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Small mammal tooth enamel carbon isotope record of C₄ grasses in late Neogene China



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ABSTRACT

The spatiotemporal pattern of the late Cenozoic spread of C₄ vegetation is an important indicator of environmental change that is intertwined with the uplift of the Himalaya and Tibetan Plateau, and the development of the East Asian monsoons. To explore the spread of C4 vegetation in China and shed new light on regional climatic evolution, we measured δ^{13} C values of more than 200 small mammal teeth (primarily rodents and lagomorphs) using a laser ablation isotope ratio mass spectrometry approach. Small mammals are highly sensitive indicators of their environment because they have limited spatial ranges and because they have minimal time-averaging of carbon isotope signatures of dietary components. The specimens originate from four classic Late Miocene fossil localities, Lufeng, Yuanmou, Lingtai, and Ertemte, along a southwest-northeast transect from Yunnan Province to Inner Mongolia. In Yunnan (Lufeng, Yuanmou) and on the Loess Plateau (Lingtai), the small mammal δ^{13} C values record nearly pure C₃ ecosystems, and mixed but C₃-based ecosystems, respectively, in agreement with previous studies based on carbon isotopes of large herbivores and soil carbonates. In Inner Mongolia, the micromammalian tooth enamel δ^{13} C record picks up the presence of C₄ vegetation where large mammal samples do not, indicating a mixed yet C₃-dominated ecosystem at ~6 Ma. As a whole, the results support a scenario of northward increasing C₄ grass abundance in a pattern that mirrors northward decreasing precipitation of the summer monsoon system. The results highlight differences between large and small mammals as indicators of C_4 vegetation in ancient ecosystems, particularly the ability of small mammal $\delta^{13}C$ values to detect the presence of minor components of the vegetation structure.

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1. Introduction

One of the most striking ecological transformations of the Cenozoic was the global emergence and expansion of C₄ grasses during the late Miocene and Pliocene, and the attendant transformation of many lineages of mammalian herbivores, with decline of forest-dwelling browsers and success of grazers of the open plains (Cerling et al., 1997; Janis et al., 2000, 2004; see Strömberg, 2011 for a review). These changes are iconic examples of the overall trend of cooling and aridification that occurred since the Middle Miocene Climatic Optimum ca. 15 Ma ago. Yet, the global signal includes considerable temporal and regional variation, with some regions changing faster than others or even temporarily reversing the direction of change. In the Eurasian realm, for example, precipitation increased during the late Late Miocene in northern China while the Mediterranean experienced arid conditions

* Corresponding author. E-mail address: laura.arppe@helsinki.fi (L. Arppe). (Fortelius et al., 2002; Eronen et al., 2010; Fortelius et al., 2014). Aridification was, in turn, more profound and occurred earlier in North America than in Europe (Eronen et al., 2012). In Asia, one of the main factors responsible for this heterogeneity is the regional development of monsoonal climate systems (Wang et al., 2005; Molnar et al., 2010; Tang et al., 2015).

The East Asian Monsoon is the most important monsoon system of the late Neogene. This system can be traced back to the early Cenozoic (Sun and Wang, 2005), and although considerable uncertainty remains, there is evidence from multiple proxies that a marked strengthening of the system occurred in the late Miocene (An et al., 2001; Molnar et al., 2010; Tang, 2013). Since the fundamental characteristic of a monsoon system is seasonal fluctuation between two opposing states, e.g., of precipitation and dominant wind direction (Ramage, 1971; Webster et al., 1998), a strengthened monsoon almost inevitably means strengthened seasonality. Seasonality of rainfall and temperature, in conjunction with mean values of those parameters, is the dominant control on the basic nature of prevailing ecosystems, for example, evergreen forest, deciduous forest, savanna, or grassland (e.g. Breckle, 2002). One of the most intensely studied phenomena that is often placed in a context of monsoon history is the expansion of grasses using C_4 photosynthesis, the so-called C_4 grasses, which are favored in regions with warm growing seasons, and in conditions of reduced atmospheric CO_2 concentration (Ehleringer and Björkman, 1977; Ehleringer et al., 1997). The rise of C_4 grasses dates back to the early Oligocene (but see Vicentini et al., 2008) with rare evidence of occurrence from the Oligocene to middle Miocene (Strömberg, 2005; Tipple and Pagani, 2007; Urban et al., 2010). Today, C_4 grasses dominate mid- to low-latitude grasslands in warm-temperate, subtropical, and tropical regions and make up ca. 20–30% of terrestrial primary production (Lloyd and Farquhar, 1994; Ehleringer et al., 1997; Fung et al., 1997; Still et al., 2003).

A dramatic increase in global C₄ biomass took place during the late Miocene, with a clear pattern of expansion earlier at lower latitudes and later in higher latitudes (Cerling et al., 1997; Ségalen et al., 2007; Passey et al., 2009). In many regions, the emergence and expansion of C₄ grasses appear to have followed a two-stage pattern: a late Miocene appearance and a post-Miocene growth to dominance (e.g. Fox and Koch, 2004; Ségalen et al., 2007; Martin et al., 2008; Strömberg and McInerney, 2011; Biasatti et al., 2012; Fox et al., 2012). C₄ grasses became a significant part of herbivore diets between 9.9 and 6.5 Ma in Eastern Africa (3°S–5°N), 7 and 3 Ma in Central Africa (~16–22°S), and 5 and 3 Ma in Southern Africa ($24-25^{\circ}$ S), but ecosystem-level open C₄ grasslands were established only in the Pleistocene (Cerling et al., 1997; Ségalen et al., 2007; Uno et al., 2011). In Argentina, C₄ plants are present in the ecosystem already at 9 Ma, and the main C₄ expansion took place 8-5 Ma (MacFadden et al., 1996; Hynek et al., 2012). In the Great Plains of North America, soil carbonate δ^{13} C records (Fox and Koch, 2003, 2004; Fox et al., 2012) indicate a presence of C₄ grass throughout Miocene with increased C₄ abundance after 6 Ma, while herbivore diets below 37°N show a significant C₄ component only after 8-6 Ma (Cerling et al., 1997; Passey et al., 2002). In South Asia, proxy records from the Siwaliks of Pakistan (Quade et al., 1989, 1992; Quade and Cerling, 1995; Cerling et al., 1997; Behrensmeyer et al., 2007; Kimura et al., 2013), India (Sanyal et al., 2004; Singh et al., 2011, 2013) and Nepal (Quade et al., 1995) show the change from C₃ to C₄ biomass taking place 8–4.5 Ma. Notwithstanding claims of up to 60% C₄ grass in the early Oligocene (Urban et al., 2010), C₄ grasses have not established a significant presence in western Europe or the eastern Mediterranean region during the past 20 Myr (Cerling et al., 1997). Isotope records from China indicate a C₄ component in the landscape from ca. 8-7 Ma, and subsequent expansion during Pliocene to Pleistocene times (Gaboardi et al., 2005; Zhang et al., 2009; Passey et al., 2009; Biasatti et al., 2012).

