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Combined oxygen and sulphur isotope analysis—a new tool to unravel vertebrate (paleo)-ecology

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Abstract

Reconstructing the living environment of extinct vertebrates is often challenging due to the lack of proxies. We propose a new proxy to the living environment based on the combined oxygen and sulphur stable isotope analysis of vertebrate hydroxyapatite. We tested this isotopic proxy to 64 biogenic apatite (bones) samples that represent a wide spectrum of the extant vertebrate phylogenetic diversity including crocodiles, snakes, turtles, mammals, birds, lizards, fish and amphibians. We show that the combination of these two isotopic systems allows the living environment of all these vertebrates to be unambiguously distinguished between freshwater (aquatic vs semi-aquatic), seawater (aquatic vs semi-aquatic) and terrestrial. The main goal of this study is to provide a present-day isotopic reference frame and to discuss methodological issues that will serve to interpret future oxygen and sulphur isotope results obtained either from fossil or modern skeletal material. This new isotopic approach of combined oxygen and sulphur isotope analysis will be particularly useful to document major aquatic-terrestrial transitions in the fossil record but also to better constrain the living environment of some present-day species.

Keywords Geochemistry · Stable isotope · Biogenic apatite · Ecology · Fossil

Introduction

Background information

Vertebrate evolution has been many times punctuated by ecological transitions between terrestrial and aquatic (freshwater and seawater) environments resulting in major radiation events: during the Late Devonian-Early Carboniferous, early tetrapods left

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François Fourel francois.fourel@univ-lyon1.fr aquatic environments and colonised terrestrial ones (Ahlberg and Milner 1994); during the Jurassic-Cretaceous, various crocodylomorphs belonging to the thalattosuchians, the pholidosaurids, the dyrosaurids and the eusuchians, radiated in the marine environments (Martin et al. 2014); One hundred million years later, during the Cenozoic (Eocene), early cetaceans also experienced a secondary adaptation to aquatic environments (Gingerich et al. 2001). Reconstruction of a thorough picture of

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these ecological transitions requires detailed knowledge of the living environment of the extinct taxa involved.

Terrestrial, freshwater and marine environments have different physical and chemical properties such as density, viscosity and salinity, resulting in specific morphological and physiological adaptations of living species. Consequently, the morphofunctional analysis of skeletal remains of fossil taxa has often been used to elucidate their living environment (Coates and Clack 1990; Fernández and Gasparini 2000; Pierce et al. 2012; Spoor et al. 2002). However, skeletal remains sometimes may be incomplete or may have lost their original shape during postdepositional events such as burial and tectonic deformation or compaction. Such processes preclude a reliable interpretation of anatomical features in terms of morpho-functionality. Furthermore, soft tissues indicative of specific environments such as salt glands are easily degraded, and delicate ossified structures such as the semicircular canal system of the inner ear are rarely preserved in the fossil record. Finally, morphological features can predate functional adaptation (exaptation process) so that it can be misinterpreted in terms of living environment.

The sediments in which vertebrate fossils are embedded also constitute an important source of information. The detailed study of the lithology, petrology and geochemistry, along with sedimentary structures, allows precise reconstruction of the environmental conditions that prevailed during the deposition of the sediments. However, the living environment of vertebrates does not necessarily represent the depositional environment in which they were embedded (e.g. anoxic bottom waters). This is particularly true for vertebrates that travel long distances or migrate (e.g. anadromous and catadromous fish). Furthermore, carcasses can be transported over long distances after death resulting in a mismatch between the environment deduced from the sediment of the taphocoenosis and the genuine living environment.

Those problems have raised the need for other methods to reconstruct living environments independently of vertebrate morphology and depositional environments. For instance, stable carbon, oxygen or strontium isotope compositions of bones and teeth have been used as direct tracers of the living environment and applied to fossilised remains, such as those of early tetrapods (Goedert et al. 2018), early cetaceans (Roe et al. 1998; Clementz et al. 2006) or crocodilian taxa (Martin et al. 2016), to get a better picture of these major ecological transitions. Here, we propose a new method to determine past living environments of vertebrates based on the combined analysis of oxygen and sulphur isotope compositions of their biogenic apatite.

