate? *Earth Planet. Sci. Lett.* 142, 1–6. [http://dx.doi.org/10.1016/0012-821x\(96\)00093-3](http://dx.doi.org/10.1016/0012-821x(96)00093-3).
- Iacumin, P., Nikolaev, V., Ramigni, M., 2000. C and N stable isotope measurements on Eurasian fossil mammals, 40 000 to 10 000 years BP: herbivore physiologies and palaeoenvironmental reconstruction. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 163, 33–47. [http://dx.doi.org/10.1016/s0031-0182\(00\)00141-3](http://dx.doi.org/10.1016/s0031-0182(00)00141-3).
- Irace, A., Monegato, G., Tema, E., Martinetto, E., Gianolla, D., Vassio, E., Bellino, L., Violanti, D., 2015. Unconformity-bounded stratigraphy in the Plio-Pleistocene continental record: new insights from the Alessandria Basin (NW Italy). *Geol. J.* 2015 <http://dx.doi.org/10.1002/gj.2744> [Published online].
- Ji, Q., Luo, Z.-X., Yuan, C.-X., Wible, J.R., Zhang, J.-P., Georgi, J.A., 2002. The earliest known eutherian mammal. *Nature* 416, 816–822. <http://dx.doi.org/10.1038/416816a>.
- Kahlke, R.-D., García, N., Kostopoulos, D.S., Lacombat, F., Lister, A.M., Mazza, P.P.A., Spassov, N., Titov, V.V., 2011. Western Palaearctic palaeoenvironmental conditions during the Early and early Middle Pleistocene inferred from large mammal communities, and implications for hominin dispersals in Europe. *Quat. Sci. Rev.* 30, 1368–1395. <http://dx.doi.org/10.1016/j.quascirev.2010.07.020>.
- Klotz, S., Fauquette, S., Combourieu-Nebout, N., Uhl, D., Suc, J.-P., Mosbrugger, V., 2006. Seasonality intensification and long-term winter cooling as a part of the Late Pliocene climate development. *Earth Planet. Sci. Lett.* 241, 174–187. <http://dx.doi.org/10.1016/j.epsl.2005.10.005>.
- Koch, P.L., 1998. Isotopic reconstruction of past continental environments. *Annu. Rev. Earth Planet. Sci.* 26, 573–613. <http://dx.doi.org/10.1146/annurev.earth.26.1.573>.
- Koch, P.L., Tuross, N., Fogel, M.L., 1997. The effects of sample treatment and diagenesis on the isotopic integrity of carbonate in biogenic hydroxylapatite. *J. Archaeol. Sci.* 24, 417–429. <http://dx.doi.org/10.1006/jasc.1996.0126>.
- Kocsis, L., 2011. Geochemical compositions of marine fossils as proxies for reconstructing ancient environmental conditions. *Chimia* 65, 787–791. <http://dx.doi.org/10.2533/chimia.2011.787>.
- Kocsis, L., Ozsvart, P., Becker, D., Ziegler, R., Scherler, L., Codrea, V., 2014. Orogeny forced terrestrial climate variation during the late Eocene–early Oligocene in Europe. *Geology* 42, 727–730. <http://dx.doi.org/10.1130/g35673.1>.
- Kohn, M.J., 1996. Predicting animal delta O-18: Accounting for diet and physiological adaptation. *Geochim. Cosmochim. Acta* 60, 4811–4829. [http://dx.doi.org/10.1016/s0016-7037\(96\)00240-2](http://dx.doi.org/10.1016/s0016-7037(96)00240-2).
- Kohn, M.J., 2010. Carbon isotope compositions of terrestrial C3 plants as indicators of (paleo)ecology and (paleo)climate. *Proc. Natl. Acad. Sci.* 107, 19691–19695. <http://dx.doi.org/10.1073/pnas.1004933107>.
- Kohn, M.J., Cerling, T.E., 2002. Stable isotope compositions of biological apatite. *Rev. Mineral. Geochem.* 48, 455–488. <http://dx.doi.org/10.2138/rmg.2002.48.12>.
