Regional endothermy as a trigger for gigantism in some extinct macropredatory sharks

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Abstract

Otodontids include some of the largest macropredatory sharks that ever lived, the most extreme case being Otodus (Megaselachus) megalodon. The reasons underlying their gigantism, distribution patterns and extinction have been classically linked with climatic factors and the evolution, radiation and migrations of cetaceans during the Paleogene. However, most of these previous proposals are based on the idea of otodontids as ectothermic sharks regardless of the ecological, energetic and body size constraints that this implies. Interestingly, a few recent studies have suggested the possible existence of endothermy in these sharks thus opening the door to a series of new interpretations. Accordingly, this work proposes that regional endothermy was present in otodontids and some closely related taxa (cretoxyrhinids), playing an important role in the evolution of gigantism and in allowing an active mode of life. The existence of regional endothermy in these groups is supported here by three different approaches including isotopic-based approximations, swimming speed inferences and the application of a novel methodology for assessing energetic budget and cost of swimming in extinct taxa. In addition, this finding has wider implications. It calls into question some previous paleotemperature estimates based partially on these taxa, suggests that the existing hypothesis about the evolution of regional endothermy in fishes requires modification, and provides key evidence for understanding the evolution of gigantism in active macropredators.

Introduction

Otodontids or megatooth sharks are an extinct family of lamniform apex predators that lived from the Early Paleocene to Pliocene [1]. This group is only known from disarticulated remains, mainly isolated teeth but also some sets of associated teeth and vertebral centra [2–5]. The large size of these remains, including teeth that reach up to 168 mm in height [6], has aroused the interest in the group for some time. In consequence, several authors have tried to establish accurate allometric relationships between tooth height (or vertebral centra width) and body length and between body length and body mass in extant sharks with the goal of inferring sizes and masses of otodontids [6–8]. The most conservative estimates for body
length of the popularly known “Megalodon” (*Otodus (Megaselachus) megalodon* following Capetta [1]) are about 16 meters, which makes it the largest megapredatory shark known among both fossil and extant taxa [6]. However, the scarcity of studies dealing with the factors that allowed megatooth sharks to reach such large sizes is striking. In this sense, some works (e.g., [6,9,10]) suggest that the gigantism as well as the distribution patterns, evolution and extinction of otodontids were closely linked to climatic factors and/or the abundance of their potential prey after the radiation of the cetaceans in the Paleogene.

Most of these previous hypotheses, however, presupposed otodontids as ectothermic fishes without taking into account some possible size-related metabolic constraints on the macropredatory lifestyle [11,12]. In this sense, there is a clear tendency for the largest marine vertebrates to be slow filter-feeding planktivores, whereas active macropredators are much smaller, suggesting that some size-related factors determine their activity and feeding strategy [13]. Most of the biggest ectothermic fishes, such as the whale shark (*Rhincodon typus*), the basking shark (*Cetorhinus maximus*) and the megamouth shark (*Megachasma pelagios*), are in fact filter-feeding chondrichthynes that range in size from 5.5 to 21 meters [14]. Strikingly, the only marine vertebrates with sizes and lifestyles comparable to those of the biggest megatooth sharks are large odontocete cetaceans with well-developed homeothermy, such as the killer whale (*Orcinus orca*) and the sperm whale (*Physeter catodon*) [15].

Contrary to all classical interpretations, Ehret [5] questioned the ectothermic nature of otodontids and suggested possible endothermy based on the lack of a correlation between their growth rates and the sea temperature through the Cenozoic, which would be expected if they were ectotherms. More recently, Pimiento et al. [16] have also proposed regional endothermy for *O. megalodon* on the basis of paleobiogeographical data and the range of water temperatures this species inhabited. Regional endothermy is the ability of some fish lineages to maintain certain body areas at higher temperatures than the surrounding water by means of vascular countercurrent heat exchangers or specialized thermogenic organs [17,18]. This adaptation involves an active mode of life and much higher metabolic rates than those of the ectothermic fishes of the same size [19,20]. Within osteichthynes, it has appeared independently in at least three different groups including tunas (tribe Thunnini, Scombridae), billfishes (Xiphiidae and Istiophoridae) and *Gasterochisma* [17]. Recently, whole-body endothermy has also been described in the opah (*Lampris guttatus*) [21]. In chondrichthynes, regional endothermy is restricted to lamniform sharks, where it is found in two of the three species of alopids [22–24] and in all species of the family Lamnidae ([25] and references therein).

Interestingly, regional endothermy of otodontids is rather parsimonious considering that they seem to be phylogenetically closely related to lamnids [5], according to two hypotheses about the relationships between the groups. The first considers otodontids as lamnids [6,9,26–29], while the second considers them as a separate family, which is either sister-group, or closely related to laminids [1,5,30–36]. In fact, some works have suggested that both groups could have evolved from representatives of other Late Cretaceous-Paleocene lamniform sharks, traditionally included in the family Cretoxhirinidae [1,9,26,31] (see Results and discussion section for a more detailed revision of the phylogenetic affinities of cretoxyrhinids).

Here it is proposed that regional endothermy was present in otodontids and some close related taxa (cretoxyrhinids), playing a key role in their evolutionary history, allowing gigantism and the maintenance of active macropredatory modes of life. Three approaches provide evidence for regional endothermy in these groups: (1) testing the degree to which their body temperature is dependent on that of the surrounding water by analysing δ¹⁸O differences between teeth of cretoxyrhinids-otodontids and associated ectotherms; (2) assessing whether burst swimming speeds of cretoxyrhinids-otodontids, estimated assuming ectothermy and regional endothermy, were fast enough for hunting successfully on their potential prey; and
(3) assessing the energetic viability of ectothermy in cretoxyrhinids by calculating whether their costs of swimming were long-term sustainable by their metabolic budget. In addition, a number of multidisciplinary implications derived from this finding are discussed. These include: (1) the possible overestimation of some previous paleotemperature estimates based partially on these taxa, (2) the reinterpretation of some pre-existing hypotheses about the evolution of regional endothermy in fishes, and (3) the contribution of this work to the understanding of the evolution of gigantism in active macropredators.