Oxygen isotope composition of vertebrate apatite

Oxygen isotope composition of surface waters ($\delta^{18}O_w$) is mainly controlled by evaporation and condensation processes during which isotopic fractionation takes place (Craig and Gordon 1965; Dansgaard 1964). Marine environments have relatively uniform $\delta^{18}O_w$ values of $0 \pm 1\%$ except at high latitudes, where $\delta^{18}O_w$ values are lower, ranging from – 3 to – 1‰ due to mixing with ice melt, and at tropical latitudes where high evaporation rates result in positive $\delta^{18}O_w$ values ranging from + 1 to + 2‰, especially in closed tropical and subtropical seas like the Red Sea, the Dead Sea, Mediterranean Sea or Caribbean Sea (Craig and Gordon 1965; Gat 1984; Gat et al. 1996). Hypersaline lagoons or sabkhas (but also inland lakes, such asf in East Africa) can also reach $\delta^{18}O_w$ values higher than + 2‰ (e.g., Gat and Levy 1978).

The $\delta^{18}O_w$ values of freshwaters mainly derive from those of meteoric waters (groundwater contributions being possible) whose ultimate source is seawater. Evaporation of seawater at low latitudes, distillation and cooling of the humid air mass during its transport towards high latitudes are responsible for the negative δ^{18} O values of meteoric waters (Dansgaard 1964). At the global scale, the higher the latitude and altitude, the lower the δ^{18} O values of rainfall and snow. These values are comprised between -6 and -2% at low latitudes and decrease down to about -15% at high latitudes, polar caps excluded. Oxygen isotope compositions of vertebrate biogenic apatite phosphate $(\delta^{18}O_p)$ are linearly correlated with the oxygen isotope composition of their environmental waters (Longinelli 1984; Luz et al. 1984). Consequently, vertebrates living or ingesting different environmental waters will record in their bones distinct oxygen isotope compositions. Nonetheless, it is worth to note that physiological factors such as evaporative transcutaneous water loss and thermo-metabolism, which are species-specific, also impact the oxygen isotope compositions recorded in bioapatites (e.g. Kohn 1996; Levin et al. 2006).

Sulphur isotope composition of vertebrate apatite

Sulphur isotope composition of sulphates (δ^{34} S) is highly variable in modern aquatic environments. Marine environments have high and relatively uniform sulphate δ^{34} S values close to + 21.0% (Böttcher et al. 2007). Most freshwater environments (e.g. rivers, lakes, ponds, precipitations) have comparatively lower sulphate δ^{34} S values, ranging from – 20.0 to + 20.0% (Krouse 1980; Kaplan 1983; Nehlich 2015). It has been shown that the sulphur isotope composition of food is recorded in vertebrate organic tissues (e.g. muscles, hairs) or molecules (e.g. bone collagen) with low isotopic fractionation (+ 0.5% + 2.4%, Nehlich 2015), especially when compared to the oxygen isotopic system. A recent study also measured very low sulphur isotope fractionation values between the collagen of sub-fossil red fox and that of its preys (ranging from -0.54 to $+0.03\%_0$, with a mean analytical error of ± 0.4 ; Krajcarz et al. 2019). Notably, this study further allows such low sulphur isotope fractionation to apply for carnivores.

Sulphur isotope analysis of vertebrate organic tissues is, therefore, particularly relevant to differentiate between freshwater and seawater environments. In particular, this method has been successfully used to determine the living environment exploited by

fish at the species and population levels (Fry 2002; Fry and Chumchal 2011; Hesslein et al. 1991; Nehlich et al. 2013; Trembaczowski 2011) or in archaeological studies to know if ancient human populations relied on freshwater or marine food resources (e.g. Bocherens et al. 2016). More generally, terrestrial environments (including freshwater ones) and animals living there have generally relatively low δ^{34} S values compared to marine environments. Nonetheless, it is worth to note that coastal or island environments may be substantially influenced by sulphate from marine environments, which can be redeposited as rain or aerosols (the so-called 'sea spray' effect) with sulphate δ^{34} S values close to those of marine environment (+ 20.3%); Nielsen 1974; Norman et al. 2006). Consequently, the δ^{34} S values of vertebrates living in those terrestrial environments submitted to sea spray effect can be relatively high and may complicate interpretation concerning the living environment.