- Kohn, M.J., Schoeninger, M.J., Barker, W.W., 1999. Altered states: effects of diagenesis on fossil tooth chemistry. *Geochim. Cosmochim. Acta* 63, 2737–2747. [http://dx.doi.org/10.1016/s0016-7037\(99\)00208-2](http://dx.doi.org/10.1016/s0016-7037(99)00208-2).
- Kovács, J., Szabó, P., Kocsis, L., Vennemann, T., Sabol, M., Gasparik, M., Virág, A., 2015. Pliocene and Early Pleistocene paleoenvironmental conditions in the Pannonian Basin (Hungary, Slovakia): stable isotope analyses of fossil proboscidean and perissodactyl teeth. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 440, 455–466. <http://dx.doi.org/10.1016/j.palaeo.2015.09.019>.
- Kürschner, W.M., 2010. C-isotope composition of fossil sedges and grasses. *Geophys. Res. Abstr.* 12. <http://dx.doi.org/10.1029/EGU2010-8012-1>.
- Lacombat, F., Mors, T., 2008. The northernmost occurrence of the rare Late Pliocene rhinoceros *Stephanorhinus jeanvireti* (Mammalia, Perissodactyla). *Neues Jahrb. Geol. P.-A.* 249, 157–165. <http://dx.doi.org/10.1127/0077-7749/2008/0249-0157>.
- Leethorp, J.A., Sealy, J.C., Vandermerwe, N.J., 1989. Stable carbon isotope ratio differences between bone collagen and bone apatite, and their relationship to diet. *J. Archaeol. Sci.* 16, 585–599. [http://dx.doi.org/10.1016/0305-4403\(89\)90024-1](http://dx.doi.org/10.1016/0305-4403(89)90024-1).
- Levin, N.E., Cerling, T.E., Passey, B.H., Harris, J.M., Ehleringer, J.R., 2006. A stable isotope aridity index for terrestrial environments. *Proc. Natl. Acad. Sci.* 103, 11201–11205. <http://dx.doi.org/10.1073/pnas.0604719103>.
- Liang, Y., Blake, R.E., 2007. Oxygen isotope fractionation between apatite and aqueous-phase phosphate: 20–45 degrees C. *Chem. Geol.* 238, 121–133. <http://dx.doi.org/10.1016/j.chemgeo.2006.11.004>.
- Loftus, E., Stewart, B.A., Dewar, G., Lee-Thorp, J., 2015. Stable isotope evidence of late MIS 3 to middle Holocene palaeoenvironments from Sehonghong Rockshelter, eastern Lesotho. *J. Quat. Sci.* 30, 805–816. <http://dx.doi.org/10.1002/jqs.2817>.
- Longinelli, A., 1984. Oxygen isotopes in mammal bone phosphate: a new tool for paleohydrological and paleoclimatological research? *Geochim. Cosmochim. Acta* 48, 385–390. [http://dx.doi.org/10.1016/0016-7037\(84\)90259-x](http://dx.doi.org/10.1016/0016-7037(84)90259-x).
- Longinelli, A., Selmo, E., 2003. Isotopic composition of precipitation in Italy: a first overall map. *J. Hydrol.* 270, 75–88. [http://dx.doi.org/10.1016/S0022-1694\(02\)00281-0](http://dx.doi.org/10.1016/S0022-1694(02)00281-0).
- Luz, B., Kolodny, Y., 1985. Oxygen isotope variations in phosphate of biogenic apatites. 4. Mammal teeth and bones. *Earth Planet. Sci. Lett.* 75, 29–36. [http://dx.doi.org/10.1016/0012-821x\(85\)90047-0](http://dx.doi.org/10.1016/0012-821x(85)90047-0).
- Luz, B., Kolodny, Y., Horowitz, M., 1984. Fractionation of oxygen isotopes between mammalian bone-phosphate and environmental drinking water. *Geochim. Cosmochim. Acta* 48, 1689–1693. [http://dx.doi.org/10.1016/0016-7037\(84\)90338-7](http://dx.doi.org/10.1016/0016-7037(84)90338-7).