**Data source and methodology**

**Oxygen isotopic approach**

Vertebrate oxygen isotopic data have been compiled from the literature including records from 20 different sedimentary beds, ranging in age from the Cretaceous to the Miocene, where remains of Cretoxyrhinidae and Otodontidae have been recovered in association with presumed ectothermic taxa (Table 1 and S1 Table). Following the methodology established by Bernard et al. [37], the difference in δ¹⁸O values between coexisting cretoxyrhinids/otodontids and ectothermic taxa has been calculated in all beds, plotting them as a function of the ectothermic taxa δ¹⁸O value (as a proxy of seawater temperature) and adjusting all points to a regression line whose slope is an indicator of the degree of thermoregulation. Slope values close to -1 imply that body temperature is independent of that of the water, slope values close to 0 imply a complete dependence, and intermediate slope values indicate some degree of independence. Three different regression analyses were performed: (1) including all cretoxyrhinids-otodontids and all associated ectothermic taxa; (2) including all cretoxyrhinids-

### Table 1. Mean δ¹⁸O values of cretoxyrhinids-otodontids and associated ectotherms from different sedimentary beds.

| Code | Locality | Age          | Cretoxyrhinid or otodontid taxa | Cretoxyrhinid or otodontid δ¹⁸O | Ectotherm δ¹⁸O | Ectotherm δ¹⁸O (without bentic genera) | Source |
|------|----------|--------------|---------------------------------|---------------------------------|---------------|----------------------------------------|--------|
|      |          |              |                                 | n δ¹⁸O SD  | n δ¹⁸O SD | n δ¹⁸O SD                              |        |
| 1    | Cerro la Bruja, Peru | Miocene  | Otodus megalodon | 1 20.90 | 2 21.00 | 0.42 | 2 21.00 | 0.42 | [39] |
| 2    | Sidi Daoui, Morocco | Upper Maastrichtian | Cretolamna biauriculata marocana | 1 19.98 | 1 20.15 |       |       |       |        |
| 3    | Oued Erguita, Morocco | Lower Maastrichtian | Cretolamna | 1 18.95 | 1 18.66 |       |       |       |        |
| 4    | Ouled Abdoun, Morocco | Upper Maastrichtian | Cretolamna | 2 20.35 | 0.92 | 1 19.50 | 1 19.50 |       |        |
| 5    | Ouled Abdoun, Morocco | Thanetian | Cretolamna sp. | 1 19.47 | 1 19.98 |       |       |       |        |
| 6    | Ouled Abdoun, Morocco | Thanetian | Otodus sp. | 2 20.92 | 0.08 | 3 21.65 | 0.61 | 3 21.65 | 0.61 | [38] |
| 7    | Ouled Abdoun, Morocco | Thanetian | Otodus sp. | 2 20.49 | 0.04 | 1 20.57 |       | 1 20.57 |       |        |
| 8    | Ouled Abdoun, Morocco | Maastrichtian | Cretolamna marocana | 1 20.56 | 1 19.87 |       | 1 19.87 |       |        |
| 9    | Ganntour Basin, Morocco | Maastrichtian | Cretolamna marocana | 1 21.04 | 1 21.19 |       | 1 21.19 |       |        |
| 10   | Ganntour Basin, Morocco | Maastrichtian | Cretolamna marocana | 2 20.67 | 0.48 | 1 21.03 |       | 1 21.03 |       |        |
| 11   | Ganntour Basin, Morocco | Maastrichtian | Cretolamna marocana | 2 20.44 | 0.65 | 2 20.53 | 0.35 | 2 20.53 | 0.35 | [38] |
| 12   | Ganntour Basin, Morocco | Maastrichtian | Cretolamna marocana | 2 20.41 | 0.65 | 2 20.77 | 0.11 | 2 20.77 | 0.11 | [38] |
| 13   | Ganntour Basin, Morocco | Maastrichtian | Cretolamna marocana | 2 20.26 | 0.22 | 1 19.62 |       | 1 19.62 |       |        |
| 14   | Ganntour Basin, Morocco | Maastrichtian | Cretolamna marocana | 2 20.25 | 0.10 | 2 20.39 | 0.37 | 2 20.39 | 0.37 | [38] |
| 15   | Ganntour Basin, Morocco | Maastrichtian | Cretolamna marocana | 2 20.76 | 0.18 | 4 20.94 | 0.23 | 4 20.94 | 0.23 | [38] |
| 16   | Yonne, France | Upper Campanian | Cretolamna appendiculata | 1 21.00 | 1 21.10 |       | 1 21.10 |       |        |
| 17   | Ardèche, France | Upper Aptian | *Otodus sp. | 1 20.10 | 1 20.70 |       |       |       |        |
| 18   | Skotniki, Poland | Lower Cenomanian | *Otodus appendiculatus | 1 20.52 | 2 19.35 | 0.77 | 2 19.35 | 0.77 | [42] |
| 19   | Asen, Sweden | Uppermost Lower Campanian | Cretolamna appendiculata | 1 20.94 | 1 19.62 |       | 1 19.62 |       |        |
| 20   | Benguerir, Morocco | Maastrichtian | Cretolamna marocana | 6 19.43 | 0.31 | 5 19.98 | 0.51 | 5 19.98 | 0.51 | [42] |

