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Oxygen and carbon isotope compositions of middle Cretaceous vertebrates from North Africa and Brazil: Ecological and environmental significance

Romain Amiot ^{a,b,*}, Xu Wang ^c, Christophe Lécuyer ^b, Eric Buffetaut ^d, Larbi Boudad ^e, Lionel Cavin ^f, Zhongli Ding ^c, Frédéric Fluteau ^g, Alexander W.A. Kellner ^h, Haiyan Tong ^d, Fusong Zhang ^c

^a Key Laboratory of Evolutionary Systematics of Vertebrates, Institute of Vertebrate Paleontology and Paleoanthropology, CAS, 142 Xi Zhi Men Wai Da Jie, Beijing 100044, China

^b CNRS UMR 5125, Paléoenvironnements & Paléobiosphère, Université Lyon 1, 69622, Villeurbanne, France

^c Key Laboratory of Cenozoic Geology and Environment, Institute of Geology and Geophysics, Chinese Academy of Sciences, Beijing 100029, China

^d CNRS UMR 8538, Laboratoire de Géologie de l'Ecole Normale Supérieure, 24 rue Lhomond, 75231 Paris Cedex 05, France

^e Université Moulay Ismail, Laboratoire des Formations Superficielles, BP.509 Boutalamine, 52000 Errachidia, Morocco

^f Department of Geology and Palaeontology, Muséum d'Histoire naturelle, CP 6434, 1211, Genève 6, Switzerland

^g Institut de Physique du Globe de Paris, 2 place Jussieu, F-75005, Paris, France

h Paleovertebrate Sector, Department of Geology and Paleontology, Museu Nacional/Universidade Federal do Rio de Janeiro, Quinta da Boa Vista, São Cristóvão,20940-040 Rio de Janeiro, RJ, Brazil

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ABSTRACT

In order to investigate mid-Cretaceous terrestrial climates of low paleolatitudes, Moroccan, Tunisian and Brazilian vertebrate apatites have been analyzed for their oxygen and carbon isotope compositions of phosphates ($\delta^{18}O_p$) and carbonates ($\delta^{18}O_c$, $\delta^{13}C_c$). At each site, coexisting theropod dinosaurs, titanosaurid sauropods, pterosaurs, crocodilians, turtles and fish have distinct $\delta^{18}O_p$ and $\delta^{13}C_c$ values reflecting their ecologies, diets and foraging environments. Oxygen isotope compositions of surface waters ($\delta^{18}O_w$) estimated from turtle and crocodile $\delta^{18}O_p$ values range from $-5.0 \pm 1.0\%$ to $-2.4 \pm 1.0\%$, which do not differ from mean annual rainwater values occurring today under inter-tropical sub-arid to arid climates. High water temperatures ranging from 21 ± 6 °C to 34 ± 2 °C deduced from fish $\delta^{18}O_p$ values are in agreement with those published for mid-Cretaceous low latitudes. Temporary or seasonal droughts are inferred from high $\delta^{18}O_p$ values of lungfish teeth, even though lower reptile $\delta^{18}O_p$ values suggest the use of distinct and most likely larger or regularly renewed bodies of water. Environmental conditions of the studied low latitude regions during the Aptian–Cenomanian interval were somewhat similar to those experienced today under semi-arid to arid tropical or equatorial climates, but with higher mean surface temperatures than present–day ones. © 2010 Elsevier B.V. All rights reserved.

1. Introduction

During the Cretaceous period (-135 to -65 Ma), important thermal fluctuations have been identified with an optimum during the mid-Cretaceous (Albian–Turonian), and cold episodes characterized by near freezing to freezing conditions in polar regions (e.g. Frakes, 1999; Price, 1999; Pucéat et al., 2003; Amiot et al., 2004). Elevated partial pressure of atmospheric CO₂, changes in oceanic circulation patterns in relation to plate motion and high sea levels most likely accounted for these peculiar climatic conditions (Haq et al., 1988; Barron et al., 1995; Berner and Kothavala, 2001). During the mid-Cretaceous thermal optimum, low equator-to-pole thermal gradient

* Corresponding author. CNRS UMR 5125, Paléoenvironnements & Paléobiosphère, Université Lyon 1, 69622, Villeurbanne, France.

E-mail address: romain.amiot@univ-lyon1.fr (R. Amiot).

(Huber et al., 1995), very high temperatures at low latitudes (Schouten et al., 2003) and episodically extreme climatic conditions in Saharan environments have been proposed (Russell and Paesler, 2003). According to these authors, mid-Cretaceous Saharan environments were characterized by a decline in productivity of terrestrial vegetation inferred from the relatively small size of herbivorous dinosaurs, highly productive waters deduced from the large size of fish and crocodilians, apparently a low vertebrate diversity and episodic climatic extremes taking mid-Cretaceous Saharan ecosystems near tolerance limits. If mid-Cretaceous Saharan ecosystems suffered such conditions, it raises the question of how the rather diverse and complex faunal associations shared water and food resources. Therefore, we analyzed the oxygen and carbon isotope compositions of Aptian to Cenomanian freshwater reptiles and fish from two North African and two Brazilian deposits in order to investigate climatic conditions as well as the habitats and foraging preferences of vertebrate faunas.

The first step consists in estimating mean air temperatures where dinosaurs, non-dinosaurian reptiles and fish coexisted. According to the studies performed by Barrick et al. (1999) and Amiot et al. (2007),

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the oxygen isotope composition of ambient waters can be inferred from that of turtle and crocodile apatite phosphate, which in turn can be used to calculate water temperature when combined to the isotopic composition of coexisting fish phosphate according to fractionation equations determined by Longinelli and Nuti (1973) and Kolodny et al. (1983). These water temperatures can be further compared to air temperatures estimated from the oxygen isotope compositions of ambient freshwaters that indirectly reflect the compositions of terrestrial vertebrates for feeding on natural resources can be approached by the use of the carbon isotope composition of apatite carbonate that reflects the isotopic composition of food (e.g. Sillen and Lee-Thorp, 1994; Koch, 1998; Cerling and Harris, 1999; Passey et al., 2005 ; Fricke and Pearson, 2008; Tütken, in press).

1.1. Studied areas

Four deposits from inter-tropical paleolatitudes are considered (Fig. 1). Most samples were recovered from Moroccan deposits belonging to the lower Kem Kem beds (or Ifezouane Formation) in the Tafilalt region where extensive field exploration and collecting were carried out. This Formation, dated as Lower Cenomanian, represents fluvial deposits in deltaic environments (see Cavin et al. (2010) for a complete review). The other North African studied deposit is the Chenini Member of the Aïn Guettar Formation of Tunisia from which oxygen isotope compositions of phosphatic remains have been published (Amiot et al., 2006). The Chenini Member, dated as Upper Aptian to Lower Albian (Le Loeuff et al., 2010), also represents fluviodeltaic environments with possibly mixed freshwater and marine influences (Anderson et al., 2007). Two sites were studied in Brazil, which are the locality of Laje do Coringa, constituted by estuarine deposits belonging to the Alcântara Formation of Lower Cenomanian age (Pedrão et al., 1993; Kellner et al., 2009), and the Araripe Basin from which a few samples come from the Crato and Romualdo formations, dated as Aptian to Albian (Pons et al., 1991; Fara et al., 2005).

1.2. Oxygen isotope composition of vertebrate apatites

The oxygen isotope compositions of apatite phosphate ($\delta^{18}O_p$) and carbonate ($\delta^{18}O_c$) from vertebrate bones, teeth and fish scales are a function of the $\delta^{18}O_{bw}$ value of the animal's body water as well as of its body temperature (Kolodny et al., 1983; Longinelli, 1984; Luz et al., 1984). The $\delta^{18}O_{bw}$ value is related to the $\delta^{18}O_w$ value of ingested water and to the animals' ecology and physiology. For most continental vertebrates, the main source of ingested oxygen is drinking or plant

water, which is meteoric water or water derived from it (D'Angela and Longinelli, 1990; Cormie et al., 1994; Kohn et al., 1996; Straight et al., 2004). As the $\delta^{18}O_w$ value of meteoric water depends on climatic parameters such as air temperature, humidity and amount of precipitation (Dansgaard, 1964; Gat, 1996; Grafenstein et al., 1996; Fricke and O'Neil, 1999), vertebrates thus indirectly record in their phosphatic tissues the climatic conditions of their living environment. It is noteworthy that the $\delta^{18} O_w$ value of surface waters can differ from that of precipitations due to local processes such as evaporation, mixing with ground waters or with rivers draining catchments with different water oxygen isotope compositions. Such processes complicate the interpretations in terms of climatic reconstructions. Physiological adaptations to a specific habitat (aquatic, semi-aquatic or terrestrial) also affect the $\delta^{18}O_{bw}$ value by controlling the magnitude of body input and output oxygen fluxes, some of them being associated with oxygen isotopic fractionations between apatite and water (Luz and Kolodny, 1985; Bryant and Froelich, 1995; Kohn, 1996). From living and fossil communities of mammals, reptiles and fish, it has been observed that differences in the range of $\delta^{18}O_{p}$ values or in some cases differences in mean $\delta^{18}O_p$ values between coexisting aquatic or semi-aquatic vertebrates and terrestrial ones were related to their habitat use (Fricke and Rogers, 2000; Clementz and Koch, 2001; Clementz et al., 2003; Amiot et al., 2006, 2010). Ecological specificities such as plant-water use among herbivorous communities also affect the $\delta^{18} O_p$ value of vertebrates. Indeed, large differences in $\delta^{18}O_p$ values have been observed between coexisting herbivorous mammals that drink surface waters and those that only rely on water in plants, usually enriched in ¹⁸O by several per mils relative to surface waters (Kohn et al., 1996). The $\delta^{18}O_w$ values of ingested water can be estimated from $\delta^{18}O_p$ values of fish, turtles and crocodilians by using PO₄-water fractionation equations established for extant species (e.g. Kolodny et al., 1983; Barrick et al., 1999; Amiot et al., 2007), the applicability of these equations to Mesozoic faunas having been tested successfully (Barrick et al., 1999; Amiot et al., 2004, 2009).

1.3. Carbon isotope composition of vertebrate apatites

Carbon isotope compositions of apatite from air-breathing vertebrates primarily reflect animal diets, with a ¹³C-enrichment relative to ¹²C that varies among animals. Indeed, ¹³C-enrichments between apatite and diet of herbivorous and carnivorous terrestrial mammals were estimated in the range 10–15‰ and close to 9‰, respectively (Lee-Thorp et al., 1989; Tieszen and Fagre, 1993; Sillen and Lee-Thorp, 1994; Koch, 1998; Cerling and Harris, 1999; Passey et al., 2005). Similar ¹³C-enrichments have been observed between reptile apatite and diet, namely 8–9‰ and 12‰ for carnivorous and herbivorous



Fig. 1. Paleogeographic map of the mid-Cretaceous showing the geographic position of studied localities. Numbers refer to the following deposits: 1: Ifezouane Formation, Morocco; 2: Chenini Member, Tunisia; 3: Crato and Romualdo formations, Brazil; 4: Alcântara Formation, Brazil.

marine turtles, respectively, and about 9% estimated for crocodilians (Stanton, 2006). Higher ¹³C-enrichment of 16‰ has been determined for large non flying birds (Von Schirnding et al., 1982; Johnson et al., 1998). According to some studies, variations in ¹³C-enrichment between apatite and diet are primarily a result of differences in digestive physiology, rather than differences in the magnitude of fractionation between mineral and body fluid (Passey et al., 2005). In aquatic environments, the relationship between fish carbonate and diet δ^{13} C values is complicated as primary producers constituting the base of fish food webs have $\delta^{13}C$ values that can vary due to differences in productivity, dissolved CO₂ concentration, bicarbonate utilization, as well as variations in the mixing of atmospheric and respired CO₂ under different flow conditions (see Clementz and Koch (2001) for a review). Moreover, for some fish carbonate biogenic components such as otoliths, a substantial amount, if not all, of the carbon may also be derived from dissolved inorganic carbon (DIC) of the ambient water (McConnaughey et al., 1997; Thorrold et al., 1997). However, ¹³C-enrichments of 10–16‰ between carbonate and diet have been observed for carnivorous sharks (Rau et al., 1983; Vennemann et al., 2001). Given the similarity with terrestrial mammals ¹³C-enrichments between carbonate and diet, it was suggested that the carbon in the carbonate of shark teeth is compatible with a predominantly dietary source of carbon (Vennemann et al., 2001).

2. Sample collection

One hundred and one apatite remains consisting of theropod dinosaur teeth, titanosaurid sauropod teeth, pterosaur teeth, crocodilian teeth and osteoderms, turtle bony plates and fish teeth and scales, were recovered from two Brazilian (Araripe Basin, Crato and Romualdo formations, and Laje Do Coringa, Alcântara Formation) and six Moroccan localities (Tafilalt region, Kem Kem beds). Samples were collected as isolated specimens, cleaned and analyzed for their oxygen and carbon isotope compositions of phosphate and carbonate. Published $\delta^{18}O_p$ values of similar vertebrates remains coming from Tunisia (Chenini Member), Morocco (Kem Kem beds) and Brazil (Alcântara Formation) were added to the dataset (Amiot et al., 2006, 2010). Except for small teeth and scales for which bulk analyses were performed, enamel was preferentially selected. As reptile teeth are continuously replaced and take several months to grow (Erickson, 1996a,b), they can record seasonal variations in the $\delta^{18}O_w$ values of ingested surface water. In order to retrieve mean annual values of local waters, enamel was sampled from the base to the apex of each tooth and several teeth from each locality were analyzed. For each sample, the most mineralized apatite part was selected, namely reptile tooth enamel, dense bone layers from turtle shells and crocodilian osteoderms, and ganoine (an enamel-like apatitic tissue) covering the surface of fish scales. Sample description is reported in Table 1.