- Luz, B., Cormie, A.B., Schwarcz, H.P., 1990. Oxygen isotope variations in phosphate of deer bones. *Geochim. Cosmochim. Acta* 54, 1723–1728. [http://dx.doi.org/10.1016/0016-7037\(90\)90338-7](http://dx.doi.org/10.1016/0016-7037(90)90338-7).

- 10.1016/0016-7037(90)90403-8.
- Magri, D., Di Rita, F., Palombo, M.R., 2010. An Early Pleistocene interglacial record from an intermontane basin of central Italy (Scoppito, L'Aquila). *Quat. Int.* 225, 106–113. <http://dx.doi.org/10.1016/j.quaint.2009.04.005>.
- Marchina, C., Bianchini, G., Natali, C., Pennisi, M., Colombani, N., Tassinari, R., Knoeller, K., 2015. The Po river water from the Alps to the Adriatic Sea (Italy): new insights from geochemical and isotopic ($\delta^{18}\text{O}$ - δD) data. *Environ. Sci. Pollut. Res.* 22, 5184–5203. <http://dx.doi.org/10.1007/s11356-014-3750-6>.
- Martin, C., Bentaleb, I., Kaandorp, R., Iacumin, P., Chatri, K., 2008. Intra-tooth study of modern rhinoceros enamel $\delta^{18}\text{O}$: is the difference between phosphate and carbonate $\delta^{18}\text{O}$ a sound diagenetic test? *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 266, 183–189. <http://dx.doi.org/10.1016/j.palaeo.2008.03.039>.
- Martinelli, L.A., Devol, A.H., Victoria, R.L., Richey, J.E., 1991. Stable carbon isotope variation in C3 and C4 plants along the Amazon River. *Nature* 353, 57–59.
- Martinetto, E., 2015. Monographing the Pliocene and Early Pleistocene carpal floras of Italy: methodological challenges and current progress. *Palaeontogr. Abt. B* 293, 57–99.
- Martinetto, E., Momohara, A., Bizzarri, R., Baldanza, A., Delfino, M., Esu, D., Sardella, R., 2015. Late persistence and deterministic extinction of humid thermophilic plant taxa of East Asian affinity (HUTEA) in southern Europe. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* <http://dx.doi.org/10.1016/j.palaeo.2015.08.015>. Online version.
- Mazza, P., 1988. The Tuscan early Pleistocene rhinoceros *Dicerorhinus etruscus*. *Palaeontogr. Ital.* 75, 1–87.
- Metcalfe, J.Z., Longstaffe, F.J., 2012. Mammoth tooth enamel growth rates inferred from stable isotope analysis and histology. *Quat. Res.* 77, 424–432. <http://dx.doi.org/10.1016/j.yqres.2012.02.002>.
- Metcalfe, J.Z., Longstaffe, F.J., White, C.D., 2009. Method-dependent variations in stable isotope results for structural carbonate in bone bioapatite. *J. Archaeol. Sci.* 36, 110–121. <http://dx.doi.org/10.1016/j.jas.2008.07.019>.
- Metcalfe, J.Z., Longstaffe, F.J., Zazula, G.D., 2010. Nursing, weaning, and tooth development in woolly mammoths from Old Crow, Yukon, Canada: implications for Pleistocene extinctions. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 298, 257–270. <http://dx.doi.org/10.1016/j.palaeo.2010.09.032>.
- Metcalfe, J.Z., Longstaffe, F.J., Jass, C.N., Zazula, G.D., Keddie, G., 2016. Taxonomy, location of origin and health status of proboscideans from Western Canada investigated using stable isotope analysis. *J. Quat. Sci.* 31, 126–142. <http://dx.doi.org/10.1002/jqs.2849>.
- Meyers, S.R., Hinnov, L.A., 2010. Northern Hemisphere glaciation and the evolution of Plio-Pleistocene climate noise. *Paleoceanography* 25, PA3207. <http://dx.doi.org/10.1029/2009pa001834>.