Due to technical difficulties, sulphur isotope analyses have been only applied to organic tissues that easily degrade after animal death and are rarely preserved in the fossil record. A new method has been recently developed to measure the sulphur isotope ratios (${}^{34}S/{}^{32}S$) of sulphate compound in calcium phosphate minerals (analytical precision equals $0.5\%(1\sigma)$) with a low-S concentration (0.14% to 1.19%) such as vertebrate bioapatites (Fourel et al. 2015; Goedert et al. 2016). Previous results indicated that sulphur isotope compositions of environmental waters are recorded in vertebrate inorganic tissues (bone apatite) with low isotopic fractionation ($0.8\%_{0} \pm$ $0.8\%_{0}$, n = 5; Goedert et al. 2018). Therefore, sulphur isotope analysis of bioapatite from extinct vertebrates can provide estimates of the salinity of their aqueous environments (Goedert et al. 2018).

Material and methods

Sixty-four vertebrate bone apatite samples have been collected and analysed in this study (Online Information 1). Samples were selected to encompass a broad ecological and taxonomic spectrum of vertebrates (crocodiles, snakes, turtles, mammals, birds, lizards, fish and amphibians). For each taxonomic group, vertebrates of distinct ecology such as aquatic (freshwater vs marine), semi-aquatic and terrestrial were selected (Online Information 2). Oxygen and sulphur isotope analyses have been performed on each bone sample of the 64 vertebrates.

Forty vertebrate bone apatite samples were collected in the osteological collections of the 'Musée des Confluences' of Lyon, France. Samples were further selected in historical collections to ensure a wild provenance. Specimens with a labelled precise localisation were prioritised when possible. In addition, 24 vertebrate bone apatite samples for which sulphur isotope composition have been previously published (Goedert et al. 2016, 2018; cf. Online Information 1) have been added

to the dataset and their oxygen isotope composition measured in this study. For each specimen, about 100 mg of bone powder was sampled using a spherical diamond-tipped drill bit. The surface of the bone, which may have been chemically treated for curatorial purpose (samples coming from the 'Musée des Confluences'), was removed prior to sampling.

All statistical tests were performed using Past 3.22 software. We used Mann-Whitney U test to compare the different median values and give the associated P value (P). Data of Figs. 1 and 2 were plotted using KaleidaGraph 4.5.3 software. Figures were drawn using Inkscape 0.92.3.

Oxygen isotope analysis

Bone apatite samples were treated following the wet chemistry protocol described by Crowson et al. (1991) and slightly modified by Lécuyer et al. (1993). This protocol consists in the isolation of phosphate (PO_4^{3-}) from apatite using acid dissolution and anion-exchange resin. For each sample, 30 mg of enamel powder was dissolved in 2 mL of 2 M HF overnight. The CaF₂ residue was separated by centrifugation, and the solution was neutralised by adding 2.2 mL of 2 M KOH. 2.5 mL of Amberlite[™] anionexchange resin was added to the solution to separate the PO_4^{3-} ions. After 24 h, the solution was removed and the resin was eluted with 27.5 mL of 0.5 M NH₄NO₃. After 4 h, 0.5 mL of NH₄OH and 15 mL of an ammoniacal solution of AgNO₃ were added, and the samples were placed in a thermostated bath at 70 °C during 7 h, allowing the precipitation of silver phosphate (Ag₃PO₄) crystals. When only a few mg of apatite powders could be collected, the wet chemistry procedure was adapted following Bernard et al. (2009) for small sample weights (about 3 mg).

Oxygen isotope compositions were measured using a high-temperature pyrolysis (Py) technique involving a VarioPYROcube[™] elemental analyser (EA) interfaced in continuous flow (CF) mode to an IsoprimeTM isotopic ratio mass spectrometer (IRMS) (EA-Py-CF-IRMS technique (Fourel et al. 2011; Lécuyer et al. 2007) at the Laboratoire de Géologie de Lyon (UMR 5276, Université Claude Bernard Lyon 1). For each sample, 5 aliquots of 300 µg of Ag₃PO₄ were mixed with 300 µg of pure graphite powder and loaded in silver foil capsules. Pyrolysis was performed at 1450 °C. Measurements were calibrated against the NBS120c (natural Miocene phosphorite from Florida: $\delta^{18}O = 21.7\%$ (V-SMOW); Lécuyer et al. 1993) and the NBS127 (barium sulphate, BaSO₄: $\delta^{18}O = 9.3\%$ (V-SMOW); Hut 1987). Silver phosphate samples precipitated from standard NBS120c were repeatedly analysed $(\delta^{18}O_{p} = 21.6\%; 1\sigma = 0.4; n = 16)$ along with the silver phosphate samples derived from vertebrate bioapatites to ensure that no isotopic fractionation took place during the