Calculation of past geographic coordinates of the sampling sites was made using the Apparent Polar Wander Path (APWP) of South American and African continents calculated by Besse and Courtillot (2002). This APWP was constructed using a selection of the best paleomagnetic poles available for South America and Africa that were averaged over 20-My time windows. Paleolatitudes and associated uncertainties are shown in Table 2. These uncertainties depend on the quality and number of magnetic poles used to build the APWP.

3. Analytical techniques

Measurements of oxygen isotope compositions of apatite phosphate consist in isolating the phosphate ions using acid dissolution and anion-exchange resin, according to a protocol derived from the original method published by Crowson et al. (1991) and slightly modified by Lécuyer et al. (1993). Silver phosphate was quantitatively precipitated in a thermostatic bath set at a temperature of 70 °C. After filtration, washing with double deionised water, and drying at 50 °C, 15 mg of Ag₃PO₄ was mixed with 0.8 mg of pure powder graphite. $^{18}\text{O}/^{16}\text{O}$ ratios were measured by reducing silver phosphates to CO₂ using graphite reagent (O'Neil et al., 1994; Lécuyer et al., 1998). Samples were weighed into tin reaction capsules and loaded into quartz tubes and degassed for 30 min at 80 °C under vacuum. Each sample was heated at 1100 °C for 1 min to promote the redox reaction. The CO₂ produced was directly trapped in liquid nitrogen to avoid any kind of isotopic reaction with quartz at high temperature. CO₂ was then analyzed with a Thermo-Finnigan MAT253 mass spectrometer at the Institute of Geology and Geophysics, Chinese Academy of Sciences. Isotopic compositions are quoted in the standard δ notation relative to V-SMOW. Silver phosphate precipitated from standard NBS120c (natural Miocene phosphorite from Florida) was repeatedly analyzed ($\delta^{18}O = 21.7 \pm 0.2\%$; n = 31) along with the silver phosphate samples derived from the fossil vertebrate remains.

For the oxygen and carbon isotope analysis of carbonate, about 10 mg of enamel, dentine or bone powder was pre-treated according to the procedure of Koch et al. (1997). Powders were washed with a 2% NaOCl solution to remove organic matter, followed by a 0.1 M acetic acid solution to remove diagenetic carbonates. The powder/ solution ratio was kept constant to 0.04 g mL^{-1} for both treatments. Each treatment lasted for 24 h and samples were rinsed five times with distilled water. From the pre-treated powders a 2 mg aliquot was analyzed using a Thermo Finnigan Gasbench II following a procedure adapted after Spoetl and Vennemann (2003). Five drops of 100% orthophosphoric acid were added allowing the samples to react at 72 °C for 1 h under a He atmosphere before starting 10 measurement cycles of the isotopic composition of the produced CO₂ with a Finnigan MAT 253 continuous flow isotope ratio mass spectrometer. The measured carbon and oxygen isotopic compositions were normalized relative to the NBS-19 calcite standard. The normalization incorporates the CO₂-carbonate acid fractionation factor for calcite. Reproducibility for the carbon and oxygen isotopic compositions of apatite carbonate is better than $\pm 0.1\%$ and $\pm 0.2\%$, respectively. The carbon and oxygen isotopic compositions are expressed as δ -values relative to V-PDB for carbon and V-SMOW for oxygen.

4. Results

Oxygen and carbon isotope measurements of apatite phosphate and carbonate are reported in Table 1. The whole dataset ranges from 16.5% to 22.9% in $\delta^{18}O_p$ values, from 21.5% to 29.3% in $\delta^{18}O_c$ values and from -14.0% to -1.7% in $\delta^{13}C_c$ values. For each of the 9 localities, mean $\delta^{18}O_p$ values for each taxonomic group of reptiles are plotted against their calculated $\delta^{18}O_w$ value (Fig. 2). Significant $\delta^{18}O_p$ value differences are observed at any given locality between the various groups of reptiles, with range from 16.4% to 22.2% for dinosaurs (16.4% to 22.2% for spinosaurid theropods, 18.2% to 22.2% for other theropods and 19.2% to 19.9% for sauropods), from 18.3% to 22.9% for pterosaurs, from 17.5% to 19.9% for crocodilians and from 16.7% to 19.7% for turtles. Fish $\delta^{18}O_p$ values range from 19.2% to 22.3%.

Oxygen isotope compositions of apatite carbonate at Laje do Coringa and Moroccan localities are roughly linearly correlated to $\delta^{18}O_p$ values, with a slope close to unity, whereas $\delta^{18}O_p$ and $\delta^{18}O_c$ values at Araripe basin locality show a large scattering (Fig. 3). Offsets can also be observed between mean $\delta^{13}C_c$ values of apatite carbonate from different taxonomic groups, especially between fish taxa that have $\delta^{13}C_c$ values ranging from -11.1% to -1.5%, as shown in Fig. 4 where they are plotted against their corresponding $\delta^{18}O_p$ values. Reptile $\delta^{13}C_c$ values range from -12% to -7%, indicating that they were feeding on C₃ plant-based foodwebs.

Table 1

Taxon, phosphatic tissue, location, and age are given along with oxygen and carbon isotope compositions of apatite phosphate and carbonate of mid-Cretaceous vertebrate samples.

Sample no.	Skeletal tissue	Taxon	Locality name	Country	Formation	Age	$\delta^{18}O_P$	$\delta^{18}O_C$	$\delta^{13}C_{C}$	Data origin	
							(% V-SMOW)	(% V-SMOW)	(‰ V-PDB)		
BR13	Tooth bulk	cf. Carcharodontosaurus	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	20.4	24.7	-11.1	This study	
BR14	Tooth bulk	cf. Carcharodontosaurus	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	20.9	26.6	-10.2	This study	
BR15	Tooth bulk	cf. Carcharodontosaurus	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	22.2	29.3	-8.4	This study	
BR16	Tooth bulk	Spinosauridae	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	20.4	25.3	-7.9	This study	
BR17	Tooth bulk	Spinosauridae	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	19.5	24.9	- 7.5	This study	
BR18	Tooth bulk	Spinosauridae	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	19.6	26.6	-4.3	This study	
BR19	Tooth bulk	Spinosauridae	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	19.9	25.5	-9.0	This study	
BR20	Tooth bulk	Spinosauridae	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	19.4	25.4	-6.8	This study	
BR21	Tooth bulk	Spinosauridae	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	19.4	25.6	-7.1	This study	
BR22	Tooth enamel	Spinosauridae	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	19.1	25.3	-6.4	This study	
BR23	Tooth bulk	Spinosauridae	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	18.7	25.0	-4.0	This study	
BR24	Tooth enamel	Crocodylia	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	19.2	26.1	-10.1	This study	
BR25	Tooth bulk	Crocodylia	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	19.7	26.5	-9.8	This study	
BR26	Tooth enamel	Crocodylia	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	19.1	25.5	-9.0	This study	
BR27	Tooth bulk	Ceratodontiformes indet.	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	20.2	26.8	-8.0	This study	
BR28	Tooth bulk	Ceratodontiformes indet.	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	19.5	25.8	-6.9	This study	
BR29	Fin ray bone	Mawsonia	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	19.1	25.0	-6.9	This study	
BR30	Fin ray bone	Mawsonia	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	18.8	24.9	-6.3	This study	
BR31	Fin ray bone	Mawsonia	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	20.2	25.5	-10.1	This study	
BR32	Fin ray bone	Mawsonia	Laje do Coringa	Brazil	Alcântara Fm.	Early Cenomanian	20.2	26.5	-5.1	This study	
BR01	Tooth enamel	Anhanguerid pterosaur	Araripe Basin	Brazil	Crato Fm.	Aptian—Albian	21.0	26.7	-9.5	This study	
BR02	Tooth bulk	Anhanguerid pterosaur	Araripe Basin	Brazil	Crato Fm.	Aptian—Albian	20.4	23.2	-8.3	This study	
BR03	Tooth bulk	Anhanguerid pterosaur	Araripe Basin	Brazil	Crato Fm.	Aptian—Albian	20.7	25.1	-10.2	This study	
BR04	Tooth bulk	Anhanguerid pterosaur	Araripe Basin	Brazil	Crato Fm.	Aptian—Albian	21.5	26.3	-9.8	This study	
BR05	Tooth bulk	Anhanguerid pterosaur	Araripe Basin	Brazil	Crato Fm.	Aptian—Albian	21.2	25.9	-10.0	This study	
BR06	Tooth bulk	Anhanguerid pterosaur	Araripe Basin	Brazil	Crato Fm.	Aptian—Albian	21.4	24.8	-8.9	This study	
BR07	Tooth bulk	Anhanguerid pterosaur	Araripe Basin	Brazil	Crato Fm.	Aptian—Albian	21.2	21.7	-7.5	This study	
BR08	5 bulk teeth	Neoproscinetes penalvai	Araripe Basin	Brazil	Romualdo Fm.	Aptian—Albian	22.5	26.1	-1.7	This study	
BR09	Tooth enamel	Neoproscinetes penalvai	Araripe Basin	Brazil	Romualdo Fm.	Aptian—Albian	22.5	22.6	-3.0	This study	
BR10	Tooth enamel	Neoproscinetes penalvai	Araripe Basin	Brazil	Romualdo Fm.	Aptian—Albian	21.9	22.4	- 1.5	This study	
BR11	Scales	Vinctifer	Araripe Basin	Brazil	Romualdo Fm.	Aptian—Albian	19.9	24.5	-9.3	This study	
BR12	Shell bone	Araripemys	Araripe Basin	Brazil	Romualdo Fm.	Aptian—Albian	19.7	21.5	-9.6	This study	
M-BL-022	Tooth enamel	Carcharodontosaurus saharicus	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	19.0	-	-	Amiot et al. (2006)	
M-BL-023	Tooth enamel	Carcharodontosaurus saharicus	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	18.2	-	-	Amiot et al. (2006)	
M-BL-024	Tooth bulk	Theropoda	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	19.1	25.7	- 8.5	This study	
M-BL-025	Tooth bulk	Theropoda	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	19.0	25.6	-11.1	This study	
M-BL-036	Tooth enamel	Spinosaurus	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	19.3	-	-	Amiot et al. (2006)	
M-BL-041	Tooth enamel	Spinosaurus	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	20.6	27.6	- 10.8	This study	
M-BL-042	Tooth enamel	Spinosaurus	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	19.7	25.9	-9.5	This study	
M-BL-043	Tooth enamel	Spinosaurus	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	19.6	26.3	-10.0	This study	
M-BL-044	Tooth enamel	Spinosaurus	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	19.8	26.5	-14.0	This study	
M-BL-045	Tooth enamel	Spinosaurus	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	19.1	25.9	-12.1	This study	
M-BL-003bis	Tooth enamel	Anhanguerid pterosaur	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	18.3	24.3	-11.3	This study	
M-BL-013	Osteoderm	Crocodylia	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	16.7	-	-	Amiot et al. (2006)	
M-BL-038a	Tooth enamel	cf. Sarcosuchus	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	18.4	-	-	Amiot et al. (2006)	
M-BL-040	Tooth enamel	Crocodylia	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	17.4	-	-	Amiot et al. (2006)	
M-BL-014	Shell bone	Chelonia	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	16.7	-	-	Amiot et al. (2006)	
M-BL-016	Shell bone	Chelonia	Bou Laalou	Morocco	Kem Kem beds	Early Cenomanian	17.2	-	-	Amiot et al. (2006)	
M-KS-013	Tooth bulk	Carcharodontosaurus saharicus	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	18.6	26.3	-9.4	This study	
M-KS-023	Tooth enamel	Carcharodontosaurus saharicus	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	20.7	-	-	Amiot et al. (2006)	
M-KS-024	Tooth enamel	Carcharodontosaurus saharicus	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	19.9	-	_	Amiot et al. (2006)	
M-KS-028	Tooth enamel	Carcharodontosaurus saharicus	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	18.9	-	-	Amiot et al. (2006)	
M-KS-015	Tooth bulk	Theropoda	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	18.6	25.6	-11.2	This study	
M-KS-016	Tooth bulk	Theropoda	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	19.8	26.7	-7.6	This study	