- Mishra, S., Tripathi, A., Tripathi, D.K., Chauhan, D.K., 2015. Role of sedges (Cyperaceae) in wetlands, environmental cleaning and as food material: Possibilities and future perspectives. In: Azooz, M.M., Ahmad, P. (Eds.), *Plant-environment Interaction: Responses and Approaches to Mitigate Stress*. John Wiley & Sons Ltd., Chichester, pp. 327–338. <http://dx.doi.org/10.1002/9781119081005.ch18>.
- Montanari, S., Louys, J., Price, G.J., 2013. Pliocene Paleoenvironments of South-eastern Queensland, Australia inferred from stable isotopes of marsupial tooth enamel. *Plos One* 8, e66221. <http://dx.doi.org/10.1371/journal.pone.0066221>.
- O'Leary, M.H., 1988. Carbon isotopes in photosynthesis. *BioScience* 38, 328–336. <http://dx.doi.org/10.2307/1310735>.
- Palombo, M.R., 2007. What is the boundary for the Quaternary period and Pleistocene epoch? The contribution of turnover patterns in large mammalian complexes from north-western Mediterranean to the debate. *Quaternaire* 18, 35–53.
- Passey, B.H., Robinson, T.F., Ayliffe, L.K., Cerling, T.E., Sponheimer, M., Dearing, M.D., Roeder, B.L., Ehrlinger, J.R., 2005. Carbon isotope fractionation between diet, breath CO_2 , and bioapatite in different mammals. *J. Archaeol. Sci.* 32, 1459–1470. <http://dx.doi.org/10.1016/j.jas.2005.03.015>.
- Pellegrini, M., Snoeck, C., 2016. Comparing bioapatite carbonate pre-treatments for isotopic measurements: Part 2 — impact on carbon and oxygen isotope compositions. *Chem. Geol.* 420, 88–96. <http://dx.doi.org/10.1016/j.chemgeo.2015.10.038>.
- Pellegrini, M., Lee-Thorp, J.A., Donahue, R.E., 2011. Exploring the variation of the delta O-18(p) and delta O-18(c) relationship in enamel increments. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 310, 71–83. <http://dx.doi.org/10.1016/j.palaeo.2011.02.023>.
- Petronio, C., Bellucci, L., Martinetto, E., Pandolfi, L., Salari, L., 2011. Biochronology and palaeoenvironmental changes from the middle Pliocene to the late Pleistocene in Central Italy. *Geodiversitas* 33, 485–517. <http://dx.doi.org/10.5252/g2011n3a4>.
- Pontini, M.R., Bertini, A., 2000. Late Pliocene vegetation and climate in Central Italy: high-resolution pollen analysis from the Fosso Bianco succession (Tiberino Basin). *Geobios* 33, 519–526. [http://dx.doi.org/10.1016/s0016-6995\(00\)80024-8](http://dx.doi.org/10.1016/s0016-6995(00)80024-8).
- Pryor, A.J.E., Stevens, R.E., O'Connell, T.C., Lister, J.R., 2014. Quantification and propagation of errors when converting vertebrate biomineral oxygen isotope data to temperature for palaeoclimate reconstruction. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 412, 99–107. <http://dx.doi.org/10.1016/j.palaeo.2014.07.003>.
- Pushkina, D., Bocherens, H., Ziegler, R., 2014. Unexpected palaeoecological features of the Middle and Late Pleistocene large herbivores in southwestern Germany revealed by stable isotopic abundances in tooth enamel. *Quat. Int.* 339, 164–178. <http://dx.doi.org/10.1016/j.quaint.2013.12.033>.
- Quade, J., Cerling, T.E., Barry, J.C., Morgan, M.E., Pilbeam, D.R., Chivas, A.R., Leethorp, J.A., Vandermerwe, N.J., 1992. A 16-Ma record of paleodiet using carbon and oxygen isotopes in fossil teeth from Pakistan. *Chem. Geol.* 94, 183–192. [http://dx.doi.org/10.1016/0168-9622\(92\)90011-x](http://dx.doi.org/10.1016/0168-9622(92)90011-x).