As of yet, the mechanisms driving the spread of C_4 grasslands are not well understood and are actively debated, but the near-synchronous expansion of C_4 grasslands suggests a combination of a global driving mechanism modified by region-specific conditions. The list of possible factors includes the long-term evolution of atmospheric partial pressure of CO_2 (e.g. Cerling et al., 1997; Zhang et al., 2009; Singh et al., 2011), tectonics (Quade et al., 1989), temperature, seasonality and aridity (Pagani et al., 1999), fire frequency (Keeley and Rundel, 2005; Scheiter et al., 2012), and in the specific case of eastern Asia, the East Asian Monsoon system (Quade et al., 1989, 1995; Passey et al., 2009).

The history of the global spread of C₄ biomass is known from the distinct isotopic signature of C₄ plants in the fossil record. The carbon fixation strategy of C₄ plants (most tropical grasses and sedges) results in reduced fractionation of carbon isotopes between atmospheric CO₂ and photosynthates. C₄ plants typically have δ^{13} C values from -15 to 10‰, significantly higher than those of C₃ vegetation (all trees, most shrubs, some cool/temperate climate grasses) ranging from -35 to -22‰. This large difference in isotopic composition allows for unambiguous detection of C₄ plants in the ecosystem, provided existence of suitable proxy materials that record the plant carbon isotopic signatures. In addition to ancient soil carbonates (e.g. Quade et al., 1989; Quade and Cerling, 1995; Latorre et al., 1997) and phytoliths (McInerney et al., 2011; Strömberg and McInerney, 2011), the carbon isotopic composition of herbivore tooth enamel has served as a window to the distant past (Cerling et al., 1993; Wang et al., 1994; MacFadden et al., 1994, 1996; Cerling et al., 1997; Passey et al., 2002; Gaboardi et al., 2005; Wang and Deng, 2005; Passey et al., 2009; Biasatti et al., 2012; Kita et al., 2014). Offset from the δ^{13} C values of the diet by 11–14‰ (Cerling et al., 1997; Cerling and Harris, 1999; Passey et al., 2005; Podlesak et al., 2008), herbivore teeth and bones reflect the isotopic composition of the consumed vegetation.

Due to sample size requirements for δ^{13} C analysis, the use of small mammal specimens for stable isotope paleoecological studies has been very limited, and investigations have relied primarily on thickenameled large mammals. Yet micromammals are common in the fossil record and make up an important component of terrestrial ecosystems, and with their generally high fecundities, high local densities, and population growth rates, they hold tremendous potential as sensitive and quickly responding recorders of environmental changes. Other than size-imposed restrictions, micromammalian isotopic proxy records show no disadvantage compared to larger animals (Gehler et al., 2012) and have proven useful in a number of pioneering studies (e.g. Grimes et al., 2005; Hopley et al., 2006; Yeakel et al., 2007; Hynek et al., 2012; Kimura et al., 2013). Hynek et al. (2012) recently demonstrated the value of micromammalian δ^{13} C values in Argentina, which enabled detection of the presence of C₄ vegetation in the ecosystem 1-2 Ma earlier than indicated by large mammal and soil carbonate δ¹³C records. The sensitivity of small mammals as recorders of their environment is linked to their physiologies. The small mass of dietary intake leads to a greater probability of sampling end-member isotope compositions, and the relatively quick growth rate of micromammalian dental tissue (Passey et al., 2005; Podlesak et al., 2008) leads to decreased dampening and homogenization of isotope signals. Thus, the possibilities of capturing the full range of isotope values present in the environment are heightened using a sampling of many small mammal taxa. The small areal ranges of individual micromammals promote an "in situ" nature of the record and thus affords potential for the reconstruction of landscape patchiness.

Taking advantage of laser-based methods (Sharp and Cerling, 1996; Passey and Cerling, 2006) allowing routine analyses of very small samples, we explore the potential of the rich micromammalian fossil faunas in four late Neogene localities across China and attempt to shed new light on the evolution of climate and spread of C₄ vegetation in eastern Asia. The abundance of micromammalian remains allows for high sample replication (in terms of numbers of individual tooth specimens analyzed) and comprehensive, representative sampling of the herbivore small mammal community. We present a data set from more than 200 small mammal tooth specimens and compare the micromammalian dietary record with that of large mammals, discussing the potential reasons for the observed differences and their implications for future isotope work on small mammals. Based on the new δ^{13} C data, we update our current knowledge - largely based on paleosol and large herbivore isotope proxies - on regional C₄ signal histories and view these histories in the larger perspective of late Neogene climatic and ecological evolution in East Asia.