* Cretolamna specimens

https://doi.org/10.1371/journal.pone.0185185.t001
otodontids and only associated pelagic ectothermic taxa (minimizing variability caused by the thermal gradient of the water column); and (3) including only cretoxyrhinids-otodontids and associated pelagic ectothermic taxa from Kocsis et al. [38], where remains considered as contemporaneous come from well delimited individual layers (reducing variability associated with low stratigraphic and temporal precision). Data from specimens doubtfully assigned to *Cretolamna* and *Otodus* or specimens where diagenetic alterations have been demonstrated were not included in the analyses. In order to avoid the inclusion of endothermic taxa as a proxy of water temperature, the following cases were also discarded: (1) data from expected endothermic taxa, such as mammals, ichthyosaurs, plesiosaurs, mosasaurs or some fish groups (see Introduction) and (2) data from chondrichthyans whose taxonomic assignment does not rule out the possibility of pertaining to one of the regional endothermic families.

In parallel, measurements of internal body organs/tissues and surrounding water temperature have also been compiled for three extant species of regional endothermic sharks from Lowe and Goldman [25] (S2 Table). Data were represented in a similar way that was done for isotopic data, calculating the differences in temperature between each pair of measurements for all sharks, plotting them as a function of the seawater temperature and adjusting all points to a regression line. Slopes were then contrasted with those obtained for δ^{18}O values of cretoxyrhinids and otodontids in order to evaluate if the degree of thermoregulation is comparable between living and extinct groups.

Finally, latitudinal changes in δ^{18}O tooth values of cretoxyrhinids and presumed ectothermic fishes were also compared from Campanian-Maastrichtian data compiled in Pucéat et al. [42] (S3 Table). δ^{18}O values were represented as a function of latitude, regression lines were obtained for both groups and ANCOVA analysis was performed by means of PASW software. The same analyses were carried out considering the sea surface temperatures predicted from the δ^{18}O values (S3 Table). Data from undetermined teeth were not included in either analysis.

Cruise and burst speed inferences

Records of cruise and burst relative swimming speeds of living fishes have been compiled from the literature including both ectothermic and regional endothermic species, trying to cover a wide range of sizes and taxonomic groups (S4 Table). Data were log-transformed and plotted as a function of total fish length. Then, the scaling of both swimming speeds was studied in ectotherms and regional endotherms by linear regression analysis and differences between both groups were tested by ANCOVA analysis using PASW software. Cruise and burst swimming speeds of some cretoxyrhinids (*Cretolamna* and *Cretosyrhina*) and otodontids (*Megalolamna*, *Otodus* (*Megaselacus*), *Otodus* (*Otodus*) and *Parotodus*) were estimated from their total body lengths by interpolation of the regression analyses. Estimates were carried out considering them both as ectotherms and regional endotherms and 95% individual prediction intervals were calculated for all swimming speeds. Total body lengths of 17.9 m, 9.2 m, 6.8 m, 5.0 m, 6.4 m and 3.0 m have been considered here for *Otodus* (*M*.), *Otodus* (*O*.), *Parotodus*, *Megalolamna*, *Cretosyrhina* and *Cretolamna* respectively. Body length estimations of cretoxyrhinids (*Cretosyrhina* and *Cretolamna*) were taken from previous studies based on articulated or semi-articulated specimens [43,44]; whereas those of otodontids (*Megalolamna*, *Parotodus*, *Otodus* (*O*.) and *Otodus* (*M*.) were based on analysis of isolated teeth figured in the literature. Following the methods described in Pimiento et al. [45], once a range of plausible positions was assigned to each tooth, crown height was used to calculate body lengths from the position-specific regressions established in white sharks by Shimada [8] (Table 2). Only one tooth per species was selected representing the largest or one of the largest records for its position in the dental series. This allows a conservative estimate of the body sizes of these taxa to be obtained, but
larger individuals probably existed. In any case, determining the maximum body lengths of these sharks is beyond the scope of this paper and considering larger body size estimates would only favour the endothermic scenario (see Results and discussion section).

Independent estimations of cruise and burst swimming speed of *Cretoxyrhina* were obtained following the models developed by Sambilay [48] on the basis of the relationship between swimming speed and caudal fin aspect ratio:

\[
\log_{10}(S_c) = -0.828 + 0.6196 \times \log_{10}(PCL) + 0.3478 \times \log_{10}(AR)
\]

\[
\log_{10}(S_b) = -0.0659 + 0.6196 \times \log_{10}(PCL) + 0.3478 \times \log_{10}(AR)
\]

Considering \(AR = H^2 / S\)

Where \(S_c\) is cruising swimming speed in kilometres \(\times\) hours\(^{-1}\), \(S_b\) is burst swimming speed in kilometres \(\times\) hours\(^{-1}\), \(PCL\) is precaudal body length (i.e., body length excluding the caudal fin or standard length) in centimetres, \(AR\) is aspect ratio, \(H\) is height of the caudal fin, and \(S\) is surface area of the caudal fin (note that \(H^2\) and \(S\) should be expressed in the same units because \(AR\) is dimensionless).

\(PCL\) of *Cretoxyrhina* was calculated assuming 640 cm TL from Mollet and Cailliet’s [49] equation:

\[PCL = -0.09195 + 0.8535 \times TL\]

The aspect ratio of *Cretoxyrhina mantelli* was inferred from two different caudal fin variables (Cobb’s angle and hypochordal ray angle, S1A Fig) measured on a well-preserved specimen (CMN 40906, S1B Fig). The relationship between such metric variables and \(AR\) was previously established in living lamniform sharks by linear regression from data provided in Kim et al. [50] (see S5 Table).