MAX 50 MX 500 MX 50	M-KS-009	Tooth enamel	Spinosaurus	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	19.8	26.6	-10.7	This study
MAX 500 Toole names Spinsonanta Relation shifts Marcer at 20019 Andrer at 20019 MAX 5007 Toole names Spinsonanta Relation shifts Marcer at 20039 MAX 5007 Toole names Spinsonanta Relation shifts Marcer at 20039 MAX 5007 Toole names Spinsonanta Relation shifts Marcer at 20039 MAX 5007 Toole names Toole names Relation shifts Marcer at 20039 MAX 5007 Toole names Toole names Relation shifts Marcer at 20039 MAX 5007 Toole names Toole names Relation shifts Marcer at 20039 MAX 5007 Toole names Cocolpia Marcer at 20039 Marcer at 20039 MAX 5007 Stole names Cocolpia Marcer at 20039 Marcer at 20039 MAX 5008 Stole names Cocolpia Marcer at 20039 Marcer at 20039 MAX 5008 Stole names Cocolpia Marcer at 20039 Marcer at 20039 MAX 5008 Stole names Cocolpia Marcer at 20039 Marcer at 20039	M-KS-014	Tooth enamel	Spinosaurus	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	19.4	26.9	-8.9	This study
MALSADTook examesyntownesNethils ShineMerceKen Mern KeFully Cremunian17.8A hard et al. (2005)MA KG MUTook exameSyntownesKirelii ShineMerceKirelii ShineKirelii ShineMarceKirelii Shine	M-KS-006	Tooth enamel	Spinosaurus	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	16.9	-	-	Amiot et al. (2010)
MA-5680Totak manufSymmauryMettilla ShinaMaccaKan Kan DabaLally Commania9.3Anac Ket al. (200)MAS287Totak meanufSymmauryRettilla ShinaMaccaKan Kan DabaEndy Commania10.3Anac Ket al. (200)MAS281Totak meanufSymmauryRettilla ShinaMaccaKan Kan DabaEndy Commania10.3Anac Ket al. (200)MAS281Totak meanufCorcalylaRettilla ShinaMaccaKan Ken DabaEndy Commania15.5Anac Ket al. (200)MAS281Shina LawCorcalylaRettilla ShinaMaccaKen Ken DabaEndy Commania15.7Anac Ket al. (200)MAS281Shina LawCorcalylaRettilla ShinaMaccaKen Ken DabaEndy Commania16.7Anac Ket al. (200)MAS281Shina LawCorcalylaRettilla ShinaMaccaKen Ken DabaEndy Commania <td< td=""><td>M-KS-007</td><td>Tooth enamel</td><td>Spinosaurus</td><td>Khetitila Srhira</td><td>Morocco</td><td>Kem Kem beds</td><td>Early Cenomanian</td><td>17.8</td><td>-</td><td>_</td><td>Amiot et al. (2006)</td></td<>	M-KS-007	Tooth enamel	Spinosaurus	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	17.8	-	_	Amiot et al. (2006)
MASK 2017 Tothe examel Spinonizmic Decitial Strini Morecol Rom beds Endity Communiant 20.3	M-KS-008	Tooth enamel	Spinosaurus	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	19.8	-	_	Amiot et al. (2010)
MAS-4901 Tools name Spinsouring Identità Spinsouring In Spinsouring 1/3 - - Americal (2005) MAS-4901 Tools name Cascolyla Identità Spinso Ken Ken bods Entry Commanian 18.8 - - Americal (2005) MAS-4901 Tools name Cascolyla Identità Spinso Ken Ken bods Entry Commanian 18.8 - - Americal (2005) MAS-4901 Stolob Cascolyla Identità Spinso Ken Ken bods Entry Commanian 16.7 - - Americal (2005) MAS-1918 Stolob Cascolyla Identità Spinso Ken Ken bods Entry Commanian 16.7 - - Americal (2005) MAS-1918 Stolob Carlpiolers Identità Shina Merce Ken Ken bods Entry Commanian 16.7 - - - Americal (2005) MAS-1918 Stolob Carlpiolers Identità Shina Merce Ken Ken bods Entry Commanian 16.7 - - - -	M-KS-036	Tooth enamel	Spinosaurus	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	20.3	-	_	Amiot et al. (2006)
MA-B-301 Toole name Tunnosurinde / Subsection Kleicilla Shina Morece Korn Konb deal Bits - - And tel (2005) MK-SG1 Toole named Corcodyla Kleitili Shina Morrece Kenn Ken block Eably Genomaina 18.8 - - Antot et al (2005) MK-SG40 Toole named Corcodyla Keitili Shina Morrece Kenn Ken block Eably Cenomaina 18.4 - - Antot et al (2005) MK-SG40 Toole named Corcodyla Keitili Shina Morrece Kenn Ken block Eably Cenomaina 18.4 25.7 -3.3 This study MK-SG400 Sole panior Lippideer Keitili Shina Morrece Kenn Ken block Eably Cenomaina 18.4 25.7 -3.3 This study MK-SG40 Toole block Lippideer Keitili Shina Morrece Kein Kein block Eably Cenomaina 18.3 25.7 -3.3 This study MK-SG40 Toole block Lippideer Keitili Shina Morrece Kein Kei	M-KS-037	Tooth enamel	Spinosaurus	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	17.3	-	_	Amiot et al. (2006)
MA-S-2011 Toch enamel Creacylyia Kleritia Shrina Morece Ken Ken besh Euly Cenaumian 18.8 Arrier Cel. (2006) MK-S040 Toch bulk Creacylyia Kheritia Shrina Morece Ken Ken besh Euly Cenaumian 17.8	M-KS-001a	Tooth enamel	Titanosauridae	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	19.9	26.7	-9.5	This study
Mi-SG-14 Tosh ename Cocoodyla Klerithi Shrian Morace Ken kem beb Exty commanian 16.5 - - Amate et al. (2006) Mi-SG-101 Skell born Circoodyla Kentriki Shrian Morace Kent Kem beb Exty commanian 16.7 - - Amate et al. (2006) Mi-SG-101 Sale bulk Cirplater Kentriki Shrian Morace Kentriki Shrian Morace <t< td=""><td>M-KS-031</td><td>Tooth enamel</td><td>Crocodylia</td><td>Khetitila Srhira</td><td>Morocco</td><td>Kem Kem beds</td><td>Early Cenomanian</td><td>18.8</td><td>-</td><td>_</td><td>Amiot et al. (2006)</td></t<>	M-KS-031	Tooth enamel	Crocodylia	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	18.8	-	_	Amiot et al. (2006)
M-KS-041 Took hulk Conception Khertiki Schin More Cell Schin Instruct el. (2006) M-KS-012 Scale pannice C. Epideers Khertiki Schin More Cell Schin 16.7 - - Ambre el. (2006) M-KS-010 Scale pannice C. Epideers Khertiki Schin More Cell Schin Biol 25.7 - 3.8 11.8 42.9 - - Ambre el. (2006) M-KS-010 Scale pannice C. Epideers Khertiki Schin More Cell Schin Scale pannice - - Ambre el. (2007) M-KS-020 Scale pannice C. Epideers Khertiki Schin More Cell Schin Scale Schin - - - - Ambre el. (2006) M-KS-020 Tooch hulk Dipola inder. Khertiki Schin More Cell Schin Scale Schin -	M-KS-034	Tooth enamel	Crocodylia	Khetitila Srhira	Morocco	Kem Kem beds	Farly Cenomanian	16.5	_	_	Amiot et al. (2006)
NK-S012 Stell blas Chebrais Klerital Stila Manor ed. Loran Manor ed. Loran Manor ed. Loran Manor ed. Loran MK-S0198 Scale bulk cf. Ipplders Kheitial Stila Manor ed. Loran Factor ed. Lor	M-KS-040	Tooth bulk	Crocodylia	Khetitila Srhira	Morocco	Kem Kem beds	Farly Cenomanian	17.8	_	_	Amiot et al. (2006)
NH-S-010 Scale genome cf.Lppdeer Ricettia String Morece Ken Ken bed Early Commanian 21.1 28.0 -4.9 This study MK-S-0108 Scale bulk cf.Lppdeers Ricettia String Morece Ken Ken beds Early Commanian 18.4 25.4 -5.3 This study MK-S-0108 Scale bulk cf.Lppdeers Ricettia String Morece Ken Ken beds Early Commanian 20.4 2.7.2 -4.6 This study MK-S-0108 Scale genome cf.Lppdeers Ricettia String Morece Ken Ken beds Early Commanian 20.0 2.7.2 -4.6 This study MK-S-0107 Toorb bulk Dipon inder. Ricettia String Morece Ken Ken beds Early Commanian 19.9 25.0 -2.2 This study MK-S-0107 Toorb bulk Dipon inder. Chait Morece Ken Ken beds Early Commanian 21.4 28.2 -0.6 This study MK-1011 Toorb channel Spinsozurar Chait Morece	M-KS-012	Shell hone	Chelonia	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	16.7	_	_	Amiot et al. (2006)
M-KS 0198 Scale bulk cl. Leptotres Rettilla Shrin Morreco Kenn Kenn beds Larly Ceromanian 18.4 25.4 5.2 This study M-KS 0198 Scale bulk cl. Leptotres Rettilla Shrin Morreco Kenn Kenn beds Early Ceromanian 20.4 25.3 10.7 This study M-KS 0198 Scale bulk cl. Leptotres Rettilla Shrin Morreco Kenn Kenn beds Early Ceromanian 13.1 25.4 10.3 This study M-KS 0298 Conth halk Diponi indet. Rettilla Shrin Morreco Kenn Kenn beds Early Ceromanian 13.1 25.4 10.3 This study M-KS 045 Torth halk Portegriters Rettilla Shrin Morreco Kenn Kenn beds Early Ceromanian 21.4 23.3 3.3 This study M-C1-0121 Torth namel Spinsaurus Chalf Morreco Kenn Kenn beds Early Ceromanian 17.5 2 3.8 Antie et al. (2011) M-C1-010 Torth namanel Spinsaurus Ch	M-KS-019A	Scale ganoine	cf Lenidotes	Khetitila Srhira	Morocco	Kem Kem beds	Farly Cenomanian	21.1	28.0	-49	This study
NH-S-0190 Scale bulk of Lepróters Rettilla Shrin Morecco Kem Kem beds LerfV consumana 18.6 2.7.7 -5.3 This study NH-S-0190 Scale bulk of Lepróters Rettilla Shrin Morecco Kem Kem beds LerfV consumana 2.0.9 2.7.2 -4.6 This study NH-S-0190 Scale bulk Of Noermoton fracmus Kettilla Shrin Morecco Kem Kem beds LerfV consumana 1.9.2 2.5.2 -10.1 This study NH-S-020 Tooth bulk Of Noermoton fracmus Kettilla Shrin Morecco Kem Kem beds Early Consumana 1.9.2 2.5.2 -1.2 This study NH-S-040 Tooth bulk Protopicran Kettilla Shrin Morecco Kem Kem beds Early Consumana 2.9.2 -2.2 This study NH-F-0401 Tooth bulk Terropode Chadra Morecco Kem Kem beds Early Consumana 2.9.2 -2.5 This study NH-F-0401 Tooth channel Spinsaurus Chadr Morecco Kem Kem beds<	M-KS-019R	Scale bulk	cf Lepidotes	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	18.4	25.4	-82	This study
bit/S-0190 Scale balk of Lepfolder Defittilis Shira Merceo Kern Kenn beds Early Commanian 20.4 25.3 -10.7 This study M-SG-0190 Scale pancine Chapfolder Feltilis Shira Moreco Kern Kenn beds Early Commanian 19.1 25.4 -10.3 This study M-SG-020 Tooth bulk Diponi indet. Feltilis Shira Moreco Kern Kenn beds Early Commanian 19.9 25.4 -10.3 This study M-SG-021 Tooth bulk Propripers Feltilis Shira Moreco Kern Kenn beds Early Commanian 21.4 28.2 -9.6 This study M-GH-011 Tooth channel Carchinedintosonna sudirizon Chalt Moreco Kern Kenn beds Early Commanian 21.4 28.3 -8.3 This study M-GH-012 Tooth channel Carchinedintosonna sudirizon Chalt Moreco Kern Kenn beds Early Commanian 21.4 28.2 -15.1 This study M-GH-013 Tooth channel Sylinosourus <td< td=""><td>M-KS-019C</td><td>Scale bulk</td><td>cf Lepidotes</td><td>Khetitila Srhira</td><td>Morocco</td><td>Kem Kem beds</td><td>Early Cenomanian</td><td>18.6</td><td>25.7</td><td>-53</td><td>This study</td></td<>	M-KS-019C	Scale bulk	cf Lepidotes	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	18.6	25.7	-53	This study
M-KS-019 Scale gamme cf. Jupiders Kiertills Shrin Mercox Ken beds Early Genomanian 22.0 27.2 -4.6 This study M-KS-025 Tonch bulk Diposi inder, Keneritäs Shrin Mercox Ken beds Early Genomanian 19.2 25.5 -10.1 This study M-KS-025 Tonch bulk Drobe inder, Kentilla Shrin Mercox Ken Ken beds Early Genomanian 19.9 25.5 -10.1 This study M-KS-025 Tonch bulk Protoperus Kentilla Shrin Mercox Ken Ken beds Early Genomanian 19.9 25.5 -10.2 This study M-GH-013 Tonch enand Gardmodintoamus subrins Chalt Mercox Ken Ken beds Early Genomanian 11.6 23.2 -2.8 This study M-GH-024 Tonch enand Spinosums Chalt Mercox Ken Ken beds Early Genomanian 11.8 - - Americ et al. (2010) M-GH-026 Tonch enand Spinosums Chalt Mercox <td< td=""><td>M-KS-019D</td><td>Scale bulk</td><td>cf Lepidotes</td><td>Khetitila Srhira</td><td>Morocco</td><td>Kem Kem beds</td><td>Farly Cenomanian</td><td>20.4</td><td>25.3</td><td>- 10.7</td><td>This study</td></td<>	M-KS-019D	Scale bulk	cf Lepidotes	Khetitila Srhira	Morocco	Kem Kem beds	Farly Cenomanian	20.4	25.3	- 10.7	This study