2. Regional setting

Fossil material was collected from four localities (Fig. 1): two from northern China (Ertemte and Lingtai) and two from southern China (Lufeng and Yuanmou). The modern climate in all these areas is under the influence of seasonal alterations of the East Asian summer monsoon and the Siberian–Mongolian winter monsoon, and the sites in southern China are also strongly influenced by the Indian summer monsoon. During summer seasons, high pressure cells form over the Indian and Pacific oceans and give rise to the warm and humid winds migrating into the interior China. During winter seasons, the atmospheric pressure



Fig. 1. Map showing the sampled Neogene fossil localities. Additionally, the locations of five other sites discussed in the text are indicated; L = Lantian, Y = Yushe, B = Baode, BI = Bilike, G = Gaotege, Bg = Baogedawula. Dashed line indicates the approximate northwesterly extent of the present-day summer monsoon.

gradient is reversed and the cold and sometimes dust-laden winds, driven by the Siberian high pressure, migrate along the northern and eastern margins of the Tibetan Plateau to South China (An et al., 1990; An, 2000).

The Ertemte site (41°54'N 114°6'E) is located 4 km southeast of the town of Huade in the Inner Mongolian Autonomous district. The area belongs to the temperate grassland and steppe zone, and mean annual temperature and rainfall are 3.2 °C and 329 mm (WorldClim Dataset). Knowledge of the Ertemte fauna dates back to almost a century ago (Andersson, 1923; Schlosser, 1924), and it now is the most diverse and abundant Late Miocene mammalian fauna in China. The mammalian assemblage is composed of 66 species of both small and large mammals (Qiu et al., 2013b), collected from a limited stratigraphic horizon within a unit of lacustrine deposits. The age of the Ertemte fauna is based on biochronology, and it is considered to be equivalent to late MN13 (corresponds approximately to late Turolian, ca. 7-5 Ma; Storch, 1987; Qiu et al., 2006). Based on Flynn et al. (1997), Ertemte can be correlated with the basal Gaozhuang formation, which is dated magnetostratigraphically to C3r (6.033-5.235 Ma; Opdyke et al., 2013), and here we use 5.5 Ma as a rough estimate of the age of the Ertemte fauna. The Ertemte assemblage represents a typical Hipparion fauna. Though not systematically reviewed since the description by Schlosser (1924), large mammals are characterized by abundant cervids and few bovids. Micromammals are characterized by a high diversity of insectivores and myomorph rodents, and high generic diversity of ground squirrels and lophate cricetids (Qiu et al., 2013b). Our survey covers more than half of the Ertemte small mammal taxa.

The Lingtai fossil material (35° 03′N 107° 44′E) comes from a stratigraphic section in Leijiahe village, Lingtai county, in Gansu province in the central Loess Plateau (LP), ca 10 km east from Lingtai town. The area belongs to the warm-temperate deciduous-broadleaf forest zone with annual temperature and precipitation of about 10.7 °C and 620 mm (WorldClim Dataset), respectively. The late Miocene– Pliocene strata resting on the Cretaceous basement are mainly fluvial silts and sands that reach a composite thickness of ca. 60 m, overlain by Pleistocene loess and paleosol deposits. Repeated collecting activities in Leijiahe area since 1970s have resulted in numerous superposed micromammal faunas, consisting altogether of 80 micromammal forms (Zheng and Zhang, 2001; Qiu et al., 2013a), whereas the large mammal fossils collected at the site are sparse and fragmentary, dominated by herbivores like *Hipparion* and *Gazella* (Chen, 2002). Age control of the fossiliferous levels comes from magnetostratigraphy (Wei et al., 1993). The micromammalian samples in this study originate from eight different stratigraphic levels spanning 6.6–3.16 Ma in age. We have approximated the ages of the fossil beds using linear interpolation for upper and lower limits of fossil-bearing magnetozones.

The Lufeng fossil material (25° 18'N 102° 24'E) comes from near Shihuiba village in the Lufeng basin, located about 65 km northwest of the provincial capital, Kunming, Yunnan Province. The locality lies east of the Dianzhong Plateau, in a fault depression that has mean elevation of 1560 m. The area belongs to the temperate mountain forest and subtropical forest zone with the mean annual temperature of 15.2 °C and rainfall of 953 mm (WorldClim Dataset). Lufeng is renowned for the fossil hominoid Lufengpithecus lufengensis and for being the most fossiliferous and taxonomically diverse late Neogene locality in South China (Qiu and Storch, 1990). The Late Miocene exposures comprise a 20- to 30-m-thick succession (Dong and Qi, 2013) in which the fossilbearing sediments are massive lignites that are ca. 5-6 m thick, alternating with carbonaceous clays, silts, and sands with interbedded lignites (Harrison et al., 2002). Since the discovery of hominoid materials in the late 1970s, altogether 98 mammal taxa have been uncovered from the Shihuiba Formation, including three taxa of primates, and diverse ungulate and rodent fauna. Paleomagnetic ages of ca. 4 Ma (Yin, 1994) and 6.9-6.2 Ma (Dong and Qi, 2013) have been reported for the hominoid-producing horizon. However, the paleomagnetic age estimates are considered to be too young to be concordant with the mammalian fauna (cf. Flynn and Qi, 1982; Harrison et al., 2002), which compare to the Late Miocene assemblages in Europe. Furthermore, the Lufeng rhizomyd rodents are equivalent to those at 8 Ma in Siwaliks, Pakistan (Flynn and Qi, 1982). Based on these constraints, recognizing that the age is associated with large error bars, we use an approximation

of 7.5 Ma for the fossil bed for the purposes of plotting. This is also in concert with the estimated age in Biasatti et al. (2012).