- Ravazzi, C., Pini, R., Breda, M., 2009. Reconstructing the palaeoenvironments of the early Pleistocene mammal faunas from the pollen preserved on fossil bones. *Quat. Sci. Rev.* 28, 2940–2954. <http://dx.doi.org/10.1016/j.quascirev.2009.07.022>.
- Rivals, F., Mol, D., LaCombat, F., Lister, A.M., Semprebon, G.M., 2015. Resource partitioning and niche separation between mammoths (*Mammuthus rumanus* and *Mammuthus meridionalis*) and gomphotheres (*Anancus arvernensis*) in the Early Pleistocene of Europe. *Quat. Int.* 379, 164–170. <http://dx.doi.org/10.1016/j.quaint.2014.12.031>.
- Rook, L., Martínez-Narváez, B., 2010. Villafranchian: the long story of a Plio-Pleistocene European large mammal biochronologic unit. *Quat. Int.* 219, 134–144. <http://dx.doi.org/10.1016/j.quaint.2010.01.007>.
- Rozanski, K., Araguás-Araguás, L., Gonfiantini, R., 1993. Isotopic patterns in modern global precipitation. In: Swart, P.K., Lohmann, K.C., McKenzie, J., Savin, S. (Eds.), *Climate Change in Continental Isotopic Records*. American Geophysical Union, Washington, DC, pp. 1–36.
- Russo Ermolli, E., Sardella, R., Di Maio, G., Petronio, C., Santangelo, N., 2010. Pollen and mammals from the late Early Pleistocene site of Saticula (Sant'Agata de' Goti, Benevento, Italy). *Quat. Int.* 225, 128–137. <http://dx.doi.org/10.1016/j.quaint.2009.06.013>.
- Salzmann, U., Williams, M., Haywood, A.M., Johnson, A.L.A., Kender, S., Zalasiewicz, J., 2011. Climate and environment of a Pliocene warm world. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 309, 1–8. <http://dx.doi.org/10.1016/j.palaeo.2011.05.044>.
- Scherler, L., Tütken, T., Becker, D., 2014. Carbon and oxygen stable isotope compositions of late Pleistocene mammal teeth from dolines of Ajoie (Northwestern Switzerland). *Quat. Res.* 82, 378–387. <http://dx.doi.org/10.1016/j.yqres.2014.05.004>.
- Skrzypek, G., Sadler, R., Wiśniewski, A., 2016. Reassessment of recommendations for processing mammal phosphate $\delta^{18}\text{O}$ data for paleotemperature reconstruction. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 446, 162–167. <http://dx.doi.org/10.1016/j.palaeo.2016.01.032>.
- Spötl, C., Vennemann, T.W., 2003. Continuous-flow isotope ratio mass spectrometric analysis of carbonate minerals. *Rapid Commun. Mass Spectrom.* 17, 1004–1006. <http://dx.doi.org/10.1002/rclm.1010>.
- Sullivan, C.H., Krueger, H.W., 1981. Carbon isotope analysis of separate chemical phases in modern and fossil bone. *Nature* 292, 333–335. <http://dx.doi.org/10.1038/292333a0>.
- Tafforeau, P., Bentaleb, I., Jaeger, J.-J., Martin, C., 2007. Nature of laminations and mineralization in rhinoceros enamel using histology and X-ray synchrotron microtomography: potential implications for palaeoenvironmental isotopic studies. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 246, 206–227. <http://dx.doi.org/10.1016/j.palaeo.2006.10.001>.
- Teodoridis, V., Bruch, A.A., Vassio, E., Martinetto, E., Kvaček, Z., Stuchlík, L., 2015. Plio-Pleistocene floras of the Vildštejn formation in the Cheb Basin, Czech Republic — a floristic and palaeoenvironmental review. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* <http://dx.doi.org/10.1016/j.palaeo.2015.09.038>. Online version.