M-K-025 Tordt, Duik cf. Recruinds affroms Kenetidis Shina Morece Ken Ken beds Early Genomanian 19.1 25.4 -10.3 This study M-KS-026 Tooth buik d. Norceo Ken Ken beds Early Genomanian 19.9 26.1 -10.3 This study M-KS-042 Tooth buik d. Norceo Ken Ken beds Early Genomanian 19.9 26.1 -10.3 This study M-KS-042 Tooth buik Gradmontonizons subricis Chailt Morece Ken Ken beds Early Genomanian 21.2 28.2 -9.6 This study M-CH-012 Tooth buik Theoropda Chailt Morece Ken Ken beds Early Genomanian 21.0 22.2 -9.6 This study M-CH-041 Tooth buik Theoropda Chailt Morece Ken Ken beds Early Genomanian 11.0 22.2 -1.5 Annie et al. (2010) M-CH-0407 Tooth caunel Spinosaura Chailt Morece Ken Ken beds Early Genomanian 11.2 2.4 -1.5 This study M-CH-0405 Spinosaura	M-KS-019D	Scale ganoine	cf Lepidotes	Khetitila Srhira	Morocco	Kem Kem beds	Farly Cenomanian	20.4	23.5	-46	This study
NH-G-263 Tooth bulk Digrain indet. Kieruita Srinir Moreco Kem Kem beds Early Ceramanian 19.7 25.5 -10.3 This study NH-G-045 Tooth bulk Proteprins Kieruita Srinir Moreco Kem Kem beds Early Ceromanian 19.9 25.1 -10.3 This study NH-G-045 Tooth enamel Carchrondombonanes soharics Chait Moreco Kem Kem beds Early Ceromanian 21.4 28.3 -8.3 This study NG-10-013 Tooth enamel Carchrondombonanes soharics Chait Moreco Kem Kem beds Early Ceromanian 21.4 28.3 -8.3 This study NG-10-01 Tooth enamel Spinesarurs Chait Moreco Kem Kem beds Early Ceromanian 19.4 - - Aminet et al. (2010) NG-10-05 Tooth enamel Spinesarurs Chait Moreco Kem Kem beds Early Ceromanian 19.4 - - Aminet al. (2010) NG-10-05 Tooth enamel Spinesarurs Chait Mor	M-KS-025	Tooth bulk	cf Neoceratodus africanus	Khetitila Srhira	Morocco	Kem Kem beds	Farly Cenomanian	19.1	25.4	- 10 3	This study
NEG-207 Toolb bulk d. Proceemake giremus Kieritä sfürz Morrece Kem Kem bed Eury Commanian 19.9 26.1 -10.3 Tüs study M-G1-012n Toolb bulk Prorigeters Kieritä sfürz Morrece Kem Kem bed Eury Commanian 22.1 28.2 -9.6 This study M-G1-012n Toolb enamel Gardenodomosame solvarius Chaft Morrece Kem Kem bed Eury Commanian 22.4 28.2 -9.6 This study M-G1-041 Toolb enamel Spinosaruus Chaft Morrece Kem Kem bed Eury Commanian 18.4 - - Annote tal. (2010) M-G1-025 Toolb enamel Spinosaruus Chaft Morrece Kem Kem bed Eury Commanian 18.4 - - Annote tal. (2010) M-G1-025 Toolb enamel Spinosaruus Chaft Morrece Kem Kem bed Eury Commanian 19.2 25.4 -11.5 This study M-G1-030 Toolb enamel Spinosaruus Chaft Morrece Kem Ke	M-KS-025	Tooth bulk	Dippoi indet	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	10.7	25.4	_ 10.5	This study
NEX-God Tooth Dukk Propression Rest Neth Early Censmania 19.9 25.9 -1.2 This study MCL10123 Tooth enamel Carchendonizosamus subarica Chaif. Morreco Kem Kein bed Early Censmania 2.1.4 28.2 -9.5 This study MCL1013 Tooth enamel Garchendonizosamus subarica Chaif. Morreco Kem Kein bed Early Censmania 2.1.4 28.3 -8.3 This study MCL1043 Tooth enamel Spinosarus Chaif. Morreco Kem Kein bed Early Censmania 17.8 -2.2 -1.15 This study MCL1043 Tooth enamel Spinosarus Chaif. Morreco Kem Kein bed Early Censmania 19.2 2.5.4 -11.5 This study MCL10403 Tooth enamel Spinosarus Chaif. Morreco Kem Kein bed Early Censmania 19.2 2.5.4 -11.1 This study MCL10403 Tooth enamel Spinosarus Chaif. Morreco Kem Kein bed Early Censmania	M_KS_027	Tooth bulk	of Neoceratodus africanus	Khetitila Srhira	Morocco	Kem Kem beds	Early Cenomanian	10.0	25.5	- 10.1	This study
MC-10-12n Tools examel Cardinadiminatures subrices Chaif Moreco Kem Ken bels Early (comonniani 21.1 28.2 -6.8 This study MC-10-101 Tools examel Cardinadiminatures subrices Chaif Moreco Kem Ken bels Early (comonniani 21.4 28.3 -8.3 This study MC-10-01 Tools examel Spinosaurus Chaif Moreco Kem Ken bels Early (comonniani 18.4 - - Amore et al. (2010) MC-10-03 Tools examel Spinosaurus Chaif Moreco Kem Ken bels Early (comonniani 19.1 25.2 -11.5 This study MC-10-03 Tools examel Spinosaurus Chaif Moreco Kem Ken bels Early (comonniani 19.2 25.4 -11.5 This study MC-10-03 Tools examel Spinosaurus Chaif Moreco Kem Ken bels Early (comonniani 19.2 25.2 -11.1 This study MC-10-01 Tools examel Choody (la Chaif Moreco	M-KS-027	Tooth bulk	Protonterus	Khetitila Schira	Morocco	Kem Kem beds	Early Cenomanian	10.0	25.0	- 3.2	This study
nc 1-113Toth examel toth pulkConduction toth pulkChaftMorece MoreceKem Ken beds and Cenomanian21.423.3-1.3This studyM-CH-104Toth bulkTheropodChaftMorece MoreceKem Ken beds and Cenomanian20.926.2-9.6-9.6This studyM-CH-104Toth enamelSpinsournsChaftMorece MoreceKem Ken beds and Cenomanian18.4Anniet et al. (2010)M-CH-037Toth enamelSpinsournsChaftMorece MoreceKem Ken beds and Cenomanian19.125.2-11.5This studyM-CH-037Toth enamelSpinsournsChaftMorece MoreceKem Ken beds and Cenomanian19.225.2-11.1This studyM-CH-037Toth enamelSpinsournsChaftMorece MoreceKem Ken beds bedsEarly Cenomanian19.225.2-11.1This studyM-CH-0365Toth enamelAnageerid preosaurChaftMorece MoreceKem Ken beds bedsEarly Cenomanian19.0Anniet et al. (2060)M-CH-011Toth bulkCroodyliaChaftMorece MoreceKem Ken beds bedsEarly Cenomanian19.0Anniet et al. (2060)M-CH-028Shell boneCheloniaChaftMorece MoreceKem Ken beds bedsEarly Cenomanian18.0Anniet et al. (2060)M-CH-031Shell boneCheloniaChaftMorece Morec	M_CH_012n	Tooth enamel	Carcharodontosaurus sabaricus	Chaoft	Morocco	Kem Kem beds	Early Cenomanian	22.1	23.5	- 9.6	This study
$ \begin{array}{c} \mbox{rep}{res} \mbox{rep}{res} \mbox{res} \mbo$	M_CH_013	Tooth enamel	Carcharodontosaurus saharicus	Chaoft	Morocco	Kem Kem beds	Early Cenomanian	22.1	20.2	- 83	This study
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	M-CH-014	Tooth bulk	Theropoda	Chaoft	Morocco	Kem Kem beds	Early Cenomanian	21.4	26.5	- 9.5	This study
MC1407Tooth reamejoinsumChartMonecoKen bedsEarly Commanian18.4 $ -$ Aunot et al (2010)MC41035Tooth reamelSpinsumChartMonecoKem ken bedsEarly Commanian19.12.2 $ -$ Aunot et al (2010)MC41035Tooth reamelSpinsumChartMonecoKem ken bedsEarly Commanian19.2 $2.2.2$ $ -$	M-CH-004	Tooth enamel	Spinosaurus	Chaoft	Morocco	Kem Kem beds	Early Cenomanian	17.5	20.2	- 5.0	Amiot et al. (2010)
Inc. 1	M_CH_007	Tooth enamel	Spinosaurus	Chaoft	Morocco	Kem Kem beds	Early Cenomanian	18.4			Amiot et al. (2010)
Incl. or in the standJoint and the spin sourceChartMore constraintsLink <thlink< th="">Lin</thlink<>	M_CH_025	Tooth enamel	Spinosaurus	Chaoft	Morocco	Kem Kem beds	Early Cenomanian	10.4	25.2	_ 12.5	This study
In Cli 1000Tooth enamelSpinozanizaChantMorecoKem Kem bedsEarly Cenomanian19.22.5.41.1.2This StudyM-CH-005kTooth enamelAnanguerid ptensaurusChantMorecoKem Kem bedsEarly Cenomanian19.22.5.2-11.1This StudyM-CH-005kTooth enamelCrocodyliaChantMorecoKem Kem bedsEarly Cenomanian19.0Anniot et al. (2006)M-CH-010Tooth enamelCrocodyliaChantMorecoKem Kem bedsEarly Cenomanian19.0Anniot et al. (2006)M-CH-010Tooth bulkCrocodyliaChantMorecoKem Kem bedsEarly Cenomanian18.6Anniot et al. (2006)M-CH-029Shell boneCheloniaChantMorecoKem Kem bedsEarly Cenomanian18.8This studyM-CH-024Rostal tooth bulkCrocodyliaChantMorecoKem Kem bedsEarly Cenomanian19.926.3-9.1This studyM-TA-017Tooth enamelGurchardontszuurus subricusTakemoutMorecoKem Kem bedsEarly Cenomanian18.225.5-9.8This studyM-TA-026Tooth enamelSpinosaurusTakemoutMorecoKem Kem bedsEarly Cenomanian18.225.5-9.8This studyM-TA-028Tooth enamelSpinosaurusTakemoutMorecoKem Kem bedsEarly Cenomanian18.4 <td< td=""><td>M-CH-025</td><td>Tooth enamel</td><td>Spinosaurus</td><td>Chaoft</td><td>Morocco</td><td>Kem Kem beds</td><td>Farly Cenomanian</td><td>17.8</td><td>23.2</td><td>- 11.5</td><td>This study</td></td<>	M-CH-025	Tooth enamel	Spinosaurus	Chaoft	Morocco	Kem Kem beds	Farly Cenomanian	17.8	23.2	- 11.5	This study
InclusionDoth bulkTransourdageChantMoreceKern Kern bedsEarly Cenomanian19.229.3-11.3This studyM-CH-005bbTooth bunkTransourdageChantMoreceKern Kern bedsEarly Cenomanian20.925.3-8.5This studyM-CH-001Tooth bunkCrocodyliaChantMoreceKern Kern bedsEarly Cenomanian20.925.3-8.5This studyM-CH-011Tooth bunkCrocodyliaChantMoreceKern Kern bedsEarly Cenomanian20.225.8-11.2This studyM-CH-021Shell boneCheloniaChantMoreceKern Kern bedsEarly Cenomanian18.9Anniet et al. (2006)M-CH-021Shell boneCheloniaChantMoreceKern Kern bedsEarly Cenomanian18.9This studyM-TA-017Tooth enamelCarcharodontosaurus subricusTakemoutMoreceKern Kern bedsEarly Cenomanian20.126.0-9.0This studyM-TA-017Tooth enamelCarcharodontosaurus subricusTakemoutMoreceKern Kern bedsEarly Cenomanian18.9This studyM-TA-026Tooth enamelSpinosaurusTakemoutMoreceKern Kern bedsEarly Cenomanian18.626.1-10.7This studyM-TA-028Tooth enamelSpinosaurusTakemoutMoreceKern Kern bedsEarly Cenomanian18.425.7-9.8Thi	M_CH_037	Tooth enamel	Spinosaurus	Chaoft	Morocco	Kem Kem beds	Early Cenomanian	10.2	24.2	- 11.5	This study
InclusionsInternational interventionChank interventionProtect interventionCarlo interve	M-CH-0065	Tooth bulk	Titanosauridae	Chaoft	Morocco	Kem Kem beds	Early Cenomanian	10.2	25.4	- 11.J - 11.1	This study
M-CH-00005 Tooth Chain Chain More Chain Lang Chonanian Loc. Loc. Loc. And Chain M-CH-011 Tooth bulk Croodylia Chain More Co Kem Kem beds Early Cenomanian 10.0 - - Amidet 41 (2006) M-CH-013 Shell bone Chelonia Chain More Co Kem Kem beds Early Cenomanian 18.0 - - Amidet 41 (2006) M-CH-031 Shell bone Chelonia Chain More Co Kem Kem beds Early Cenomanian 18.9 - - Amidet 41 (2006) M-TA-017 Tooth enamel Carcharodontosaurus soharicus Takemout More Co Kem Kem beds Early Cenomanian 18.9 - - - This study M-TA-017 Tooth enamel Spinosaurus Takemout More Co Kem Kem beds Early Cenomanian 18.6 26.1 - 10.7 This study M-TA-027 Tooth enamel Spinosaurus Takemout More Co Kem Kem beds Early Cenoman	M_CH_005bis	Tooth enamel	Anhanguerid pterosaur	Chaoft	Morocco	Kem Kem beds	Early Cenomanian	20.0	25.2	- 85	This study
M-CH-010Tooth bulkCheckyliaChaaftMorocoKem Kem bedsEarly Cenomanian12.0Aniot et al. (2006)M-CH-029Shell boneCheloniaChaaftMorocoKem Kem bedsEarly Cenomanian18.6Amiot et al. (2006)M-CH-024Rostral tooth bulkOnchoprists numidusChaaftMorocoKem Kem bedsEarly Cenomanian18.9This studyM-TA-017Tooth enamelOnchoprists numidusChaaftMorocoKem Kem bedsEarly Cenomanian21.3This studyM-TA-019Tooth enamelCarcharodontosaurus saharicusTakemoutMorocoKem Kem bedsEarly Cenomanian19.926.3-9.1This studyM-TA-026Tooth enamelSpinosaurusTakemoutMorocoKem Kem bedsEarly Cenomanian18.626.1-10.7This studyM-TA-027Tooth enamelSpinosaurusTakemoutMorocoKem Kem bedsEarly Cenomanian18.425.7-9.9This studyM-TA-028Tooth enamelSpinosaurusTakemoutMorocoKem Kem bedsEarly Cenomanian19.425.1-7.9This studyM-TA-020Tooth bulkAnhanguerid pterosaurTakemoutMorocoKem Kem bedsEarly Cenomanian19.324.7-10.2This studyM-TA-020Tooth bulkAnhanguerid pterosaurTakemoutMorocoKem Kem bedsEarly Cenomanian19		Tooth on amal	Crocodulia	Chaoft	Morocco	Kem Kem bods	Early Conomanian	20.9	23.5	- 8.5	Amiot at al (2006)