The Yuanmou Basin (25°42′N 101°52′E) in Yunnan Province is located ca 110 km northwest of Kunming, on the northern edge of the Dianzhong plateau with an average elevation of about 1100 m a.s.l. (Dong and Qi, 2013). The area has a mean annual temperature of 19.03 °C and of rainfall 790 mm (WorldClim Dataset) with strong seasonal variation. The Neogene deposits (Xiaohe Formation) are mainly distributed in the northwestern part of the basin and comprise more than 80-m-thick series of fluvial and lacustrine sediments containing abundant mammalian fossils. Altogether 110 mammalian species have been recovered from the Xiaohe Fm (Dong and Qi, 2013). The Yuanmou hominoid fauna is dominated by carnivores, rodents, and artiodactyls (Qi et al., 2006). Qi et al. (2006) published a paleomagnetic age of ca. 7.2-8.1 Ma for the Xiaohe Fm. Based on micromammalian assemblages, Ni and Qiu (2002) conclude that the Yuanmou fauna is slightly older than that of Lufeng. Therefore, based on these constraints, we assign an estimated age of 8 Ma to the Yuanmou micromammal samples, consistent with the estimation presented in Biasatti et al. (2012).

3. Material and methods

The study material comprises screen washed rodent and ochotonid lagomorph teeth (n = 223; Table 1, Appendix 1). The majority of sampled teeth are adult M1 molars, although a few ochotonid incisors were also analyzed. All specimens belong to the collections of the Institute of Vertebrate Paleontology and Paleoanthtropology, Beijing, China. The tooth material from Ertemte (n = 142) and Lingtai (n = 63) is very well preserved, with yellow to light brown coloring, and clean, clear enamel surfaces. Prior to laser analysis, the teeth were washed in MilliQ water in an ultrasonic bath. The relative proportions of different taxa that were analyzed reflect the relative abundance of taxa at the localities, which are usually dominated by cricetids. The material from the Yunnan province (n = 18) is less well preserved, with variable coloring from light brown to black and the enamel surfaces often partially covered by sediment. In addition, some surfaces showed remains of adhesive putty. The Yunnan material was treated with trichloroethylene solution and MilliQ water under ultrasonic agitation to remove traces of putty and sediment.

The stable isotope analyses were carried out at Johns Hopkins University using a Photon-Machines Fusions 30-watt CO₂ laser coupled to a custom built partially automated extraction line following the general design of Passey and Cerling (2006). The isotope ratios of the CO₂ extracted from the tooth enamel were measured on a Thermo MAT 253 mass spectrometer. The analytical procedures and data reduction protocols follow those outlined in Passey and Cerling (2006). The δ^{13} C and δ^{18} O values reported for each specimen represent the mean of one to eight individual analyses. After data reduction, a correction of +0.5%was applied to the δ^{13} C values to account for the systematic offset observed between laser-based δ^{13} C values and those derived from the conventional phosphoric acid method (Passey and Cerling, 2006). Estimates of the relative proportions of C₃ and C₄ plants ingested (reported as percent C_4) follow the approach of Passey et al. (2002), where the δ^{13} C values of C₃ plants, water-stressed C₃ plants, and C₄ plants are calculated from temporally varying δ^{13} C values of atmospheric CO₂. To account for the isotopic fractionation between enamel and diet, an enrichment factor of 11.0% is used (c.f. Bywater-Reyes et al., 2010; Hynek et al., 2012). Whether the chosen enrichment factor is appropriate for all small bodied mammals remains untested, but recent findings of a smaller enamel-diet fractionation for wood rats $(11.0\% \pm 0.1;$ Podlesak et al., 2008) and voles (11.5 $\% \pm 0.3$; Passey et al., 2005) relative to large mammals (ca. 14‰; Cerling and Harris, 1999) add to the growing pool of evidence of smaller enrichment factors for small mammals (ca. 9-10‰ for laboratory rats and mice: DeNiro and Epstein, 1978; Ambrose and Norr, 1993; Tieszen and Fagre, 1993; Jim et al., 2004).

Table 1

Sample material, number of tooth specimens analyzed per taxon, range of enamel δ^{13} C values, and calculated contribution of C₄ grasses to diet.

		n	δ^{13} C range ^a		%С4 ^b
			max	min	
FRTEMTE INNER MONGOLIA					
Cricetidae	Sinocricetus zdanskvi	10	-9.5	-10.9	7-18
	Sinocricetus sp.	10	-5.9	-11.5	1-49
	Microtodon atavus	8	-10.6	-12.6	0-9
	Pseudomeriones	7	-4.5	-11.1	5-60
	abbreviatus				
	Kowalskia neimengensis	5	-10.3	-11.8	0-11
	Microtoscoptes	5	-11.2	-13.1	0-4
	praetermissus				
	Nannocricetus sp.	5	-8.5	-11.0	5-26
	Anatolomys teilhardi	2	-11.2	-11.3	3
Cricetidae	Prosiphneus licenti	5	-10.9	-12.2	0-6
(Myospalacinae)	-				
Dipodidae	Dipus fraudator	9	-7.5	-10.7	8-35
-	Lophocricetus sp.	8	-8.4	-11.0	1-27
	Paralactaga suni	4	-9.0	-11.3	3-18
	Prolophocricetus sp.	3	-9.3	-12.6	0-20
	Sicista sp.	1	-11.2		4
	Sinozapus sp.	1	-11.3		3
Muridae	Occitanomys pusillus	15	-5.7	-12.4	0-50
	Apodemus orientalis	7	-5.1	-12.3	0-55
	Micromys chalceus	5	-11.5	-13.4	0-1
	Karnimata hipparionum	4	-10.2	-13.2	0-12
	Orientalomys cf. similis	3	-10.9	-12.1	0-7
Ochotonidae	Ochotona lagreli	8	-9.1	-10.1	13-21
	Ochotona minor	5	-10.1	-10.7	8-13
Sciuridae	Eutamias ertemtensis	8	-10.0	-12.1	0-14
	Sinotamias sp.	1	-12.8		0
	Prospermophilus sp.	1	-10.4		11
Gliridae	Myomimus sinensis	2	-11.1	-11.7	0–5
LINGTAL GANSU PROVINCE					
Cricetidae	Pseudomeriones	15	-3.4	-14.9	0-73
	abbreviatus				
	Bahomys sp.	7	-8.7	-13.3	0-25
	Nannocricetus sp.	4	-6.0	-10.2	13-48
	Allocricetus bursae	4	-8.6	-11.1	5-27
	Chardinomys primitivus	3	-8.7	-12.7	0-26
Cricetidae	Mesosiphneus intermedius	4	-11.1	-11.8	0-5
(Myospalacinae)					
	Mesosiphneus teilhardi	3	-10.5	-11.2	5-10
	Prosiphneus licenti	3	-8.4	-9.1	19-28
Ochotonidae	Ochotona sp.	14	-7.6	-11.9	0-35
	Ochotonoides	4	-8.1	-14.2	0-30
	complicidens				
Muridae	Karnimata hipparionum	3	-9.0	-13.1	0-23
Platacanthomyidae	Platacanthomus dianonsis	2	_11 =	_17 <i>6</i>	0_1
i ididedititi0111y10de	Tunhlomus primitiuus	∠ 2	120	- 12.0	0-1
Muridae	Linomys vunnanansis	ر	- 12.0	- 13.3	0_7
withIude	Linomys yunnunensis	4	- 10.8	- 14.0	0-7
LUFENG, YUNNAN PROVINCE					
Muridae	Linomys yunnanensis	6	-8.9	-15.6	0-23
Cricetidae	Kowalskia hanae	3	-13.5	-14.5	0