Fig. 1 $\delta^{18}O_p$ and $\delta^{34}S_{apatite}$ values of modern vertebrates including (from left to right) crocodiles, snakes, turtles, mammals, birds, lizards, fish and amphibians. **a** Oxygen isotope composition of bone phosphate ($\delta^{18}O_p$) as variations in parts per mille from the ratio of ${}^{18}O_1{}^{16}O$ in Vienna Mean Ocean Water (% VSMOW) **b** Sulphur isotope composition of bone apatite ($\delta^{34}S_{apatite}$) as variations in parts per mille from the ratio of ${}^{34}S/{}^{32}S$ in Vienna Canyon Diablo Troilite (% VCDT). For **a**, **b**, each data point represents a biologically independent animal (n = 64) and

wet chemistry. The NBS120c average standard deviation equals $0.29 \pm 0.14\%$. Data are reported as $\delta^{18}O_p$ in % values vs V-SMOW.

Sulphur isotope analysis

Sulphur isotope compositions were measured using a VarioPYROcubeTM elemental analyser in NCS combustion mode interfaced in continuous-flow mode with an Isoprime 100TM isotope ratio mass spectrometer hosted by the platform 'Ecologie Isotopique' of the 'Laboratoire d'Ecologie des Hydrosystèmes Naturels et Anthropisés' (LEHNA,

corresponds to the average value of five and three repeated measurements for oxygen and sulphur isotope analysis, respectively (see "Material and Methods"). Each error bar corresponds to 1 s.d. (Online Information 1). For both panels, light blue, dark blue and green colours indicate that the species lives in freshwater, seawater or terrestrial environments, respectively (see Online Information 2). The name of each species is indicated close to the corresponding dot

UMR 5023, Villeurbanne, France). For each bone apatite sample, 3 aliquots of 7 mg of bioapatite powder were mixed with 20 mg of pure tungsten oxide (WO₃) powder and loaded in tin foil capsules. Tungsten oxide is a powerful oxidant ensuring the full thermal decomposition of apatite sulphate into sulphur dioxide (SO₂) gas (Goedert et al. 2016). Measurements have been calibrated against the NBS127 (barium sulphate, BaSO₄ δ^{34} S = + 20.3% (V-CDT); Halas and Szaran 2001) and S1 (silver sulphide, Ag₂S δ^{34} S = - 0.3% (V-CDT); Robinson 1995) international standards. For each analytical run of bone samples, we have also analysed BCR32 samples as a



Fig. 2 Reconstructed oxygen and sulphur isotope composition of the environmental waters ($\delta^{18}O_w, \delta^{34}S_w$) of the modern vertebrates. For oxygen, the isotopic composition of water was calculated using published isotopic fractionation equations for different groups of vertebrates (Online Information 3). For sulphur, the isotopic composition of water is very close to that recorded in bone apatite (i.e., almost no isotopic fractionation) and was calculated using published values of sulphur isotope composition of bone apatite and associated environmental water measured in present-day vertebrates (Goedert et al. (2018); Online Information 4). Each data point represents a biologically independent animal (n = 64) and corresponds to the average value of five and three repeated measurements for oxygen and sulphur isotope analysis, respectively (see "Material and Methods"). Each dot is numbered according to the species it represents (cf. Online Information 1). Error bars of each individual data point are given in Online Information 3 and Online Information 4 for oxygen and sulphur respectively. Results are given as variations in parts per mille from the ratio of ¹⁸O/¹⁶O in Vienna Mean Oean Water (% VSMOW) for oxygen and ³⁴S/³²S in Vienna Canyon Diablo Troilite (% VCDT) for sulphur. Species living in freshwater are represented by light blue dots; those living in seawater are represented by dark blue dots, and green dots are used for terrestrial species. (1): Crocodylus niloticus; (2): Crocodylus siamensis; (3): Gavialis gangeticus; (4): Crocodylus porosus; (5): Crocodylus suchus; (6):

compositional and isotopic standard (S% = 0.72, certified value ((Community Bureau of Reference 1982); δ^{34} S = 18.4% (V-CDT); Fourel et al. 2015; Goedert et al. 2016) to ensure that analytical conditions were optimal to perform sulphur isotope analyses of samples with low-S content. The sample average standard deviation for δ^{34} S measurements is 0.34% ± 0.34%. Data are reported as δ^{34} S in % vs V-CDT.