$ \begin{array}{c} \operatorname{Here}{\operatorname{Here}}{\operatorname{Her}}{\operatorname{Here}}{\operatorname{Here}}{\operatorname{Here}}{\operatorname{Her}}{\operatorname{He}}{\operatorname{He}}{\operatorname{He}}{\operatorname{He}}{\operatorname{He}}{\operatorname{He}}{\operatorname{He}}}{\operatorname{He}}{$	M CH 011	Tooth bulk	Crocodylia	Chaoft	Morocco	Kem Kem bods	Early Conomanian	19.0	-	- 11.2	This study
In-Cl-025Shell boneCheloniaChaitMoroccoKen Ken bedsEarly Cenomanian16.0Anite et al. (2006)M-CH-024Rostral tooth bulkOnchoprists numidusChaitMoroccoKem Kem bedsEarly Cenomanian21.3This studyM-TA-017Tooth enamelCarcharodontosaurus saharicusTakemoutMoroccoKem Kem bedsEarly Cenomanian19.926.3-9.0This studyM-TA-017Tooth enamelSpinosaurusTakemoutMoroccoKem Kem bedsEarly Cenomanian18.626.1-10.7This studyM-TA-026Tooth enamelSpinosaurusTakemoutMoroccoKem Kem bedsEarly Cenomanian18.225.5-9.8This studyM-TA-027Tooth enamelSpinosaurusTakemoutMoroccoKem Kem bedsEarly Cenomanian18.425.7-9.9This studyM-TA-028Tooth bulkAnhanguerid pterosaurTakemoutMoroccoKem Kem bedsEarly Cenomanian19.425.1-7.9This studyM-TA-021bisTooth bulkAnhanguerid pterosaurTakemoutMoroccoKem Kem bedsEarly Cenomanian19.324.7-10.2This studyM-TA-021bisTooth bulkAnhanguerid pterosaurTakemoutMoroccoKem Kem bedsEarly Cenomanian19.324.7-10.2This studyM-TA-030Tooth bulkAnhanguerid pterosaurTakemoutMoroccoKem Kem bedsEarly Cenoma	M CH 020	Shell bone	Cholonia	Chaoft	Morocco	Kem Kem bods	Early Conomanian	19.6	23.8	-11.2	Amiot at al (2006)
M-CH-021Boilt bolkDicklageChainDiractEnd KDiractEnd KDiract <td>M-CH-029</td> <td>Shell bone</td> <td>Chelonia</td> <td>Chaoft</td> <td>Morocco</td> <td>Kem Kem beds</td> <td>Early Cenomanian</td> <td>18.0</td> <td>_</td> <td>_</td> <td>Amiot et al. (2000) Amiot et al. (2006)</td>	M-CH-029	Shell bone	Chelonia	Chaoft	Morocco	Kem Kem beds	Early Cenomanian	18.0	_	_	Amiot et al. (2000) Amiot et al. (2006)
M-TA-024Notice Work in WindowsCharateMarketCharateMarketCharateMarketCharateMarketCharateCarlo marketCarlo marketCarlo marketCarlo marketCarlo marketCarlo marketCarlo marketCarlo marketCarlo marketMarketCarlo marketMarketCarlo marketMarket	M_CH_024	Rostral tooth bulk	Onchopristis numidus	Chaoft	Morocco	Kem Kem beds	Early Cenomanian	21.3			This study
M-TA-017Tooth enamelCarchardontosaurus subarusTakemoutMoreccoKenn Kenn bedsEarly Cenomanian20.120.0-9.0This studyM-TA-026Tooth enamelSpinosaurusTakemoutMoreccoKenn Kem bedsEarly Cenomanian18.626.1-10.7This studyM-TA-026Tooth enamelSpinosaurusTakemoutMoreccoKem Kem bedsEarly Cenomanian18.225.5-9.9This studyM-TA-031Tooth bulkSpinosaurusTakemoutMoreccoKem Kem bedsEarly Cenomanian19.724.9-7.7This studyM-TA-031Tooth bulkSpinosaurusTakemoutMoreccoKem Kem bedsEarly Cenomanian19.425.1-7.7This studyM-TA-021bisTooth bulkAnhanguerid pterosaurTakemoutMoreccoKem Kem bedsEarly Cenomanian19.324.7-10.2This studyM-TA-030Tooth bulkAnhanguerid pterosaurTakemoutMoreccoKem Kem bedsEarly Cenomanian19.223.9-9.1This studyM-TA-040OsteodermCrecodyliaTakemoutMoreccoKem Kem bedsEarly Cenomanian19.225.5-9.6This studyM-TA-040OsteodermCrecodyliaTakemoutMoreccoKem Kem bedsEarly Cenomanian19.225.1-10.7This studyM-TA-040Steol DoneCheloniaTakemoutMoreccoKem Kem bedsEarly Cenomanian19.322.9-11.3 <td>M TA 017</td> <td>Tooth on amal</td> <td>Carcharodontosaurus sabaricus</td> <td>Takomout</td> <td>Morocco</td> <td>Kom Kom bods</td> <td>Early Conomanian</td> <td>21.5</td> <td>26.0</td> <td></td> <td>This study</td>	M TA 017	Tooth on amal	Carcharodontosaurus sabaricus	Takomout	Morocco	Kom Kom bods	Early Conomanian	21.5	26.0		This study
In-TA-025Toolf enamelChromotoniosaurus sundruktisTakemoutMoroccoKem Kem bedsEarly Cenomanian15.920.5-9.1This StudyM-TA-027Tooth enamelSpinosaurusTakemoutMoroccoKem Kem bedsEarly Cenomanian18.225.5-9.8This studyM-TA-028Tooth enamelSpinosaurusTakemoutMoroccoKem Kem bedsEarly Cenomanian18.425.7-9.9This studyM-TA-021Tooth bulkAnhanguerid pterosaurTakemoutMoroccoKem Kem bedsEarly Cenomanian19.425.1-7.7This studyM-TA-021bisTooth bulkAnhanguerid pterosaurTakemoutMoroccoKem Kem bedsEarly Cenomanian19.425.1-7.9This studyM-TA-030Tooth bulkAnhanguerid pterosaurTakemoutMoroccoKem Kem bedsEarly Cenomanian19.425.1-9.1This studyM-TA-030Tooth bulkCrocodyliaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.223.9-9.1This studyM-TA-030Shell boneCheloniaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-TA-040OsteodermCrocodyliaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-TA-05Shell boneCheloniaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-1	M TA 010	Tooth on amal	Carcharodontosaurus saharicus	Takemout	Morocco	Kem Kem bods	Early Conomanian	20.1	20.0	- 9.0	This study
M-174-027Toold relatingSpinosaurusTakemoutMotoccoKein Kein bedsEarly Cenomanian18.020.1-10.7This studyM-174-027Tooth enamelSpinosaurusTakemoutMoroccoKem Kem bedsEarly Cenomanian18.425.7-9.9This studyM-174-027Tooth enamelSpinosaurusTakemoutMoroccoKem Kem bedsEarly Cenomanian19.724.9-7.7This studyM-174-021bisTooth bulkAnhanguerid pterosaurTakemoutMoroccoKem Kem bedsEarly Cenomanian19.324.7-10.2This studyM-174-021bisTooth bulkAnhanguerid pterosaurTakemoutMoroccoKem Kem bedsEarly Cenomanian19.324.7-10.2This studyM-174-021 bisTooth bulkAnhanguerid pterosaurTakemoutMoroccoKem Kem bedsEarly Cenomanian19.324.7-10.2This studyM-174-003Tooth bulkCrocodyliaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.223.9-9.1This studyM-174-004OsteodermCrocodyliaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-174-005Shell boneCheloniaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-174-010Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.0	M TA 026	Tooth enamel	Curcharouonitosaurus sanaricus	Takemout	Morocco	Kem Kem beds	Early Cenomanian	19.9	20.5	- 9.1	This study
InclusionInderindual prinosaurusJakenioutMoroccoKein Kein bedsLarly Ceinomanian16.225.3-9.8This studyM-TA-021Tooth bulkSpinosaurusTakemoutMoroccoKein Kein bedsEarly Cenomanian19.724.9-7.7This studyM-TA-020bisTooth bulkAnhanguerid pterosaurTakemoutMoroccoKein Kein bedsEarly Cenomanian19.324.7-10.2This studyM-TA-021bisTooth bulkAnhanguerid pterosaurTakemoutMoroccoKein Kein bedsEarly Cenomanian19.324.7-10.2This studyM-TA-030Tooth bulkCrocodyliaTakemoutMoroccoKein Kein bedsEarly Cenomanian19.223.9-9.1This studyM-TA-040OsteodermCrocodyliaTakemoutMoroccoKein Kein bedsEarly Cenomanian19.225.5-9.6This studyM-TA-004OsteodermCrocodyliaTakemoutMoroccoKein Kein bedsEarly Cenomanian19.025.1-10.7This studyM-TA-011Shell boneCheloniaTakemoutMoroccoKein Kein bedsEarly Cenomanian19.025.1-10.7This studyM-JQ-008Tooth bulkTheropodaJebel Al QablaMoroccoKein Kein bedsEarly Cenomanian19.025.1-10.7This studyM-JQ-010Tooth bulkTheropodaJebel Al QablaMoroccoKein Kein bedsEarly Cenomanian19.022.4	M TA 027	Tooth on amal	Spinosaurus	Takemout	Morocco	Kem Kem bods	Early Conomanian	10.0	20.1	- 10.7	This study
In-FA-02bToole FindnerSpinosaurusTakemoutMoroccoKeni Keni bedsEarly Cenomanian19.724.9-7.7This studyM-TA-021bisTooth bulkAnhanguerid pterosaurTakemoutMoroccoKem Kem bedsEarly Cenomanian19.324.7-10.2This studyM-TA-021bisTooth bulkAnhanguerid pterosaurTakemoutMoroccoKem Kem bedsEarly Cenomanian19.324.7-10.2This studyM-TA-030Tooth bulkCrocodyliaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.223.9-9.1This studyM-TA-040OsteodermCrocodyliaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-TA-030Shell boneCheloniaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-TA-041Shell boneCheloniaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-JQ-008Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.322.9-11.3This studyM-JQ-010Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.923.8-11.2This studyM-JQ-038Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.923.8-11.2	M TA 029	Tooth enamel	Spinosaurus	Takemout	Morocco	Kem Kem beds	Early Cenomanian	10.2	25.5	- 9.8	This study
In-TA-02DisTooth bulkJoinsduringTakemoutMoroccoKein Kein bedsEarly Cenomanian19.724.9-7.7This studyM-TA-02DisTooth bulkAnhanguerid pterosaurTakemoutMoroccoKem Kem bedsEarly Cenomanian19.324.7-10.2This studyM-TA-02DisTooth bulkCrocodyliaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.223.9-9.1This studyM-TA-030Tooth bulkCrocodyliaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.223.9-9.6This studyM-TA-040OstodermCrocodyliaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-TA-010Shell boneCheloniaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-TA-040StoedermCheloniaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-JQ-008Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.322.8-8.3This studyM-JQ-010Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.923.8-11.2This studyM-JQ-038Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.2-11.4This study	M TA 021	Tooth bulk	Spinosaurus	Takemout	Morocco	Kem Kem bods	Early Conomanian	10.4	23.7	- 9.9	This study
M-TA-021bisTooth bulkAnhangucht phrtsaulFakemoutMoroccoKem Kem bedsEarly Cenomanian19.424.7-1.0.2This studyM-TA-021bisTooth bulkCrocodyliaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.324.7-10.2This studyM-TA-040OsteodermCrocodyliaTakemoutMoroccoKem Kem bedsEarly Cenomanian20.525.5-9.6This studyM-TA-011Shell boneCheloniaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-TA-011Shell boneCheloniaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-9.6This studyM-JQ-008Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.322.9-11.3This studyM-JQ-010Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.923.8-11.2This studyM-JQ-038Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.924.6-9.7This studyM-JQ-039Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.324.0-11.4This studyM-JQ-040Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.324.0-11.4<	M_TA_020bis	Tooth bulk	Anhanguerid pterosaur	Takemout	Morocco	Kem Kem beds	Early Cenomanian	10.7	24.5	-7.9	This study
M-TA-021 DisToth bulkFind age the periodTakemoutMoroccoKem Kem bedsEarly Cenomanian19.223.9-9.1This studyM-TA-030Toth bulkCrocodyliaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.223.9-9.1This studyM-TA-040OsteodermCrocodyliaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-TA-011Shell boneCheloniaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.724.8-8.3This studyM-JQ-008Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.8-9.0This studyM-JQ-009Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.923.8-11.2This studyM-JQ-038Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.923.8-11.2This studyM-JQ-039Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.6-9.7This studyM-JQ-040Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.2-11.4This studyM-JQ-0404Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.325.3-8.5 </td <td>M-TA-020013</td> <td>Tooth bulk</td> <td>Anhanguerid pterosaur</td> <td>Takemout</td> <td>Morocco</td> <td>Kem Kem beds</td> <td>Early Cenomanian</td> <td>10.3</td> <td>23.1</td> <td>- 10.2</td> <td>This study</td>	M-TA-020013	Tooth bulk	Anhanguerid pterosaur	Takemout	Morocco	Kem Kem beds	Early Cenomanian	10.3	23.1	- 10.2	This study