The full data, including ages of sampled material, are given in the Appendix.

 $^a\,$ Values have been adjusted by +0.5% to account for the mean offset between laser-based and conventional phosphoric acid digestion-based $\delta^{13}C$ values (see Material and Methods).

^b Proportions of C_4 vegetation are calculated following Passey et al. (2002) (see Material and Methods).

The δ^{18} O values obtained by laser ablation represent a mixture of oxygen from the phosphate, carbonate, and hydroxide components of the hydroxyapatite. The isotopic offset between laser and conventional phosphoric acid digestion δ^{18} O values is systematic in direction but variable in magnitude, averaging -5.1% with a standard deviation of 1.2‰ for multiple laser-conventional comparisons reported by Passey and Cerling (2006). Due to the uncertainty of the isotopic offset, we do not attempt to reconstruct paleo-water δ^{18} O values using laser-based enamel δ^{18} O values. The rodent enamel δ^{18} O data are included in

Fig. 2 and in Appendix 1, but they are not interpreted nor discussed further in this paper.

A limitation of the laser ablation approach is that the laser will sample any material it impinges upon, including organic compounds on or in the tooth enamel (e.g., glues and cements used by curators, sedimentary organic carbon, and residual structural organics in the biomineral itself). Laser ablation of organic compounds in the helium-purged, oxygen-free atmosphere of the sample chamber results in pyrolization, visible as charring and production of smoke and other lingering vapors (Passey and Cerling, 2006). Any oxidation to CO₂ of organic carbon would be expected to lower our observed δ^{13} C values, because organic carbon typically has lower δ^{13} C values than coexisting inorganic carbon. In our study, data from all analyses associated with heavy charring and smoke production were discarded (n = 21). Some analyses resulted in production of incipient char halos or slight charring of the laser pit bottom, indicating minor contamination with organic carbon. Although Passey (2007) reports no significant difference in δ^{13} C between "minor char" and "clean" analyses, we observe a small but statistically significant difference (paired t-test; p < 0.05) in mean δ^{13} C values between minor char $(-10.8\% \pm 1.8)$ and clean $(-10.6\% \pm 1.9)$ analyses of the same tooth specimens (n = 58). However, this difference is inconsequential compared to the difference in δ^{13} C between C₃ and C₄ vegetation (ca. 15‰). Thus, because our objective is differentiating between C_3 and C_4 plant consumption, we apply no correction to the minor char analyses and include these data in the overall data set.

4. Results

The isotopic data are plotted in Figs. 2 and 3 and summarized in Table 1. The full dataset of δ^{13} C values is reported in Appendix 1. Except for the smallest specimens where molar size limited the number of analyses to one (n = 17), the reported values represent the mean of two to eight replicate analyses. The average standard deviation of sample replicates was $0.3 \pm 0.3\%$ (1 σ) and $0.2 \pm 0.3\%$ for δ^{13} C and δ^{18} O, respectively. Appendix 1 includes also δ^{13} C and δ^{18} O values of large herbivore enamel from Yunnan, previously presented only in a thesis (Passey, 2007).

The δ^{13} C values of tooth enamel fall between -15.6 and -3.4%, with a total range of 12.2‰. The majority of δ^{13} C data fall below -8%; only 14 specimens show less negative δ^{13} C values (Fig. 2).



Fig. 2. δ^{13} C and δ^{18} O values of rodent tooth enamel. Each symbol represents the mean value of one to four analyses of an individual tooth. Error bars are excluded for clarity but are generally less than 1% (1 σ) for both δ^{13} C and δ^{18} O. δ^{13} C values are adjusted by + 0.5% to account for the mean offset between laser-based and conventional phosphoric acid digestion-based δ^{13} C values (see Material and Methods). Proportions of consumed C₄ vegetation were calculated following the approach of Passey et al. (2002), using an enamel-diet fractionation of 11.0%.



Fig. 3. Records of δ^{13} C of vegetation in the diets of fossil mammals from China–Ertemte and Inner Mongolia (A), Lingtai (B), and Yunnan (C). Large mammal data for Inner Mongolia from Zhang et al. (2009) and for Yunnan from Passey (2007) and Biasatti et al. (2012). Soil carbonate data for Lingtai from Ding and Yang (2000). The dotted lines indicate model-predicted (Passey et al., 2009) δ^{13} C values of average C₃ vegetation, waterstressed plants, and average C₄ plants. The late Miocene large mammal data from Biasatti et al. (2012) and Passey (2007) are plotted according to the age estimates given in the text (Section 2 Regional Setting) instead of those used in the original publications.