### Cost of swimming and energy budget inferences

The energy budget of *Cretoxyrhina mantelli* (assessed from its routine metabolic rate) was compared with independent inferences of its locomotion energy requirements (i.e., net cost of swimming), considering different water temperatures and both ectothermy and regional endothermy. The potential habitable temperature range (i.e., temperatures at which energy budget exceeds locomotion requirements) was then evaluated for each thermophysiological strategy and compared with pre-existent paleoclimatic data and the paleobiogeographic distribution of this shark. This allowed the plausibility of the ectothermic and the endothermic scenarios to be contrasted.

### Table 2. Tooth crown heights (CH) and total body length (TL) estimates of otodontids and cretoxyrhinids

| Taxa         | Source                        | CH (mm) | Tooth position | TL (m)  |
|--------------|-------------------------------|---------|----------------|---------|
| Otodus (M.)  | [46]: Suppl. Mat. (UF 257579) | 41.2    | L5-L7          | 17.90*  |
| Otodus (O.)  | [1]: fig. 208A                 | 65.0    | a1-a2          | 9.18*   |
| Parotodus    | [1]: fig. 211A                 | 48.0    | a1-a2          | 6.76*   |
| Megalolamna  | [47]: fig. 2L                  | 38.8    | A1-A2 or a1-a2 | 5.02*   |
| Cretoxyrhina | [43]                          | -       | -              | 6.40    |
| Cretolamna   | [44]                          | -       | -              | 3.00    |

* Total length estimates calculated from the average of length estimates for the different positions where each tooth could have belonged.

https://doi.org/10.1371/journal.pone.0185185.t002
Net cost of swimming. Power-performance curves establish the relationship between relative swimming speed (U, body lengths\(^{-1}\)) and oxygen consumption (MO\(_2\), mgO\(_2\) kg\(^{-1}\) s\(^{-1}\)). Net costs of swimming at cruising speed (NCS) were calculated in *Cretoxyrhina* from power-performance curves of living sharks as the difference between the total metabolic rate (TMR, oxygen consumption at a particular swimming speed) and the standard metabolic rate (SMR, oxygen consumption at resting) (S2 Fig). Estimates were obtained from the equations established in *Carcharhinus acronotus* [51] (S2A Fig) and *Isurus oxyrinchus* [52] (S2B Fig), considered as ectothermic and regional endothermic oxygen consumption models respectively (see S1 Text for a detailed justification of this selection):

\[
\log(MO_2)_{C. acronotus} = 2.38 + 0.377U
\]

from which it is deduced that

\[
TMR_{(U>0)C. acronotus} = 10^{(2.38 + 0.377U)}
\]

\[
SMR_{(U=0)C. acronotus} = 10^{2.38}
\]

\[
\log(MO_2)_{I. oxyrinchus} = 2.0937 + 0.97U
\]

from which it is deduced that

\[
TMR_{(U>0)I. oxyrinchus} = 10^{(2.0937 + 0.97U)}
\]

\[
SMR_{(U=0)I. oxyrinchus} = 10^{2.0937}
\]

Where MO\(_2\) is oxygen consumption in mgO\(_2\) kg\(^{-1}\) s\(^{-1}\), U is relative swimming speed in body lengths\(^{-1}\) s\(^{-1}\), TMR is total metabolic rate in mgO\(_2\) kg\(^{-1}\) s\(^{-1}\), and SMR is standard metabolic rate in mgO\(_2\) kg\(^{-1}\) s\(^{-1}\).

Thus, NCS of *Cretoxyrhina* at cruising speed assuming ectothermy and regional endothermy was calculated as:

\[
NCS_{Ectothermic \ Cretoxyrhina} = TMR_{Ectothermic \ Cretoxyrhina} - SMR_{Ectothermic \ Cretoxyrhina}
\]

\[
NCS_{Ectothermic \ Cretoxyrhina} = [10^{(2.38 + 0.377Uc)}] - (10^{2.38})
\]

\[
NCS_{R. \ endothermic \ Cretoxyrhina} = TMR_{R. \ endothermic \ Cretoxyrhina} - SMR_{R. \ endothermic \ Cretoxyrhina}
\]

\[
NCS_{R. \ endothermic \ Cretoxyrhina} = [10^{(2.0937 + 0.97Uc)}] - (10^{2.0937})
\]

Where NCS is net cost of swimming in mgO\(_2\) kg\(^{-1}\) s\(^{-1}\), TMR is total metabolic rate in mgO\(_2\) kg\(^{-1}\) s\(^{-1}\), SMR is standard metabolic rate in mgO\(_2\) kg\(^{-1}\) s\(^{-1}\), and U\(_c\) is relative cruising swimming speed in body lengths\(^{-1}\) s\(^{-1}\) calculated from [48]'s model, see above.

The use of NCS was preferred here instead of TMR as they are considered a better parameter for assessing the energy expenditure of thrust generation during swimming [53] and seem to be independent of the water temperature [54–56].

Routine metabolic rate. Routine metabolic rate (RMR) is the mean metabolic rate measured in an animal that performs random physical activity over a given period [57]. Here, the scaling of RMR with body mass has been independently established in ectothermic and regional endothermic fishes and used to infer the RMR of *Cretoxyrhina* assuming both conditions. For this, 43 records of RMR of extant sharks and rays were compiled from the literature.
RMR data were temperature adjusted to 5˚C, 10˚C, 15˚C, 20˚C, and 25˚C using $Q_{10}$ of 2.3 (following [19]). RMR and body mass data were log-transformed and regression lines were fitted independently for points corresponding to ectothermic and regional endothermic species at each temperature. The RMR of Cretolamna was then inferred in all possible scenarios, assuming a body mass of 2655 kg (calculated from the exponential model proposed by Gottfried et al. [6] assuming a total body length of 6.4 m [43]).