Crocodylus suchus; (7): Acrochordus granulatus; (8): Xenochrophis flavipunctus; (9): Homalopsis buccata; (10): Hydrophis obscurus; (11): Pelamis platura; (12): Cerastes cerastes; (13): Testudo kleinmanni; (14): Dogania subplana; (15): Chelvdra serpentina; (16): Chelonia mydas; (17): Chelonia mydas; (18): Trachemys scripta elegans; (19): Trachemys scripta elegans; (20): Trachemys scripta elegans; (21): Trachemys scripta elegans; (22): Lutra lutra; (23): Platanista gangetica; (24): Monachus monachus; (25): Odobenus rosmarus; (26): Phoca vitulina; (27): Monodon monoceros; (28): Enhydra lutris; (29): Phocoena phocoena; (30): Dugong dugon; (31): Hydrodamalis gigas; (32): Hydrodamalis gigas; (33): Hippopotamus amphibius; (34): Rhinoceros sondaicus; (35): Camelus dromedarius; (36): Camelus bactrianus; (37): Ursus arctos; (38): Ursus maritimus; (39): Tapirus indicus: (40): Tapirus terrestris: (41): Castor fiber: (42): Larus argentatus; (43): Spheniscus demersus; (44): Anas platyrhynchos; (45): Buteo buteo; (46): Amblyrhynchus cristatus; (47): Cyprinus carpio; (48): Silurus glanis; (49): Oncorhynchus mykiss; (50): Salmo trutta; (51): Salvelinus fontinalis; (52): Oncorhynchus mykiss; (53): Sander lucioperca; (54): Solea solea; (55): Gadus morhua; (56): Limanda limanda; (57): Dicentrarchus labrax; (58): Oncorhynchus nerka; (59): Pelophylax ridibundus; (60): Pelophylax ridibundus; (61): Pelophylax ridibundus; (62): Pelophylax ridibundus; (63): Pelophylax ridibundus; (64): Salamandra salamandra

Results

Oxygen isotope

The different vertebrates analysed had oxygen isotope compositions ranging from + 12.1 to + 24.2% V-SMOW (Online Information 1; Fig. 1a), which mainly reflect the variability of oxygen isotope compositions of environmental waters. On the whole, vertebrates living or foraging in marine environments had significantly higher $\delta^{18}O_p$ values than animals living or foraging in continental (freshwater or terrestrial) environments (median $\delta^{18}O_p = +19.8\%$, $1\sigma = 3.0$, n = 18 vs median $\delta^{18}O = +15.4\%$, $1\sigma = 2.4$, n = 40; P = 4.244e-5; Mann-Whitney *U* test). It is also worth to note that vertebrates which live in both freshwater to seawater environment had intermediate median $\delta^{18}O_p$ values ($\delta^{18}O_p = +17.7\%$, $1\sigma = 0.9$, n = 6), although the difference was only significant compared to continental environments and not seawater ones (P = 0.01255 and P = 0.1611, respectively). One exception concerns the horned desert viper (*Cerastes cerastes*) and the Kleinmann's tortoise (*Testudo kleinmanni*), which had both recorded high oxygen isotope ratios in their bones due to their desert lifestyle.

Sulphur isotope

The different vertebrates analysed had sulphur isotope compositions of apatite (δ^{34} S_{apatite}) ranging from + 1.1 to + 22.9% V-CDT (Online Information 1; Fig. 1b). On the whole, vertebrates living or foraging in marine environments had significantly higher δ^{34} S values than those living or foraging in continental (freshwater or terrestrial) environments (median $\delta^{34}S_{apatite} = +$ $16.9\%, 1\sigma = 4.4, n = 18$ vs median δ^{34} S_{apatite} = + 10.4\%, 1\sigma = 4.4, n = 40; P = 0.0001357). This isotopic pattern reflects an almost systematic ³⁴S-enrichment of marine environments compared to continental ones. It is again worth to note that vertebrates living in freshwater to seawater environment had intermediate median $\delta^{34}S_{apatite}$ values ($\delta^{34}S = +13.8\%_o$, $1\sigma =$ 6.0, n = 6), although the difference was not significant with that of continental or marine environments (P = 0.1063 and P =0.5264). Sulphur isotope analysis of fossilised apatite can, therefore, help to detect the presence or proximity of seawater in the living environments of extinct vertebrates.