Specimens from Ertemte and Lingtai show significantly overlapping ranges and comparable mean values (t-test >0.05), with δ^{13} C values from -13.1 to -4.5% and -14.9 to -3.4, respectively. The data from Yuanmou and Lufeng in the Yunnan province plot to the lower end of the total δ^{13} C range, from -15.6 to -8.9%. They are statistically indistinguishable from each other but differ significantly (t-test p < 0.001) from the datasets of Lingtai and Ertemte.

The calculated proportions of C_4 plants ($%C_4$) in the diets of the small mammals are listed in Table 1 and illustrated in Fig. 3. Apart from one murid from Lufeng with an estimate of 23% C₄, the Yunnan rodents show C₃-based diets at ~8–7 Ma. In Lingtai and Ertemte, various levels of C₄ intake are observed. In Lingtai, C₄ contribution to diet varies from 0 to 73%, with estimates of more than 30% for only 5 out of 63 specimens. The gerbil species *Pseudomeriones abbreviatus* (n = 15), sampled from stratigraphic levels with ages 6.45 Ma, 5.8 Ma, 4.8 Ma and 4.7 Ma, shows an increasing trend in the amount of C₄ grass consumed from 6.45 Ma (0%) to 4.7 Ma (up to 73%) (Appendix). At Ertemte, C_4 consumption ranges from 0 to 60%, and only 10 out of 142 specimens indicate more than 30% C₄ intake. At both Lingtai and Ertemte, all taxa reaching a C₄ grass contribution of ca. 50% or above belong to muroids (the families Cricetidae and Muridae). Individuals in the families Ochotonidae and Dipodidae also consumed significant (\geq 30%) amounts of C₄ vegetation.

5.1. Lingtai

In Lingtai, the mean δ^{13} C values of micromammalian enamel at the sampled stratigraphic levels increase from -11.3% at 6.45–6.6 Ma to - 10.0‰ at 3.16 Ma, but this apparent change is not statistically significant. Due to the scarcity of large vertebrate fossil material in the late Miocene to Pliocene strata, prior δ^{13} C data on teeth and bones do not exist from Lingtai. However, the small mammal isotope data may be compared with a soil carbonate δ^{13} C record (Fig. 3b) from the same time period (Ding and Yang, 2000) taking into account a ca. 3% larger offset from source plants for pedogenic carbonate compared to small mammal enamel ($\Delta_{carbonate-biomass}$ ~13.5‰ at 25 °C; Cerling and Quade, 1993). Prior to 4 Ma, the soil carbonate data are interpreted to reflect a C₃-dominated ecosystem (Ding and Yang, 2000), with plant biomass δ^{13} C values of -22 to -23%. C₄ plants were, however, present at Lingtai, perhaps as spatially very limited patches, as registered by the few higher small mammal enamel δ^{13} C values at 6.6, 5.8, and 4.7 Ma. The particularly high δ^{13} C value of one specimen at 4.7 Ma illustrates this patchiness, and yet again documents the increased affinity of micromammals to register end-member isotope compositions.

Even after 4 Ma, when the soil carbonate δ^{13} C values begin to rise, C₄ plant abundance in the landscape seems to have remained limited (Ding and Yang, 2000). A single soil carbonate peak value of -4.7% at 3.4 Ma corresponds to mean plant δ^{13} C values of -19%, suggesting a maximum of ~30% C₄ biomass. The mean values of small mammal enamel match the soil carbonate record. For the sampling levels at 3.46 Ma, 3.35 Ma, and 3.16 Ma, mean micromammal-inferred plant biomass δ^{13} C values are -21.8, -21.1, and -21.0%, while the soil carbonate data indicate plant δ^{13} C values of -21.5, 20, and -21%, respectively. Translated into percent C₄ vegetation and averaged, both records suggest ~15% contribution from C₄ vegetation over the 3–3.5 Ma interval, indicating a mixed but still a clearly C₃-dominated ecosystem prevailed on the Loess Plateau.

5.2. Inner Mongolia

The micromammalian enamel δ^{13} C values from Ertemte indicate the presence of C₄ grasses in the local ecosystem at ~5.5 Ma. The spatial and temporal patterns of C₄ vegetation in Central Inner Mongolia were previously studied by Zhang et al. (2009) based on enamel δ^{13} C values of various large herbivores. They infer a first appearance of a significant C_4 component in the landscape in the 7.5 Ma Baogedawula locality ca 200 km NE of Ertemte (Fig. 3a), where the C₄ component of herbivore diets ranged from 0 to 29%. There are no large herbivore data from the intervening time period from that region, but again at 3.9 Ma horses and elephantids from the Gaotege locality (G in Fig. 1) were estimated to have incorporated 19-53% C₄ plants in their diets. For Ertemte (~5.5 Ma) and the nearby locality of Bilike (4.7 Ma), the large mammal enamel δ^{13} C data indicate a pure C₃ diet. The carbon isotopic composition of diet for the large herbivores, -27 to -24%, is considerably lower than that inferred for the micromammals, -24.4 to -15.5%. Making a note of possible inadequacy in number of samples (n = 6), they hypothesize that the pure C₃ diet of large herbivores reflects a retreat of C₄ vegetation after an initial expansion at 7.5 Ma (inferred from the Baogedawula area), or that the Ertemte area remained forested without any C₄ plants in the latest Miocene and Pliocene due to a warmer and wetter climate.