M-TR-050Tooth bulkCrocodyliaTakemoutMoroccoKem Kem bedsEarly Cenomanian15.225.5-9.6This studyM-TA-009Shell boneCheloniaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-TA-011Shell boneCheloniaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.724.8-8.3This studyM-TA-011Shell boneCheloniaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.322.9-11.3This studyM-JQ-008Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.322.9-11.3This studyM-JQ-010Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.923.8-11.2This studyM-JQ-038Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.324.6-9.7This studyM-JQ-039Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.324.6-9.7This studyM-JQ-039Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.324.6-9.7This studyM-JQ-039Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.324.6-9.7This	M_TA_030	Tooth bulk	Crocodylia	Takemout	Morocco	Kem Kem beds	Early Cenomanian	10.2	24.7	_ 0.1	This study
M-TA-OOFOstCott IIICrocotylaTakemottMorecoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-TA-011Shell boneCheloniaTakemoutMorecoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-TA-011Shell boneCheloniaTakemoutMorecoKem Kem bedsEarly Cenomanian19.025.1-10.7This studyM-JQ-008Tooth bulkTheropodaJebel Al QablaMorecoKem Kem bedsEarly Cenomanian19.322.9-11.3This studyM-JQ-009Tooth bulkTheropodaJebel Al QablaMorecoKem Kem bedsEarly Cenomanian19.322.9-11.2This studyM-JQ-010Tooth bulkTheropodaJebel Al QablaMorecoKem Kem bedsEarly Cenomanian19.923.8-11.2This studyM-JQ-038Tooth enamelSpinosaurusJebel Al QablaMorecoKem Kem bedsEarly Cenomanian19.324.0-11.4This studyM-JQ-040Tooth enamelSpinosaurusJebel Al QablaMorecoKem Kem bedsEarly Cenomanian19.724.2-11.2This studyM-JQ-0404Tooth bulkAnhanguerid pterosaurJebel Al QablaMorecoKem Kem bedsEarly Cenomanian19.325.3-8.5This studyM-JQ-040ATooth bulkCrocodyliaJebel Al QablaMorecoKem Kem bedsEarly Cenomanian19.325.3-8.5	M_TA_004	Osteoderm	Crocodylia	Takemout	Morocco	Kem Kem beds	Early Cenomanian	20.5	25.5	-96	This study
M-TA-051Shich bolicCheloniaTakemoutMoroccoKem Kem bedsEarly Cenomanian19.724.8- 10.7This studyM-JQ-008Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.322.9- 11.3This studyM-JQ-009Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.8- 8.3This studyM-JQ-009Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.923.8- 11.2This studyM-JQ-010Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.923.8- 11.2This studyM-JQ-038Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.324.0- 11.4This studyM-JQ-040Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.2- 11.2This studyM-JQ-0404Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.2- 11.2This studyM-JQ-0404Tooth bulkAnhanguerid pterosaurJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.325.3- 8.5This studyM-JQ-0408Tooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian <td< td=""><td>M_TA_009</td><td>Shell bone</td><td>Chelonia</td><td>Takemout</td><td>Morocco</td><td>Kem Kem beds</td><td>Early Cenomanian</td><td>10.0</td><td>25.5</td><td>- 10.7</td><td>This study</td></td<>	M_TA_009	Shell bone	Chelonia	Takemout	Morocco	Kem Kem beds	Early Cenomanian	10.0	25.5	- 10.7	This study
M-HornJinch bolkTheropolaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.8-0.5-0.5This studyM-JQ-008Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.322.9-11.3This studyM-JQ-010Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.322.9-9.0This studyM-JQ-010Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.923.8-11.2This studyM-JQ-038Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.324.6-9.7This studyM-JQ-039Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.324.0-11.4This studyM-JQ-040Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.2-11.2This studyM-JQ-0404Tooth bulkAnhanguerid pterosaurJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.2-11.2This studyM-JQ-040ATooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.325.3-8.5This studyM-JQ-040ATooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian<	M_TA_011	Shell bone	Chelonia	Takemout	Morocco	Kem Kem beds	Early Cenomanian	10.7	23.1	- 83	This study
M-JQ-000Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian10.522.5-11.5This studyM-JQ-001Tooth bulkTheropodaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.923.8-11.2This studyM-JQ-038Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.923.8-11.2This studyM-JQ-039Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.324.0-11.4This studyM-JQ-040Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.2-11.2This studyM-JQ-040Tooth bulkAnhanguerid pterosaurJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.2-11.2This studyM-JQ-040ATooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.2-11.2This studyM-JQ-040ATooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.325.3-8.5This studyM-JQ-040ATooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.325.2-9.8This studyM-JQ-040BTooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian <t< td=""><td>M-IO-008</td><td>Tooth bulk</td><td>Theropoda</td><td>Iebel Al Oabla</td><td>Morocco</td><td>Kem Kem beds</td><td>Farly Cenomanian</td><td>19.7</td><td>24.0</td><td>- 11 3</td><td>This study</td></t<>	M-IO-008	Tooth bulk	Theropoda	Iebel Al Oabla	Morocco	Kem Kem beds	Farly Cenomanian	19.7	24.0	- 11 3	This study
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M-JQ-038Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian18.924.6-9.7This studyM-JQ-039Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.324.0-11.4This studyM-JQ-040Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.324.0-11.4This studyM-JQ-040Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.2-11.2This studyM-JQ-040ATooth bulkAnhanguerid pterosaurJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.325.3-8.5This studyM-JQ-040ATooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.325.3-8.5This studyM-JQ-040BTooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.325.2-9.8This studyM-JQ-040BTooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian18.724.7-9.3This studyM-JQ-0407Tooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian18.724.7-9.3This studyM-JQ-0407Tooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian <td>M-IO-010</td> <td>Tooth bulk</td> <td>Theropoda</td> <td>Jebel Al Oabla</td> <td>Morocco</td> <td>Kem Kem beds</td> <td>Farly Cenomanian</td> <td>19.9</td> <td>24.5</td> <td>- 11 2</td> <td>This study</td>	M-IO-010	Tooth bulk	Theropoda	Jebel Al Oabla	Morocco	Kem Kem beds	Farly Cenomanian	19.9	24.5	- 11 2	This study
M-JQ-039Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.324.0-11.4This studyM-JQ-040Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.2-11.2This studyM-JQ-040Tooth bulkAnhanguerid pterosaurJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.2-11.2This studyM-JQ-040ATooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.325.3-8.5This studyM-JQ-040BTooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.325.2-9.8This studyM-JQ-040BTooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian18.724.7-9.3This studyM-JQ-0407Tooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian18.724.7-9.3This studyM-JQ-0407Tooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian18.724.7-9.3This studyM-JQ-0407Tooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian17.923.3-10.0This study	M-IO-038	Tooth enamel	Spinosaurus	Jebel Al Oabla	Morocco	Kem Kem beds	Farly Cenomanian	18.9	24.6	-97	This study
M-jQ-040Tooth enamelSpinosaurusJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.2-11.2This studyM-jQ-001bisTooth bulkAnhanguerid pterosaurJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.724.2-11.2This studyM-jQ-001bisTooth bulkAnhanguerid pterosaurJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian20.926.5-10.1This studyM-jQ-040ATooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian19.325.3-8.5This studyM-jQ-040BTooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian18.925.2-9.8This studyM-jQ-040CTooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian18.724.7-9.3This studyM-jQ-040CTooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian18.724.7-9.3This studyM-jQ-040CTooth bulkCrocodyliaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian18.724.7-9.3This studyM-jQ-017Shell boneCheloniaJebel Al QablaMoroccoKem Kem bedsEarly Cenomanian17.923.3-10.0This study	M-IO-039	Tooth enamel	Spinosaurus	Jebel Al Oabla	Morocco	Kem Kem beds	Early Cenomanian	19.3	24.0	-114	This study
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M-jQ-040A Tooth bulk Crocodylia Jebel Al Qabla Morocco Kem Kem beds Early Cenomanian 20.5 20.5 10.1 This study M-jQ-040A Tooth bulk Crocodylia Jebel Al Qabla Morocco Kem Kem beds Early Cenomanian 19.3 25.3 - 8.5 This study M-jQ-040B Tooth bulk Crocodylia Jebel Al Qabla Morocco Kem Kem beds Early Cenomanian 18.9 25.2 - 9.8 This study M-jQ-040C Tooth bulk Crocodylia Jebel Al Qabla Morocco Kem Kem beds Early Cenomanian 18.9 25.2 - 9.8 This study M-jQ-040C Tooth bulk Crocodylia Jebel Al Qabla Morocco Kem Kem beds Early Cenomanian 18.7 24.7 - 9.3 This study M-jQ-017 Shell bone Chelonia Jebel Al Qabla Morocco Kem Kem beds Early Cenomanian 17.9 23.3 - 10.0 This study	M-IO-001his	Tooth bulk	Anhanguerid pterosaur	Jebel Al Oabla	Morocco	Kem Kem heds	Early Cenomanian	20.9	26.5	- 10 1	This study
M-JQ-040B Tooth bulk Crocodylia Jebel Al Qabla Morocco Kem Kem beds Early Cenomanian 18.9 25.2 -9.8 This study M-JQ-040C Tooth bulk Crocodylia Jebel Al Qabla Morocco Kem Kem beds Early Cenomanian 18.9 25.2 -9.8 This study M-JQ-040C Tooth bulk Crocodylia Jebel Al Qabla Morocco Kem Kem beds Early Cenomanian 18.7 24.7 -9.3 This study M-JQ-0407 Shell bone Chelonia Jebel Al Qabla Morocco Kem Kem beds Early Cenomanian 17.9 23.3 -10.0 This study	M-IO-040A	Tooth bulk	Crocodylia	Jebel Al Oabla	Morocco	Kem Kem beds	Early Cenomanian	19.3	25.3	-85	This study
M-JQ-040C Tooth bulk Crocodylia Jebel Al Qabla Morocco Kem Kem beds Early Cenomanian 18.7 24.7 -9.3 This study M-JQ-017 Shell bone Chelonia Jebel Al Qabla Morocco Kem Kem beds Early Cenomanian 17.9 23.3 -10.0 This study	M-IO-040B	Tooth bulk	Crocodylia	Jebel Al Oabla	Morocco	Kem Kem beds	Early Cenomanian	18.9	25.2	-9.8	This study
M-IO-017 Shell bone Chelonia lebel Al Qabla Morocco Kem Kem beds Early Cenomanian 17.9 23.3 – 10.0 This study	M-IO-040C	Tooth bulk	Crocodylia	Jebel Al Oabla	Morocco	Kem Kem beds	Early Cenomanian	18.7	24.7	-9.3	This study
	M-JQ-017	Shell bone	Chelonia	Jebel Al Qabla	Morocco	Kem Kem beds	Early Cenomanian	17.9	23.3	- 10.0	This study