**Validation Test.** Following the same procedure, the thermophysiological strategy of 17 living sharks, including 14 ectothermic species and 3 regional endothermic species, was assessed from simultaneous records of their cruise swimming speeds, water temperatures and body masses (S7 Table).

### Results and discussion

#### Evidence of regional endothermy in otodontids and cretoxyrhinids

**Oxygen isotopic approach.** The oxygen isotopic composition of bioapatite phosphate and carbonate has been extensively studied in vertebrates for different purposes (see [58] for a detailed review). Given that δ$_{18}$O value of a biomineral depends both on the δ$_{18}$O value of the body fluid from which it precipitates and the temperature at which it forms [58], some authors have used oxygen isotopic data from bone, dentine or enamel to infer high body temperatures or thermoregulation in fossil taxa (e.g., [37,59–62]). Recently, Bernard et al. [37] have established a useful approach for inferring the dependence of body temperature of extinct aquatic animals on that of the surrounding water (see Data source and methodology section), demonstrating thermoregulation in some Mesozoic marine reptiles. Following this methodology, the thermophysiology of cretoxyrhinids and otodontids has been explored here from the isotopic composition of fossil remains from 20 different sedimentary beds, ranging in age from the Cretaceous to the Miocene (Table 1 and S1 Table).

The regression line slopes obtained for the analysis of *Cretolamna-Otodus* species are significantly different from zero and their values are consistent with the existence of partial independence from sea water temperature (Fig 1A–1C). These results are comparable in some aspects with those provided by Bernard et al. [37], showing similar fits in the regression lines and similar standard deviation values for isotopic data (see Table 1 and compare with [37]: table S1). Furthermore, a significantly higher number of localities has been considered here, increasing the reliability of the regression results and allowing consistent conclusions. In addition, some considerations have been taken into account in order to remove or minimize possible biases related to preservation or ecological aspects. Firstly, although diagenesis may occur in a biomineral altering the δ$_{18}$O composition, enameloid is rather resistant to postdepositional alteration and recrystallization due to its dense and slightly porous structure [58]. Accordingly, almost all δ$_{18}$O data selected here come from measurements taken in tooth enameloid (except for two turtle osteocutes and a few taxa where the whole tooth was considered, S1 Table). In fact, there are previous Rare Earth Elements analyses [40,63–65] as well as taphonomic evidence [38,39] for almost all the teeth included in this analysis supporting the lack of diagenetic processes (see [66] for a detailed review on the applicability of REE analysis as a tool for detecting diagenetic processes). Moreover, Žigaite and Whitehouse [67] have recently studied the isotopic composition of different tooth tissues in extant sharks demonstrating that enameloid shows less δ$_{18}$O variation than dentine and recommended it as a target biomineral and as a preferential biogeochemical reference for environmental and paleoenvironmental studies (see also [68]). Secondly, δ$_{18}$O differences between teeth of co-occurring individuals can also reflect differences in water column temperature due to distinct water depth preferences [40,69,70]. In order to minimize this bias, isotopic data from benthic ectothermic vertebrates were discarded.
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Fig 1. Oxygen isotopic evidence and the effect of regional endothermic taxa in previous paleotemperature estimates. (A-C) Difference in the δ¹⁸O value between coexisting cretoxyrhinids/otodontids and ectothermic taxa in several localities/sedimentary beds plotted against the ectothermic taxa δ¹⁸O value (as a proxy of seawater temperature). Three different regression analyses have been performed: (A) including all cretoxyrhinids-otodontids and all associated ectothermic taxa, (B) including all cretoxyrhinids-otodontids and only associated pelagic ectothermic taxa; and (C) including only cretoxyrhinids-otodontids and associated pelagic ectothermic taxa from Kocsis et al. [38] (regression lines are shown with associated 95% confidence intervals). Details of each fossil locality (denoted by numbers) are given in Table 1 and S1 Table. (D) Regression analyses performed in living lamnid sharks for comparative purposes, considering direct water and body temperature records compiled in Lowe and Goldman [25] (DM, deep muscle; ST, stomach). In all cases, slope values close to -1 imply that body temperature is independent from that of the water, slope values close to 0 imply a complete dependence, and intermediate slope values indicate some degree of independence. (E-F) Campanian-Maastrichtian latitudinal gradients of δ¹⁸O and seawater temperature calculated from cretoxyrhinids and ectothermic taxa. (G-H) Campanian-Maastrichtian δ¹⁸O and seawater temperature estimates calculated for three different latitudinal ranges (11°-13°, 20°-23° and 36°-49°), considering only cretoxyrhinids (pink), all taxa (black) and only ectothermic taxa (green). Significance of pairwise mean contrasts are shown in each case. Data in E-H taken from Puceat et al. [42]. ** indicates significance at the 0.05 level and * indicates significance at the 0.1 level.

https://doi.org/10.1371/journal.pone.0185185.g001
from adjusted regression lines, leaving only taxa with similar ecology to cretoxyrhinids and otodontids as water temperature proxies. As expected, the effect of water column temperature variations became evident when a wide ecological range of taxa are considered, resulting in less well fitting regression lines including benthic taxa ($R^2 = 0.45$, Fig 1A) than excluding them ($R^2 = 0.68$, Fig 1B). Finally, individuals collected from the same sedimentary bed may not be strictly contemporaneous and differences in $\delta^{18}O$ values could be reflecting seasonality or climate change. Although this problem can never be ruled out, accurate stratigraphic knowledge of layers that yield the presupposed contemporaneous remains can minimize it. In this sense, the best regression line fit is obtained taking into account only taxa included in Kocsis et al. [38] ($R^2 = 0.85$, Fig 1C), where the layer of provenance of each specimen is specified accurately (note that the stratigraphic resolution of some of the other localities is given only at bed level, S1 Table).