Discussion

Oxygen isotope composition

Oxygen isotope analysis of vertebrate biogenic apatite has been widely applied to fossilised apatite of extinct vertebrates to get information on their living environment (e.g. Clementz et al. 2003, 2006; Tütken et al. 2006; Amiot et al. 2015, 2009, 2010; Pouech et al. 2014; Guy et al. 2018). As illustrated by our results, this analysis can be particularly useful to distinguish vertebrates living or foraging in marine environments from those living or foraging in continental (freshwater or terrestrial) ones (e.g. sharks: Gates et al. 2019; mosasaurs: Makádi et al. 2012; coelacanths: Simon et al. 2003).

It can also be used to further differentiate aquatic or semiaquatic lifestyle from a terrestrial one in the case of sympatric vertebrates (e.g. Amiot et al. 2010). Indeed, terrestrial animals

lose more water than semi-aquatic animals through transcutaneous evaporation or sweat. Water lost during this process as vapour is preferentially ¹⁶O-enriched, resulting in a relative ¹⁸O-enrichment of the residual body water (Cerling et al. 2008). Although the different vertebrates sampled come from different regions of the world, it should be noted for instance that the Eurasian otter (Lutra lutra) and the Eurasian beaver (Castor fiber), both having a semi-aquatic lifestyle, have recorded lower oxygen isotope ratios in their bones than fully terrestrial mammals (Online Information 1 and Fig. 1a). This is also the case for the semi-aquatic mallard duck (Anas *platvrhynchos*), which recorded in its bones lower oxygen isotope ratios than the common buzzard (Buteo buteo) (Online Information 1 and Fig. 1). In the latter case, it is worthy to note that both specimens come from the same geographic area and therefore rely on environmental waters of comparable oxygen isotope compositions.

On the contrary, it can be used to detect desert lifestyle (Lécuyer et al. 1999). For instance, the horned desert viper (*Cerastes cerastes*) and the Kleinmann's tortoise (*Testudo kleinmanni*), had both recorded high oxygen isotope ratios in their bones.

Nonetheless, for low-latitude environments, oxygen isotope compositions of freshwater and marine environments can display significant overlap. Consequently, water oxygen isotope compositions recorded in vertebrate apatites may not always be a diagnostic tracer of their living environment (e.g. Pouech et al. 2014).

Sulphur isotope composition

Compared to oxygen, sulphur isotopes have been less applied to question the ecology of extinct vertebrates, principally due to technical difficulties. Due to the large amplitude of natural isotopic variations, particularly observed between terrestrial and marine environments, it remains a particularly relevant environmental tracer (cf. Background information).

However, as discussed in the "Introduction" section, the 'sea spray' effect may complicate interpretation concerning the living environment of vertebrates for terrestrial environment under the influenced of marine ones. Moreover, some freshwater settings may have sulphur isotope compositions close to that of marine environments. For instance, rivers draining basins in which marine evaporites are exposed may have elevated dissolved sulphate content (more than 200 mg/L for the Colorado River system; Shope and Gerner 2014) and δ^{34} S values (up to seawater-like 19.5%) for the Mackenzie River system; Hitchon and Krouse 1972). Therefore, vertebrates living in such environments are expected to have high sulphur isotope compositions that could be misinterpreted as reflecting an aqueous environment at least submitted to some marine influences. Finally, vertebrate species living in aquatic environments submitted to the influences of both fresh and

marine water, like in estuaries, may record a sulphur isotope composition in their bioapatite difficult to correctly interpret in terms of living environment.

Combined oxygen and sulphur isotope composition

On the whole, the combined use of oxygen and sulphur isotope compositions of bone apatite allows, in most cases, environmental identification for the present-day vertebrates after the conversion of the measured δ^{18} O and δ^{34} S values of apatite into environmental water δ^{18} O value and dissolved environmental sulphate δ^{34} S values using known isotopic fractionation equations (Fig. 2; Online Information 3 and 4).