(continued on next page) 43

Table 1 (continued)	
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Sample no.	Skeletal tissue	Taxon	Locality name	Country	Formation	Age	$\delta^{18} O_P$	$\delta^{18}O_C$	$\delta^{13} C_{C}$	Data origin
							(% V-SMOW)	(‰ V-SMOW)	(‰ V-PDB)	
M-JQ-018	Shell bone	Chelonia	Jebel Al Qabla	Morocco	Kem Kem beds	Early Cenomanian	18.8	23.5	-9.5	This study
M-JQ-019	Shell bone	Chelonia	Jebel Al Qabla	Morocco	Kem Kem beds	Early Cenomanian	18.7	24.0	- 9.5	This study
M-JQ-020	Tooth bulk	Neoceratodus africanus	Jebel Al Qabla	Morocco	Kem Kem beds	Early Cenomanian	20.2	24.9	-11.1	This study
M-JQ-036A	Scale ganoine	Stromerichthys	Jebel Al Qabla	Morocco	Kem Kem beds	Early Cenomanian	20.4	25.3	-2.7	This study
M-JQ-036B	Scale ganoine	Stromerichthys	Jebel Al Qabla	Morocco	Kem Kem beds	Early Cenomanian	20.1	23.8	- 3.5	This study
M-JQ-036C	Scale ganoine	Stromerichthys	Jebel Al Qabla	Morocco	Kem Kem beds	Early Cenomanian	19.2	24.7	- 5.7	This study
M-ZA-015	Tooth bulk	Theropoda	Zaouia	Morocco	Kem Kem beds	Early Cenomanian	19.8	24.2	- 10.7	This study
M-ZA-010	Tooth enamel	Anhanguerid pterosaur	Zaouia	Morocco	Kem Kem beds	Early Cenomanian	22.9	-	-	This study
M-ZA-005	Osteoderm	Crocodylia	Zaouia	Morocco	Kem Kem beds	Early Cenomanian	19.7	24.0	- 12.0	This study
M-ZA-007	Shell bone	Chelonia	Zaouia	Morocco	Kem Kem beds	Early Cenomanian	18.2	23.3	-10.3	This study
M-ZA-021A	Rostral tooth bulk	Onchopristis numidus	Zaouia	Morocco	Kem Kem beds	Early Cenomanian	19.6	23.4	-8.2	This study
M-ZA-021B	Rostral tooth bulk	Onchopristis numidus	Zaouia	Morocco	Kem Kem beds	Early Cenomanian	18.8	23.0	-6.2	This study
M-ZA-021C	Rostral tooth bulk	Onchopristis numidus	Zaouia	Morocco	Kem Kem beds	Early Cenomanian	18.8	24.8	- 8.7	This study
M-ZA-021D	Rostral tooth bulk	Onchopristis numidus	Zaouia	Morocco	Kem Kem beds	Early Cenomanian	19.6	23.1	-6.8	This study
TU-001	Tooth enamel	Theropoda	Bateun el Hmaima	Tunisia	Chenini Member	Late Aptian–Early Albian	20.0	-	-	Amiot et al. (2006)
TU-002	Tooth enamel	Theropoda	Bateun el Hmaima	Tunisia	Chenini Member	Late Aptian–Early Albian	21.2	-	-	Amiot et al. (2006)
TU-003	Tooth enamel	Theropoda	Bateun el Hmaima	Tunisia	Chenini Member	Late Aptian–Early Albian	20.3	-	-	Amiot et al. (2006)
TU-004	Tooth enamel	cf. Spinosaurus	Bateun el Hmaima	Tunisia	Chenini Member	Late Aptian–Early Albian	22.2	-	-	Amiot et al. (2006)
TU-006	Tooth enamel	cf. Spinosaurus	Bateun el Hmaima	Tunisia	Chenini Member	Late Aptian–Early Albian	19.8	-	-	Amiot et al. (2006)
TU-005	Tooth enamel	cf. Spinosaurus	Bateun el Hmaima	Tunisia	Chenini Member	Late Aptian–Early Albian	16.4	-	-	Amiot et al. (2006)
TU-007	Tooth enamel	Crocodylia	Bateun el Hmaima	Tunisia	Chenini Member	Late Aptian-Early Albian	18.2	-	-	Amiot et al. (2006)
TU-008	Tooth enamel	Crocodylia	Bateun el Hmaima	Tunisia	Chenini Member	Late Aptian-Early Albian	18.4	-	-	Amiot et al. (2006)

Table 2

Locality name, paleolatitude and reconstructed environmental water $\delta^{18}O_w$ values calculated using the Amiot et al. (2007) and Barrick et al. (1999) oxygen isotope fractionation equations.

Locality	Paleolatitude			$\delta^{18}O_{water}$ (% V-SMOW)							
	mean	+	_	From crocodilians	From turtles	Average	Std. err.				
Laje do Coringa	- 5.0	1.7	11.9	-3.2	-	- 3.2	1.0				
Araripe Basin	-10.2	8.3	1.6	_	-2.4	-2.4	1.0				
Bou Laalou	12.4	3.5	3.3	-4.7	- 5.2	- 5.0	1.0				
Khetitila Srhira	16.5	2.1	2.2	-4.6	-5.4	-5.0	1.0				
Chaaft	16.5	2.1	2.2	-3.0	-3.4	-3.2	1.0				
Takemout	16.5	2.1	2.2	-2.8	-2.8	-2.8	1.0				
Jebel Al Qabla	16.5	2.1	2.2	-3.5	-3.6	-3.6	1.0				
Zaouia	16.5	2.1	2.2	-2.9	-4.0	-3.4	1.0				
Bateun el Hmaima	16.5	2.1	2.2	-4.1	-	-4.1	1.0				

5. Discussion

5.1. Preservation of pristine isotope compositions of fossil apatites

Secondary precipitation of apatite and isotopic exchange during microbially-mediated reactions may alter the primary isotopic signal (Blake et al., 1997; Zazzo et al., 2004a). However, apatite crystals that make up tooth enamel or fish scale ganoine are large and densely packed, and isotopic exchange under inorganic conditions has little effect on the oxygen isotope composition of phosphates even at geological time scales (Kolodny et al., 1983; Lécuyer et al., 1999a). Turtle shell bone should be more susceptible to diagenetic alteration because hydroxylapatite crystals of bones are smaller and less densely intergrown than those of enamels (Kolodny et al., 1996), even though several case studies have shown that the original oxygen isotope composition seems to be preserved in Mesozoic and Cenozoic turtle remains (Barrick et al., 1999; Amiot et al., 2004, 2006; Matson and Fox, 2008; Amiot et al., 2009). Although no method is available to demonstrate definitely whether or not the oxygen isotope composition of fossil vertebrate phosphate was modified by diagenetic processes, several ways to assess the preservation state of the primary isotopic



Fig. 2. Mean oxygen isotope compositions of apatite phosphate ($\delta^{18}O_P$) from studied reptile taxa plotted against calculated environmental water $\delta^{18}O_w$ values. Numbers refer to the following localities: 1: Khetitila Srhira (Morocco); 2: Bou Laalou (Morocco); 3: Bateun el Hmaima (Tunisia); 4: Jebel Al Qabla (Morocco); 5: Zaouia (Morocco); 6: Chaaft (Morocco); 7: Laje do Coringa (Brazil); 8: Takemout (Morocco); 9: Araripe Basin (Brazil).

record have been proposed (e.g. lacumin et al., 1996; Kolodny et al., 1996; Fricke et al., 1998; Lécuyer et al., 2003; Pucéat et al., 2004; Zazzo et al., 2004b; Tütken et al., 2008). Here, the main argument supporting the preservation of the original oxygen isotope composition is the systematic offset observed between dinosaurs and ectothermic reptiles (turtles and crocodilians), most likely resulting from differences in ecology and physiology. The present dataset illustrates systematic offsets between semi-aquatic animals (turtles and crocodilians) and terrestrial ones (dinosaurs, except Spinosaurus), the latter having $\delta^{18}O_n$ values up to 2.7‰ higher than the values of crocodilians and turtles. If early diagenetic processes had occurred, they would have tended to homogenise $\delta^{18}O_p$ values of all vertebrate remains whatever the physiology and ecology of the corresponding taxa (Lécuyer et al., 2003). The second clue which favours at least a partial original preservation of both phosphate and carbonate values is the correlation between their two δ^{18} O values which roughly parallels the oxygen isotope fractionation line established for extant mammals (Iacumin et al., 1996; Zazzo et al., 2004b; Fig. 3). It is noteworthy that the large data scattering observed within Araripe Basin remains may indicate a partial modification of the pristine oxygen isotope compositions of apatite carbonate, which must therefore be considered with caution. Primary preservation of carbonate $\delta^{13}C_c$ values of apatite is more difficult to test. Here we assume an original preservation based on consistent value differences observed between taxonomic groups, and the low scattering of $\delta^{13}C_c$ values within each taxon compared to the inter-taxon variability. Indeed, mean $\delta^{13}C_c$ values range from -10.7% (spinosaurs) to -2.1% (*Neoproscinetes*) whereas standard deviations within most groups are of about $\pm 1.0\%$ with a maximum of $\pm 2.3\%$ for the ubiquitous fish Lepidotes.

5.2. Meaning of the oxygen and carbon isotope compositions of phosphate and carbonate

5.2.1. Crocodilians and turtles

Crocodilians and turtles from studied localities are identified as being semi-aquatic to aquatic species. Due to higher water turnover and lower transcutaneous water evaporation compared to terrestrial vertebrates, crocodilian and turtle $\delta^{18}O_P$ values are lower than those of coexisting terrestrial dinosaurs and pterosaurs, with two exceptions in Takemout and Zaouia localities where some crocodilian $\delta^{18}O_P$ values are close to theropod dinosaur ones. It is also noteworthy that $\delta^{18}O_P$ values of crocodilians are systematically higher than those of turtle for any given locality. This might be accounted for peculiar behaviour by crocodilians such as extensive basking in the sun or the fortuitous presence of terrestrial crocodilian remains within analyzed Moroccan samples such as *Hamadasuchus rebouli* (Buffetaut, 1994), sometimes difficult to identify from isolated osteoderms or anterior teeth. Although being ectothermic animals, and thus having body temperatures fluctuating along with environmental ones (e.g. Seebacher et al.,



Fig. 3. Mean oxygen isotope compositions of apatite phosphate ($\delta^{18}O_p$) from studied vertebrate taxa plotted against their corresponding oxygen isotope composition of carbonate ($\delta^{18}O_c$), along with the phosphate–carbonate isotopic fractionation line established for extant herbivorous mammals (lacumin et al., 1996).

2003), crocodilians and turtles synthesize bone and tooth phosphate under relatively narrow temperature ranges (Barrick et al., 1999; Amiot et al., 2007). Oxygen isotope compositions of environmental waters were estimated using empirical phosphate–water fractionation equations established for crocodilians and turtles (Barrick et al., 1999; Amiot et al., 2007). Calculated δ^{18} O values of environmental waters range from -5% to -2.4%, a range corresponding to continental surface water values observed nowadays at tropical to equatorial latitudes (IAEA/WMO, 2006). Carbon isotope compositions of both crocodilians and turtles range from -12.0% to -9.2%. It is noteworthy that these values



Fig. 4. Mean oxygen isotope compositions of apatite phosphate ($\delta^{18}O_p$) from studied vertebrate taxa plotted against their corresponding carbon isotope composition of carbonate ($\delta^{13}C_c$).

fall within the larger range of -18.1% to -4.1% measured in extant crocodilians belonging to five different genera (Lécuyer et al., 2003; Stanton, 2006). Assuming a carbon isotope fractionation of 9% between crocodilian apatite and diet (a value calculated for carnivorous mammals; Lee-Thorp et al., 1989; Tieszen and Fagre, 1993), Stanton (2006) differentiated crocodilians feeding on preys based on C₄, C₃ and mixed plant foodwebs, highest $\delta^{13}C_c$ values corresponding to pure C₄based diets. Diet of fossil turtles is difficult to assess from osteological features. Carbon isotope fractionations of 6-9% calculated between carnivorous turtle apatites and their diets, and of 12‰ between herbivorous ones and their diets have been proposed for extant marine turtles (Biasatti, 2004). Such isotopic fractionation values for both turtles and crocodiles indicate δ^{13} C values for their diet around -20%, compatible with C₃-based foodwebs. Interestingly, the Brazilian turtle Araripemys, often considered as a marine turtle due to its flattened shell and its abundant presence in the saltwater deposits of the Romualdo member of the Santana Formation (e.g. Price, 1973; De Oliveira and Kellner, 2005), has both low $\delta^{18}O_P$ and $\delta^{13}C_c$ values more compatible with freshwater affinities. As only one sample was available for this study, definitive conclusions cannot be drawn, and a larger sample set must be analyzed in order to better understand the living environment of Araripemys.

5.2.2. Dinosaurs

Theropod dinosaurs (except spinosaurs which are discussed below) were terrestrial homeotherm carnivorous dinosaur with an endotherm-like thermophysiology (Barrick and Showers, 1994; Fricke and Rogers, 2000; Amiot et al., 2006). As homeotherms and obligate drinkers, variations in their oxygen isotope compositions of phosphate and carbonate predominantly reflect variations in the drinking waters which overwhelm oxygen input from other sources such as food (Barrick and Showers, 1995; Barrick et al., 1996; Straight et al., 2004).