The existence of some decoupling between the enamel $\delta^{18}O$ composition of cretoxyrhinids-otodontids and coetaneous ectothermic organisms implies the presence of more constant body temperatures in the head and jaws of the former. Body temperature decreases towards the exterior in extant regional endotherms and is almost equal to the surrounding water temperature on the body surface [71–74]. Thus, the influence of the changes of water temperature it is expected to be more noticeable on external body parts, such as the region where teeth calcify. Unfortunately, this influence has never been quantified in this area and direct comparisons with results obtained for cretoxyrhinids and otodontids are not possible. However, elevation in deep muscle and stomach temperature over ambient temperatures has been recorded in some species of extinct regional endothermic sharks [25]. Here, the influence of water temperature on the deep muscle of *Isurus* and *Lamna* and on the stomach of *Isurus* and *Carcharodon* has been quantified from measurements reported in previous works (S2 Table). The representation of these temperature data in the same way as the isotopic data allows a comparison between the slopes obtained in both cases given that $\delta^{18}O$ values reflect water and body temperatures indirectly (Fig 1D). The influence of surrounding water temperature on teeth in formation of cretoxyrhinids and otodontids (slope value of -0.66, Fig 1C) seems comparable to that on the stomach of *Isurus* and less important than on the muscle of *Lamna* (slope values of -0.59 and -0.37 respectively; Fig 1D). In any case, although calcifying teeth are embedded in relatively superficial tissues, this result might be expected as many regional endothermic fishes have cranial endothermy. Namely, lamnids, xiphids, istiophorids and some tunas are able to elevate the temperature of their brains and nearby structures ([75] and references therein). In lamnids this is achieved mainly by means of a red muscle vein that transfers heat from red muscles to brain and orbital retia, but also by the heat supplied by some aerobic muscles, such the eye muscles, and other metabolically active tissues [76]. Interestingly, Tubbesing and Block [76] noted that jaw muscle of *Lamna nasus*, composed of slow-twitch aerobic fibbers, could also aid in local heat generation, being a closer possible heat source for dental lamina. In sum, the presence of different mechanisms that contribute to tissue warming in the head of some lamniform sharks makes the existence of high temperatures in the area where teeth calcify likely, affecting the isotopic composition of enameloid in developing teeth. In fact, the effect of cranial endothermy on the isotopic composition of other calcified structures of the head has already been demonstrated in tuna otoliths [77]. However, studies focused on recording temperatures in dental lamina of the upper and lower jaw in regional endotherms are needed in order to determine the degree of temperature independence of these tissues and the possible influence of cranial endonthermy on the oxygen isotopic composition of teeth.

**Cruise and burst swimming speed inferences.** The scaling of fish locomotion has been comprehensively studied both in terms of steady and unsteady swimming, and it is well
known that cruise and burst relative speeds decrease with body length [78–82]. The reasons underlying this phenomenon are related to physiological constraints that imply lower muscle contraction rate when body mass increases [83,84], resulting in lower tail beat frequencies in larger fishes [79]. However, although the scaling of locomotion is well known for fishes in general, there are virtually no works comparing the relationship between swimming speed and body size in ectothermic and regional endothermic taxa (with the only exception of [85]). Here, fish swimming speed data have been compiled for extant species covering a wide range of body sizes (S4 Table) and this relationship has been studied and compared in both groups. ANCOVA analysis, adjusted regression lines and their associated confidence intervals strongly support the existence of higher cruise and burst swimming capabilities in regional endotherms (Fig 2A and 2B).

Some works have previously suggested differences in swimming speeds of regional endotherms and ectotherms on the basis of indirect evidence such as the increase in power output and muscular contraction rate at high body temperatures, the enhancing of the diffusion of oxygen to the muscle mitochondria and increasing lactate turnover ([18] and references therein). Nonetheless, direct records of swimming speeds are scarce and sometimes inconclusive given the size limitations of controlled experiments and the difficulty of taking measurements in the wild. Regarding cruising speeds, some experiments carried out in water tunnels suggested similar speeds for ectothermic and regional endothermic taxa [86–91]. However, the work of Watanabe et al. [85] has recently shed light on this, demonstrating that cruising speeds of fishes with regional endothermy are greater than fishes without it using free-swimming data. They argued that previous results obtained under non-free-swimming conditions may be explained by the small size of water tunnels that do not allow the study of big adults where the

Fig 2. Scaling of swimming speed in extant fishes and swimming speed inferences in cretoxyrhinids and otodontids. (A) Cruise and (B) burst relative swimming speeds (U, body lengths*s⁻¹) against body lengths (meters) of living ectothermic and regional endothermic fishes. Adjusted regression lines are showed with associated 95% confidence intervals. (C) Cruise and (D) burst swimming speed estimates (V, km*h⁻¹) of cretoxyrhinids and otodontids, considering them as ectothermic sharks (green) or regional endothermic sharks (pink), with associated 95% individual prediction intervals. Values of absolute (V, km*h⁻¹) and relative (U, BL*s⁻¹) speed estimates are also shown for each case. ** indicates significance at the 0.05 level.

https://doi.org/10.1371/journal.pone.0185185.g002
Regional endothermy is well developed. ANCOVA analysis performed here also suggest that differences in cruise swimming capabilities of ectothermic and regional endothermic fishes increase gradually with body size (Interaction, p = 0.002; Fig 2A). The intersection of adjusted lines occurs at 24 cm total length meaning that cruising speeds of both regional endotherms and ectotherms are equal at that size. Interestingly, Dickson [92] and Dickson et al. [93] have proposed similar lengths as minimum sizes of functional endothermy in fishes. A high surface area-to-volume ratio constrains the ability to maintain warm tissues in smaller individuals and endothermy cannot be completely developed until a threshold mass is reached [94]. On the other hand, works comparing burst swimming speeds of ectothermic and regional endothermic fishes from direct measurements are virtually non-existent. Some extraordinarily high speed records of small ectothermic sphyraenids and scombrids (9.5 BL\textsuperscript{-1} s\textsuperscript{-1} and 19.4 BL\textsuperscript{-1} s\textsuperscript{-1} for Sphyraena barracuda and Acanthocybium solandri respectively) equal or exceed the maximum speeds documented in regional endotherms of similar sizes (See S4 Table). These data and the methodology by which they were obtained have been sometimes questioned (see [95–99]).