The small and large mammal δ^{13} C data are not necessarily in conflict but may be explained by their inherent characteristics. Contrary to micromammalian isotope records, δ^{13} C values of large herbivores do not necessarily represent a strictly local vegetation signal (e.g. Zazzo et al., 2010). Differences may thus reflect larger range areas or migratory behavior of large herbivores. Also, as discussed in the Introduction, micromammalian δ^{13} C records have the potential to detect smaller abundances of C_4 plants in the ecosystem due to the smaller biomass consumed and shorter temporal timespan over which the signal is averaged. Thus, it is possible that the proportion of C_4 grasses in Ertemte was small enough not to show up in large herbivore diets, suggesting low to moderate levels of C_4 vegetation in the landscape. According to a survey of local plants, C_4 grasses constitute a minor component of the presentday ecosystem in the area (Zhang et al., 2009).

5.3. Yunnan

The diets of the rodents from the Yunnan province are more ¹³Cdepleted (mean – 24‰) than those of small mammals from the northern and central parts of China (mean – 21.4‰), and with a single exception are consistent with pure C₃ vegetation. This suggests a strongly C₃-dominated environment, in agreement with other lines of evidence. For example, based on small mammal assemblages, the paleoenvironmental context at Yuanmou and Lufeng has been interpreted as a tropical forest with scattered shrub and grasslands (Ni and Qiu, 2002; Qi et al., 2006). In a habitat-type analysis of the micromammalian communities, Ni and Qiu (2002) show that over 80% of the small mammal taxa were restricted to or preferred mesic/forested habitats, while only ~3–5% lived in bush and grasslands.

The predominantly closed, forested biomes of Yunnan are also reflected as low δ^{13} C values of large herbivore taxa (Passey, 2007; Biasatti et al., 2012). Compared to ungulates from contemporaneous fossil localities on the Loess Plateau, those at Yuanmou displayed markedly lower enamel δ^{13} C values, indicative of a diet consisting solely of C₃ vegetation (Passey, 2007). This agrees with the findings of Biasatti et al. (2012), who inferred nearly pure C₃ diets for large herbivores prior to 2–3 Ma at six localities in the Yunnan province. However, C₄ grasses were evidently present in the landscape, most likely in scattered patches of more open habitat. These microhabitats left an imprint on the δ^{13} C values of paleosol carbonates and large mammal enamel (Biasatti et al., 2012) and could have served as the source of the C₄ signal observed here for a single Lufeng murid.

5.4. C₄ expansion and climate

The spatiotemporal systematics of the spread of C₄ grasses is intrinsically linked to the climatic evolution of a region. Accumulating evidence from paleosol carbonate δ^{13} C values from several localities in northern China (Ding and Yang, 2000; Jiang et al., 2002; An et al., 2005; Kaakinen et al., 2006; Passey et al., 2009; Suarez et al., 2011) indicates the presence of a spatial climatic gradient defined by northward increasing δ^{13} C values and, thus, abundance of C₄ grasses, in the late Miocene and Pliocene (~7–2.7 Ma; Passey et al., 2009; Suarez et al., 2011) as a contrast to the late Pleistocene and present-day spatial pattern of southward increasing abundance of C₄ grass on the loess Plateau.

This reversed late Miocene to Pliocene pattern finds further support from large herbivore enamel δ^{13} C records across China (Passey et al., 2009). The following is a brief summary of the large herbivore δ^{13} C records for ~8-4 Ma (the period of good spatial data coverage), with the ages in the parentheses representing the total age span of the fossil specimens at each site. The maximum C₄ abundance is recorded in Yushe (6.3-2 Ma), east of the Loess Plateau (see Fig. 1), where herbivore diets commonly consisted of 50% C₄ forage (Passey, 2007; Passey et al., 2009). To the north, Baode (7-5.5 Ma) on the northern Loess Plateau and Baogedawula (7.5 Ma), Ertemte (5.5 Ma), Bilike (4.7 Ma), and Gaotege (3.9 Ma) in Inner Mongolia are C3 dominated, but animal diets at Baode, Baogedawula, and Gaotege show moderate levels of C₄ grass (Passey, 2007; Passey et al., 2009; Zhang et al., 2009; Kaakinen et al., 2013; Eronen et al., 2014). For the 7.5-4.7 Ma ungulates from Baode and Baogedawula, C₄ grasses contributed 0–30% to their diet, while the 3.9 Ma large mammal diets in Gaotege show a C₄ component of up to 50%. To the south of the C₄ maximum zone at Yushe, the late Miocene localities at Lantian (9.95-6.6 Ma) in the central Loess Plateau

also indicate C₃-based ecosystems with little C₄ vegetation (Passey et al., 2009). Only 6 out of 55 ungulate individuals incorporated up to 10% of C₄ in their diets. Passey et al. (2009) suggest that Lantian might have been a relatively forested milieu during the late Miocene. Even further south in Yunnan, large herbivores at Yuanmou, Lufeng, Baoshan, and Zhaotong (8.5–1.75 Ma, 7.5 Ma, 5 Ma–9 ka, 4 Ma, respectively; Biasatti et al., 2012; Passey, 2007) show little to no evidence of C₄ grass in their diets. However, from 3 Ma, C₄ vegetation became a significant part in the diets of herbivores in Yunnan (Biasatti et al., 2012), indicating a substantial change in regional environmental parameters taking place between 3.5 and 2.5 Ma. Incidentally, a contemporaneous change in climatic regime is also observed on the Loess Plateau and Inner Mongolia. In Lingtai and Lantian, the δ^{13} C sequences of pedogenic carbonate display a peak between 3.4 and 2.7 Ma (Ding and Yang, 2000;

An et al., 2005; Kaakinen et al., 2006; Suarez et al., 2011) suggesting transient improved conditions for C₄ growth during that time. In Inner Mongolia moderate levels of C₄ grasses are still observed at 3.9 Ma in Gaotege, while presently, they account for an insignificant amount of the local biomass (Zhang et al., 2009) suggesting a retreat or at least a significant reduction of C₄ vegetation in the area.