- Thorp, J.L., Van Der Merwe, N.J., 1987. Carbon isotope analysis of fossil bone apatite. *S. Afr. J. Sci.* 83, 712–715.
- Tipple, B.J., Meyers, S.R., Pagani, M., 2010. Carbon isotope ratio of Cenozoic CO_2 : a comparative evaluation of available geochemical proxies. *Paleoceanography* 25, <http://dx.doi.org/10.1029/2009pa001851>.
- Tütken, T., Vennemann, T.W., Janz, H., Heimann, E.P.J., 2006. Palaeoenvironment and palaeoclimate of the Middle Miocene lake in the Steinheim basin, SW Germany: a reconstruction from C, O, and Sr isotopes of fossil remains. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 241, 457–491. <http://dx.doi.org/10.1016/j.palaeo.2006.04.007>.
- Tütken, T., Furrer, H., Vennemann, T.W., 2007. Stable isotope compositions of mammoth teeth from Niederenningen, Switzerland: implications for the Late Pleistocene climate, environment, and diet. *Quat. Int.* 164–165, 139–150. <http://dx.doi.org/10.1016/j.quaint.2006.09.004>.
- Tütken, T., Kaiser, T.M., Vennemann, T., Merceron, G., 2013. Opportunistic feeding Strategy for the earliest old world hypsodont equids: evidence from stable isotope and dental wear proxies. *Plos One* 8, e74463. <http://dx.doi.org/10.1371/journal.pone.0074463>.
- Van der Merwe, N.J., Medina, E., 1991. The canopy effect, carbon isotope ratios and foodwebs in Amazonia. *J. Archaeol. Sci.* 18, 249–259. [http://dx.doi.org/10.1016/0305-4403\(91\)90064-V](http://dx.doi.org/10.1016/0305-4403(91)90064-V).
- Vennemann, T.W., Fricke, H.C., Blake, R.E., O'Neil, J.R., Colman, A., 2002. Oxygen isotope analysis of phosphates: a comparison of techniques for analysis of Ag_3PO_4 . *Chem. Geol.* 185, 321–336. [http://dx.doi.org/10.1016/s0009-2541\(01\)00413-2](http://dx.doi.org/10.1016/s0009-2541(01)00413-2).
- Wang, Y., Cerling, T.E., 1994. A model of fossil tooth and bone diagenesis: implications for paleodiet reconstruction from stable isotopes. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 107, 281–289. [http://dx.doi.org/10.1016/0031-0182\(94\)90100-7](http://dx.doi.org/10.1016/0031-0182(94)90100-7).
- Woodard, S.C., Rosenthal, Y., Miller, K.G., Wright, J.D., Chiu, B.K., Lawrence, K.T., 2014. Antarctic role in northern hemisphere glaciation. *Science* 346, 847–851. <http://dx.doi.org/10.1126/science.1255586>.
- Zanazzi, A., Judd, E., Fletcher, A., Bryant, H., Kohn, M.J., 2015. Eocene-Oligocene latitudinal climate gradients in North America inferred from stable isotope

- ratios in perissodactyl tooth enamel. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* 417, 561–568. <http://dx.doi.org/10.1016/j.palaeo.2014.10.024>.
- Zazzo, A., Lecuyer, C., Mariotti, A., 2004a. Experimentally-controlled carbon and oxygen isotope exchange between bioapatites and water under inorganic and microbially-mediated conditions. *Geochim. Cosmochim. Acta* 68, 1–12. [http://dx.doi.org/10.1016/s0016-7037\(03\)00278-3](http://dx.doi.org/10.1016/s0016-7037(03)00278-3).
- Zazzo, A., Lecuyer, C., Sheppard, S.M.F., Grandjean, P., Mariotti, A., 2004b. Diagenesis and the reconstruction of paleoenvironments: a method to restore original delta O-18 values of carbonate and phosphate from fossil tooth enamel. *Geochim. Cosmochim. Acta* 68, 2245–2258. <http://dx.doi.org/10.1016/j.gca.2003.11.009>.