However, high burst speeds could be expected for these and other tropical or subtropical species inhabiting warm waters as the contraction time of the anaerobic muscles involved in burst swimming becomes shorter when temperature rises [100]. In any case, here, ANCOVA analysis detects differences between burst speeds of both groups even when these bizarre speeds are taken into account (Interaction, p = 0.966; Ecto/Endo, p < 0.000; data not shown in Fig 2B).

Scaling relationships of cruise and burst swimming speeds for both ectotherms and regional endotherms fit well into linear models (Fig 2A and 2B). Comparatively low dispersion of the data has made it possible to obtain narrow individual confidence intervals allowing swimming speed predictions in new taxa accurate enough for the purpose of the present study. Cruise and burst swimming speed estimates have been obtained for extinct cretoxyrhinids and otophodonts from their total body lengths (Fig 2C and 2D). Previous estimates of swimming speeds of otophodonts have only been provided by Jacoby et al. [101], predicting a cruising swimming speed for Ototodus (Megaselachus) megalodon (1.34 m\textsuperscript{s}\textsuperscript{-1}) very similar to the values obtained here assuming regional endothermy (1.40 m\textsuperscript{s}\textsuperscript{-1}). On the other hand, there are no studies so far aiming to assess the burst swimming speeds of cretoxyrhinids and otophodonts, this work being the first attempt in this sense. Unsteady swimming locomotion (i.e., burst speed) is extremely important in prey-predator interactions, mainly when prey/predator length ratio is high (similar to or higher than 10\textsuperscript{-1}) and feeding strategy implies a whole body attack [102,103]. Otophodonts and cretoxyrhinids are thought to have been active macropredators hunting on relatively big and fast swimming prey. This interpretation is based in functional analyses of their dentitions [1,31], trophic level inferences from isotopic data [104] and direct evidence such as coprolites with fish remains [105] or bite marks, fractures and embedded teeth in fossil cetacean, sirenian and marine reptile bones [6,9,106–115]. As a most extreme case, some of the largest representatives of the group were potential predators of big-sized marine mammals ([16] and references therein), possibly exerting an important control on their communities and even playing a significant role in the evolutionary history of big filter-feeding whales [116] (although see also [115]). Consequently, it seems obvious that reaching high swimming speeds should be also crucial for the hunting success of cretoxyrhinids and otophodonts. In this sense, the range of burst swimming speeds inferred here considering them as ecoholic sharks (6.7–7.9 km\textsuperscript{h}\textsuperscript{-1}) seems to be too low for such active macropredators. This is especially drastic for the case of the biggest species because their absolute speeds correspond to extremely low relative speeds (e.g., 0.12 body lengths\textsuperscript{s}\textsuperscript{-1} in O. megalodon) and hunting success depends largely on the later [117]. In contrast, burst swimming speeds estimated considering both groups as regional endotherms (30.6–37.2 km\textsuperscript{h}\textsuperscript{-1}) appear to fit better with their presupposed lifestyles, probably being high enough for active pursuit and hunting of fast
prey (burst swimming speeds of extant big and medium-sized odontocetes, sirenians and otariids range approximately between 20 km$^{-1}$ and 30 km$^{-1}$ [118–121]).

In addition, the squamation pattern and some morphofunctional interpretations of cretoxyrhinids also support an adaptation to fast swimming, more congruent with values estimated for the scenario of regional endothermy. Shimada [122] noted that the body surface of Cretoxyrhina was covered by scales with parallel keels separated by U-shaped grooves where the average interkeel distance was approximately 45 microns (see [43]: fig 5 and [122]: fig 8B). These aspects evidence a clear role in drag reduction and allow the unequivocal assignation of Cretoxyrhina scales to a morphotype that is exclusive to the fastest living species of sharks [123–127]. In the same way, some other evident aspects of the squamation of Cretoxyrhina, such as the high density of scales together with a notable overlapping ([122]: fig. 8A and B) or the presence of scales with crown thinning ([43]: fig. 5A and C), have also been interpreted as an adaptation for enhancing hydrodynamic efficiency in fastest pelagic species ([128] and references therein). A few smooth rounded scales have also been described in Cretoxyrhina but they were probably restricted to the snout and possibly other regions exposed to high abrasive stress (scales of Type A in [122]). Similarly, functional interpretations of caudal fin remains also support the idea of Cretoxyrhina as a fast pelagic hunter. Specimen CMN 40906 exhibits the highest Cobb’s and hypochordal ray angles ever recorded in lamniform sharks (49˚ and 133˚ respectively), implying fast swimming capabilities in Cretoxyrhina [50]. Thomson and Simanek [129] noted that Cobb’s angles above 30˚ are characteristic of fast swimming pelagic sharks typified by the endothermic sharks within the family Lamnidae. In fact, the swimming speed calculated from morphological variables of the caudal fin of Cretoxyrhina ($\approx$70 km$^{-1}$, see S2 Text) is clearly incompatible with those predicted here under the assumption that it was an ectothermic shark ($\approx$7 km$^{-1}$). Finally, some other aspects, like the conical head and the shape and number of vertebra in Cretoxyrhina [122] and the absence of dorso-ventral flattening in Cretolamna vertebra [44], also point towards the existence of fusiform bodies in cretoxyrhinids compatible with an active pelagic lifestyle.