The complementarity of these two isotopic systems lies in the different abundance ratios of oxygen and sulphur, respectively, in seawater and freshwater bodies. Indeed, oxygen is equally present (as H_2O) in both marine and freshwater reservoirs whereas sulphur content (as SO_4^{2-}) of seawater is generally 100 to 1000 times higher than in freshwater (Fry and Chumchal 2011). Consequently, sulphur isotopes will be particularly relevant to detect the presence of seawater in the environment, even if only a small quantity of seawater intrudes freshwater environment, and oxygen isotopes will be relevant to quantify the amount of freshwater in the environment, in particular in aquatic environments where freshwater and seawater are mixing, like in deltas or estuaries (Goedert et al. 2018).

Vertebrates living or foraging in marine environments tend to have higher oxygen and sulphur isotope compositions recorded in their bone apatite than those from freshwater and terrestrial habitats. This rule is especially valid when we compare vertebrates of close phylogenetic affinity. For instance, the wild gharial (Gavialis gangeticus), living in freshwater streams, and the two captive specimens of desert crocodiles (Crocodylus suchus), kept in freshwater at the Zoo of Lyon, have recorded in their bone apatite $\delta^{18}O_p$ and $\delta^{34}S_{apatite}$ values (+ 12.1% and + 15.9%, + 14.5% and + 8.2%, and + 13.9% and + 8.5%, respectively) lower than those measured in bones of the wild Nile crocodile (Crocodilus niloticus; +17.5% and +20.1%) and saltwater crocodile (Crocodilus porosus; +16.8% and +16.3%), both known to undertake incursions in brackish waters to seawaters (cf. Online information 2). Similarly, the sea otter (Enhydra *lutris*), fully adapted to life in seawater, has higher $\delta^{18}O_{p}$ and $\delta^{34}S_{a patite}$ values (+ 17.4% and + 18.1%) than those of the Eurasian otter (*Lutra lutra*) ($\delta^{18}O_p = +14.0\%$ and $\delta^{34}S = +$ 12.8%), inhabiting freshwater environments. In a similar way, the marine narwhal (Monodon monoceros) has higher $\delta^{18}O_p$ and δ^{34} S_{apatite} values (+ 17.0% and + 16.0%) than those of the South Asian river dolphin (Platanista gangetica; +14.3% and + 10.2%%).

The general picture we have of major ecological transitions that took place during vertebrate evolution are incomplete and potentially biased as it corresponds to the final stages of these transitions. For instance, the colonisation of terrestrial environments by early tetrapods at the beginning of the Carboniferous gave rise to a wide evolutionary radiation of terrestrial tetrapods that are still present on lands today. Similarly, the multiple iterations of secondary adaptation to the aquatic environment are well illustrated by the numerous species of vertebrates belonging to different groups (crocodiles, snakes, turtles, lizards, birds and mammals), which live in present-day aquatic environments. All these vertebrates testify that different groups adapted to new environments from a common ancestor. However, the way these major ecological transitions proceeded, especially during their early stages, is difficult to infer and often remained elusive. Indeed, morpho-functional adaptations to a specific environment can be diachronous with its effective use (exaptation); the diagnose of living environment of vertebrates from morpho-functional analysis is thereby limited. Therefore, the combined use of ¹⁸O/¹⁶O and ³⁴S/³²S ratios of skeletal apatite should be particularly promising and powerful to document major ecological transitions in the fossil record for any phylogenetic group of vertebrates. For instance, this method has already been successfully applied to determine the aquatic environment of some Devonian early tetrapods and their associated vertebrate fauna (Goedert et al. 2018). Furthermore, it could also help to precise the ecology of some present-day aquatic vertebrates and shed light on the modalities of transition between terrestrial and aquatic environments during the course of vertebrate evolution over the Phanerozoic. It is also worthy to note that this method has the potential to shed light on the ecology of numerous present-day vertebrates living in transitional environments, and for which the ecology remains unclear.

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Author contribution statement All authors contributed to the study conception and design. Material preparation and data collection were performed by J. Goedert, D. Berthet and R. Amiot. Material analysis were performed by J. Goedert, F. Fourel and L. Simon. The first draft of the manuscript was written by J. Goedert, R. Amiot and C. Lécuyer, and all authors commented on previous versions of the manuscript. All authors read and approved the final manuscript.

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Data availability All data generated or analysed during this study are included in this published article [and its online information files].

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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