The micromammalian δ^{13} C data, suggesting C₃-based environments with some occurrence of C₄ grasses at both Lingtai and Ertemte, and pure C₃ vegetation for Yunnan, are consistent with the spatiotemporal pattern emerging from the soil carbonate and large mammal δ^{13} C records. According to this pattern, a zone of optimum C₄ growth was located somewhere south of Baode and the Inner Mongolian localities at 8–4 Ma, and C₄ abundance decreases both to the north and south of this maximum zone (Passey et al., 2009).



Fig. 4. Recorded occurrences of the "high-C₄" genera Apodemus, Occitanomys, and Pseudomeriones for 11.2–8.2 Ma (black dots) and 8.2–5.3 Ma (white dots) in the NOW database (Fortelius, 2014).

This pattern and the environmental changes reflected in the soil carbonate records and Yunnan herbivore diets at 3.5-2.5 Ma can be explained by a conceptual model proposed by Passey et al. (2009). In this scenario, the environmental gradient maintained by the East Asian Monsoon system, characterized by C₃ forests in the southeast changing to $C_3 + C_4$ steppe, and finally to the increasingly arid and cool C₃ + C₄ biomes in the northwest, was shifted toward the north during a period of strengthened East Asian summer monsoon. A subsequent weakening of the moisture-laden summer monsoons during 3.5-2.5 Ma caused a southward retreat of the forest-steppe-desert transition, moving the C₄ maximum steppe zone further away from the northern localities in Inner Mongolia, through the central and southern Loess Plateau localities Lingtai and Lantian. This shift might have been related to the establishment of the present state of focused summer rains with little spring, autumn, and winter precipitation (Suarez et al., 2011).

5.5. Late Miocene immigrants to China

The hypothesis of Passey et al. (2009) assumes active operation of the summer monsoon system during late Miocene to early Pliocene in East Asia, and it finds support from several other lines of evidence pointing toward humid and warm conditions during this time (e.g. Rea et al., 1998; Ding et al., 1999; Fortelius et al., 2002; Jia et al., 2003; Ma et al., 2005; Fortelius and Zhang, 2006; Wu et al., 2006; Wang et al., 2006; Kaakinen et al., 2006; Rao et al., 2008; Sun et al., 2010; Xing et al., 2012). The Late Miocene humidity of Northern China is in stark contrast to the global trend of mid-latitude drying at that time (Fortelius et al., 2002), and it seems to have created a favorable setting for the influx of immigrant herbivorous land mammals from several directions, and the later development of endemic taxa from both native and immigrant origin (Fortelius and Zhang, 2006; Mirzaie Ataabadi et al., 2013).

Similar to large mammals, rodents show a pattern of endemism and immigrations during the later late Miocene, the Baodean Land Mammal Age (Fortelius and Zhang, 2006). According to recorded occurrences in the New and Old Worlds Database of Fossil Mammals (NOW database; dataset downloaded 13th November 2013; Fortelius, 2014), three of the 32 small mammal genera included in our analyses made their first appearance in China in the late Miocene, later than their first occurrences in the West. They are Apodemus, Occitanomys, and Pseudomeriones, incidentally the same three genera whose members were observed to incorporate >50% C₄ in their dietary intake (Table 1). A fourth "high-C₄" cricetid, Sinocricetus, is endemic to East Asia. The distributions of the three "high-C₄" taxa at time periods 11.2–8.2 Ma and 8.2–5.3 Ma are illustrated in Fig. 4, which shows that after 8.2 Ma, these genera, especially Occitanomys and Apodemus, expanded considerably in Europe and extended their distribution to the Far East. The significance of this pattern is hard to assess on its own, since the small mammal record of the earlier late Miocene is relatively sparse in China. However, a similar pattern of apparent eastwards dispersal was observed for large mammals by Mirzaie Ataabadi et al. (2013) though in this case the pattern could be anchored in dental ecometrics suggesting that the dispersal was driven by increased humidity. The eastwards dispersal of these small mammal taxa might well be part of the same continental-scale biogeographic process.

If the pattern is not an artifact of a temporal sampling bias, it is inviting to hypothesize that the onset of strong summer monsoons and the establishment of C_4 ecosystems likely opened new ecological opportunities for these "high- C_4 " taxa.

6. Conclusions

The δ^{13} C data from Late Miocene and Pliocene small mammal teeth from China are consistent with prior Chinese soil carbonate and large mammal isotope data establishing a scenario of northward increasing

 C_4 abundance that in turn mirrors the climatic gradient set up by the East Asian monsoons. The carbon isotope data suggest an essentially pure C_3 environment for Yunnan around 7.5 Ma, in agreement with other lines of evidence indicating dense, forested biomes. For Lingtai, on the Loess Plateau, the small mammal $\delta^{13}C$ values indicate the presence of C_4 vegetation amidst a clearly C_3 -dominant landscape between 6.6 and 3.16 Ma. The small mammal record from Ertemte detects the presence of a C_4 component where large mammal diet data indicate a pure C_3 environment. In Lingtai and Ertemte, 7–8% of the small mammals incorporated more than 30% C_4 in their diets. The results draw attention to differences in the nature of $\delta^{13}C$ records generated from small versus large mammals and suggest that small mammals are ideal for detecting the first, faint signals of C_4 emergence in ecosystems.

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Appendix A. Supplementary data

The Appendix is an Excel-file containing the full data set of lasergenerated δ^{13} C and δ^{18} O values of small mammal tooth enamel, and δ^{13} C and δ^{18} O data of large herbivore tooth enamel from Yunnan by Passey (2007) discussed in Sections 5.3 and 5.4. Supplementary data associated with this article can be found in the online version, at [doi:http://dx.doi.org/10.1016/j.gloplacha.2015.09.003].

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