Spinosaurs were theropod dinosaurs with peculiar morphologies, the most striking features being their almost conical, crocodile-like teeth, elongated and narrow snout and their extremely tall vertebral neural spines indicating the presence of a dorsal sail. From these anatomical characteristics, these animals were supposed to be piscivorous, feeding on fish the way crocodilians do. The discovery of partially digested fish scales in the stomach region of a nearly complete skeleton of *Baryonyx* supported the hypothesis that these theropods ate preferentially fish (Charig and Milner, 1997), but not exclusively, as several direct evidence indicate that some other dinosaurs (Charig and Milner, 1997) and pterosaurs (Buffetaut et al., 2004) were also part of their diet. Although anatomical features indicate that spinosaurs were terrestrial and bipedal dinosaurs such as other theropods, it has been demonstrated that their $\delta^{18}O_P$ values are lower than the ones of coexisting terrestrial theropods and values within the range of crocodilians were most likely due to a semiaquatic lifestyle consisting in spending most of daily time immersed (Amiot et al., 2009, 2010). Compared to other spinosaurs (including the ones from the Brazilian site of Laje do Coringa), those recovered from Morocco and Tunisia may have had a more complicated lifestyle, as their $\delta^{18}O_P$ values range from semi-aquatic reptile to terrestrial theropod ones. It is possible that peculiar environmental or ecological stresses forced them to alternate between semi-aquatic and terrestrial lifestyles (Amiot et al., 2010). It is noteworthy that the higher the $\delta^{18}O_w$ values of surface waters, the lower the spinosaur $\delta^{18}O_P$ values (Fig. 2). This may suggest that under dryer conditions, which are inferred from higher δ^{18} O_w values, spinosaurs would spend more time in water to circumvent drought-related stress. Titanosaurid sauropods from Morocco have $\delta^{18}O_P$ values similar to those of theropod dinosaurs, which strongly suggest that most of their oxygen input was from drinking water. If they had mostly relied on plant water, usually ¹⁸O-enriched by several per mils relative to surface waters, their $\delta^{18}O_P$ values would have been significantly higher than those of coexisting theropods.

Studied dinosaurs have a restricted range in carbon isotope compositions from -12% to -7%, which is expected for animals feeding on C₃ plant-based foodwebs. By comparing mean $\delta^{13}C_c$ values of both dinosaur tooth enamels and associated bulk sedimentary organic matter or fossil plant leaves, ¹³C-enrichments of 18‰ have been proposed for hadrosaurs and ceratopsians (Fricke and Pearson, 2008), and 16‰ for sauropods (Tütken, in press). These ¹³C-enrichments that are 3‰ to 8‰ higher than those of extant herbivorous mammals may account for distinct digestive physiologies including the incorporation of specific plant compounds and higher production of methane from the gastrointestinal tract of herbivorous dinosaurs. Indeed, the rate of ¹²C-rich methane production and its loss during digestion and rumination can have an important influence on the enamel-diet ¹³C-enrichment factor (Passey et al., 2005). Tütken (in press) also noted that the enamel-diet ¹³C-enrichment of sauropod dinosaurs is similar to the bone-diet ¹³C-enrichment of extant ostriches, both animals belonging to the same dinosaur lineage (Saurischia). Titanosaurid sauropods from Morocco have a mean $\delta^{13}C_c$ value of $-10.3 \pm 0.8\%$, suggesting that they were feeding on plants having a δ^{13} C value of about -26%, falling within the C₃ plant range.

5.2.3. Pterosaurs

Studied Moroccan and Brazilian pterosaur teeth belong to the family Anhangueridae (Wang et al., 2009). According to their skull and tooth morphologies, these flying reptiles are considered to be fish eaters, fishing on the wing by seizing prey with their pointed jaws while flying above water surface (Campos and Kellner, 1985; Wellnhofer, 1991; Veldmeijer et al., 2007). As expected, their high $\delta^{18}O_P$ values are within the range of other terrestrial animals. Moreover, the large range of $\delta^{18}O_P$ values (from 18.3‰ to 22.9‰) suggests that they drank waters from different sources, including main river streams, evaporated water ponds and possibly marine waters while feeding. Moreover, their low and restricted $\delta^{13}C_c$ values ranging from -11.3% to -7.5% point to a diet with at least an important part of freshwater fish, as marine fish usually have more positive and variable $\delta^{13}C_c$ values (see discussion about fish $\delta^{13}C$ values below). However, as bulk teeth (enamel + dentine) only have been analyzed for their carbon isotope compositions, the possibility that diagenetic alteration is responsible for such δ^{13} C values cannot be totally excluded, dentine being more prone to such an alteration than enamel.

5.2.4. Fish

Early Cenomanian fish from Morocco and Brazil have high $\delta^{18}O_P$ values indicating that their phosphate precipitated from ¹⁸O-enriched waters relative to those calculated from isotopic compositions of coexisting crocodilians and freshwater turtles. Indeed, the use of the fish phosphate-water temperature scale (Kolodny et al., 1983) gives anomalously low precipitation temperatures ranging from 3 °C to 13 °C assuming that fish were living in waters with $\tilde{\delta}^{18}O_w$ values of -5% to -2.4% as deduced from crocodilians and turtle $\delta^{18}O_P$ values. Therefore these fish $\delta^{18}O_P$ values suggest distinct aquatic environments. The euryhaline sawfish Onchopristis numidus, the semionotiform Lepidotes and the coelacanth Mawsonia might have shared their living environments between fresh and shallow marine waters. A similar lifestyle can be proposed for the poorly known Stromerichthys, which was also found in Cenomanian continental deposits of France (Vullo and Neraudeau, 2008), a geographic distribution suggesting its ability to venture in the open ocean. Therefore, we propose that their $\delta^{18} O_P$ values reflect seawater $\delta^{18} O_{sw}$ values (between -1% and +1%) at temperatures ranging from 21 ± 6 °C (including the value of 16 °C inferred from one *Onchopristis* sample) to 34 ± 2 °C (Table 3), which is in good agreement with calculated Cenomanian sea surface temperatures of Western Tethys (Pucéat et al., 2003). Their $\delta^{13}C_c$

Ma	rine wat	er temperature:	s estimated fr	om the	δ18Op	values	of fish	using	the equation	of Kolo	odny et al	. (1983)).
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Locality	Mawsonia		Neoproscinetes		Vinctifer		Lepidotes			Onchopristis			Stromerichthys					
$\delta^{18}O_{water}$	-1	1		-1	1		-1	1		-1	1		-1	1		-1	1	
(‰ V-SMOW)	T°C		Std. err.	T°C		Std. err.	T°C		Std. err.	T°C		Std. err.	T°C		Std. err.	T°C		Std. err.
Laje do Coringa Araripe Basin Khetitila Srhira Chaaft Jebel Al Qabla Zaouia	23	32	3	11	20	1	22	30	-	21	30	6	16 25	24 34	- 2	22	31	2

values, higher than those of other continental vertebrates may derive from a carbon of marine origin such as dissolved inorganic carbon and organic carbon, usually having higher δ^{13} C values than freshwater carbon sources (Clayton and Degens, 1959). The two Moroccan lungfishes Protopterus and Neoceratodus, as well as the indeterminate ceratodontiform from Laje do Coringa (Brazil) also have high $\delta^{18}O_{n}$ values. As marine living environments for these fishes seem unlikely considering that fossil lungfish have been found quite exclusively in continental deposits since the Triassic (Cavin et al., 2007), it seems that their high $\delta^{18}O_P$ values might be due to temporary drought of their living environment, or to estivation behaviour. Indeed, African and South American lungfish are capable of surviving seasonal drying out of their habitats by burrowing into mud and estivating throughout the dry season (Fishman et al., 1986). According to this, they might have precipitated some parts of their tooth phosphate in equilibrium with ¹⁸O-enriched evaporated waters. Assuming mean temperatures of continental environments in the 30-35 °C range (Russell and Paesler, 2003), their $\delta^{18}O_P$ values could reflect isotopic equilibrium with environmental waters having mean $\delta^{18}O_w$ values in the +1% to + 2‰ range. Such differences in calculated $\delta^{18}O_w$ values between fish and crocodilian and turtles have also been observed in Early Cretaceous deposits of northeast Thailand. Indeed, these isotopic differences between fish and reptiles may reflect peculiar conditions experienced by fish during apatite mineralization such as temporary isolations from larger water bodies followed by intense evaporation. During periods of isolation of water bodies, tetrapods can move to other water bodies, whereas fish are trapped and record environmental changes in their phosphatic tissues (Amiot et al., 2009). Low $\delta^{13}C_c$ values of the Moroccan and Brazilian lungfishes are compatible with their freshwater living environments with the exception of the very positive value obtained from one Protopterus tooth that remains enigmatic in the framework of this study. The two fishes Vinctifer and Neoproscinetes from the Romualdo Formation of Brazil are considered to be either marine or species that lived in high saline waters. Their high $\delta^{18}O_P$ values are in agreement with their marine life, and the differences in both $\delta^{18}O_p$ and $\delta^{13}C_c$ values suggest differences in foraging environment and diets. Neoproscinetes is a durophageous fish according to its tooth morphology as well as its preserved gut contents found in some specimens (Kriwet, 2001), whereas Vinctifer possesses a well developed basket of gill rakers suggesting a filtering mode of feeding (Maisey, 1994). As marine shelled mollusks have significantly higher $\delta^{13}C_c$ values than marine plankton (Rau et al., 1989; Lécuyer et al., 2004; McConnaughey and Gillikin, 2008), such offset in diet $\delta^{13}C$ values is in good agreement with the difference observed between Vinctifer and Neoproscinetes $\delta^{13}C_c$ values.

5.3. Remarks on mid-Cretaceous Moroccan and Brazilian environmental conditions

Vertebrate $\delta^{18}O_p$ values are surprisingly not high enough to account for the expected hot or dry conditions that might have prevailed in Saharan environments during the mid-Cretaceous. Terrestrial dinosaurs would have undergone elevated transcutaneous

evaporation, along with ingestion of waters from evaporated ponds (at least intermittently), thus leading to high $\delta^{18}O_p$ values. Such expected extremely positive values have been measured in phosphatic remains of extant desert faunas such as the Sahara sand viper Cerastes vipera which displayed $\delta^{18}O_P$ values up to 30% (Lécuyer et al., 1999b), or reptiles and mammals from Sibiloi National Park (Kenya), some of which having $\delta^{18}O_P$ values up to 32‰ (Schoeninger et al., 2000). Morevover, elevated $\delta^{18}O_P$ values of 26% were measured from theropod dinosaur teeth recovered from the Early Cretaceous of northeast Thailand where seasonal drought took place during this period (Amiot et al., 2009). Oxygen isotope compositions of environmental waters estimated from freshwater crocodilians and turtles are in the range of, or slightly lower than, values measured today in tropical and equatorial regions with semi-arid to arid climates (IAEA/WMO, 2006). It is also noteworthy that the $\delta^{13}C_c$ values of sauropods suggest a diet with a δ^{13} C value of -26 ± 1 %, which corresponds to C_3 plants not affected by stress conditions. Under environmental stresses such as elevated temperatures, light, salt or aridity, plant leaves tend to have higher δ^{13} C values as a result of reduced stomatal conductance leading to lower carbon isotope discrimination during photosynthesis (Farquhar et al., 1989). For example, δ^{13} C values as high as -20% have been measured from Mesozoic and Cenozoic fossil plants that lived under environmental stress (Bocherens et al., 1993). On the other hand, high $\delta^{18}O_p$ values measured in lungfish from Morocco and Brazil suggest that their water environments dried up temporally or probably seasonally. From these considerations, it seems that the studied tropical to equatorial mid-Cretaceous environments were characterized by elevated temperatures reaching 34 °C (as deduced from high fish $\delta^{18}O_p$ values) in agreement with mean air temperatures previously proposed (Russell and Paesler, 2003). Water availability does not seem to have been a critical factor for plants and tetrapods as suggested by their carbon and oxygen isotope compositions, but seasonal or temporary drought might have trapped fish in small evaporating bodies of water where lungfish could have survived by burrowing into the mud.

6. Concluding remarks

Moroccan, Tunisian and Brazilian vertebrate apatites have been analyzed for their oxygen and carbon isotope compositions of phosphates ($\delta^{18}O_p$) and carbonates ($\delta^{18}O_c$, $\delta^{13}C_c$) in order to better understand mid-Cretaceous terrestrial climates of low paleolatitudes and to investigate the response of vertebrate faunas in terms of ecology. The following results are underlined:

- Oxygen isotope compositions of surface waters ($\delta^{18}O_w$) estimated from turtle and crocodile $\delta^{18}O_p$ values range from $-5.0 \pm 1.0\%$ to $-2.4 \pm 1.0\%$, which do not differ from mean annual rainwater values occurring today under inter-tropical sub-arid to arid climates.
- High water temperatures ranging from 21 ± 6 °C to 34 ± 2 °C deduced from fish $\delta^{18}O_p$ values are in agreement with those published for mid-Cretaceous low latitudes.

- At each site, coexisting theropod dinosaurs, titanosaurid sauropods, pterosaurs, crocodilians, turtles and fish have distinct $\delta^{18}O_p$ and $\delta^{13}C_c$ values reflecting their ecologies (terrestrial vs. aquatic), diets (water strategies) and foraging environments (marine vs. freshwater).
- Temporary or seasonal droughts are inferred from high $\delta^{18}O_p$ values of lungfish teeth, even though lower reptile $\delta^{18}O_p$ values suggest the use of distinct and most likely larger or regularly renewed bodies of water.
- Environmental conditions of the studied low latitude regions during the Aptian–Cenomanian interval were somewhat similar to those experienced today under semi-arid to arid tropical or equatorial climates, but with higher mean surface temperatures than present-day ones.
- The extreme conditions proposed by Russell and Paesler (2003) appear therefore unlikely for the studied areas and may have occurred further inland in the Aptian deposits of Niger where it has been suggested that dinosaurs were adapted to circumvent stress of heat and aridity (Russell and Paesler, 2003).

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