On the other hand, besides burst speed, some other unsteady swimming performance parameters, such as acceleration and manoeuvrability, are also size-constrained and decrease with body length [103]. For that reason, hunting strategies implying ambushing behaviour and/or attacks under specific ambient conditions could be reasonable for massive otodontids minimizing prey reaction time in a similar way to living great white sharks [130]. In fact, Godfrey and Altman [131], based on the analysis of a Miocene cetacean vertebra with a partially healed compression fracture, suggested that O. megalodon could display comparable predatory strategies to those of extant great white sharks. Similarly, predatory strategies implying group hunting are also effective when feeding on prey of similar or larger size than the predator [103]. However, although some modern marine vertebrates display examples of cooperative attacks (i.e., the killer whale Orcinus orca) (see for example [132]), this behavior has never been reported so far in big sharks and it is quite likely that O. megalodon was a solitary hunter [133]. In any case, any interpretation on the hunting strategies of large otodontids is highly speculative.

**Costs of swimming and energy budget inferences.** The comparison of locomotion costs with the metabolic ceilings of ectotherms and endotherms (i.e., energy budget) has been revealed as a useful approach for assessing the thermophysiological strategies of fossil groups [134]. Following a similar procedure, here the thermophysiology of Cretoxyrhina mantelli has been assessed by inferring and comparing estimates of its net costs of swimming and energy budget under ectothermic and regional endothermic scenarios.

Power-performance curves of continuously active elasmobranch species have been used by several authors for different purposes: (1) determining standard metabolic rates (SMR)
extrapolating to 0 velocity (e.g., [51, 135, 136]) and (2) determining field total metabolic rates (TMR) at a given measured speed [137, 138]. In the present study, for the first time, power-performance curves of living sharks have been used to estimate the locomotion energy requirements in an extinct taxon, Cretoxyrhina mantelli. A cruise swimming speed of 12 km h\(^{-1}\) (0.53 body lengths s\(^{-1}\)) has been inferred here for this species after applying Sambilay’s [48] model assuming a total body length of 6.4 m and aspect ratio of 4.3 (See Fig 3 and S2 Text for further details of the aspect ratio and swimming speed inferences). At this speed, the net cost of swimming is 373 gO\(_2\) kg\(^{-1}\) h\(^{-1}\) and 622 gO\(_2\) kg\(^{-1}\) h\(^{-1}\) assuming ectothermy and regional endothermy respectively (Table 3). On the other hand, energetic budgets and metabolic ceilings in animals are usually assessed by calculating sustained metabolic rate (SusMR, metabolic level that can be sustained long-term by an animal) [139]. Unfortunately, there are virtually no works dealing with SusMR of sharks and the data available in the literature are insufficient for establishing predictive models. However, SusMR is considered as broadly equivalent in ecological terms to another much more frequently measured parameter, the routine metabolic rate (RMR) [140, 141]. The well-fitted scaling relationships between the RMR and body mass obtained here for ectothermic and regional endothermic living sharks has allowed reliable predictions of RMR to be obtained in Cretoxyrhina (as an approximation of its metabolic budget) within narrow values of confidence (Table 3).

Metabolic ceilings have been proposed several times as constraining the distribution of plants and animals, accounting for their observed latitudinal and altitudinal limits [142–144]. Differences in thermal physiology between ectotherms and endotherms affect their global distributions differently, with ambient temperature being the main constraining for ectothermic organisms [145–147]. Accordingly, this work suggests differences in the potential habitable temperature range of Cretoxyrhina depending on its thermoregulatory capabilities. When
considering *Cretoxyrhina* as an ectothermic shark the model predicts that its energy expenditure of locomotion would not be sustainable over time in waters below 20˚C (note in Fig 4A that NCS exceeds RMR below this temperature in the ectothermic scenario). In contrast, such costs would be probably sustainable long-term by a regional endothermic shark in a wider range of water temperatures (RMR could exceed NCS at all the considered temperatures in the regional endothermic scenario, Fig 4A).

*Cretoxyrhina* remains have been documented from the Albian to the Cenomanian (Lower to Upper Cretaceous) over a period when sea surface mean temperatures were higher than nowadays and latitudinal gradients were less important [148]. Despite that, the fossil record of this shark reveals a cosmopolitan distribution with a paleolatitudinal range wide enough to suggest a broad tolerance to different water temperatures (see [1] for a summary of fossil occurrences). Paleobiogeographical data within the Western Interior Sea (USA) prove that *Cretoxyrhina* was able to inhabit conditions ranging from subtropical to cool temperate waters [149] and probably to migrate through the Boreal ocean [150]. In fact, the most northern fossil assemblages where *Cretoxyrhina* remains have been recovered demonstrate that this shark was, indeed, able to inhabit circumboreal seas where the mean annual surface temperature has been estimated to have been between 5˚C and 10˚C according to [151]. In consequence, the scenario of *Cretoxyrhina* as an ectothermic shark does not fit the paleobiogeographical and paleoclimatic evidence. The model presented here predicts that, in the absence of regional endothermy, only swimming speeds around or below to 0.1 body lengths s$^{-1}$ would be sustainable long-term for sharks of comparable body mass at 5˚C (Fig 4A). Accordingly, very similar speeds have been reported for *Somniosus microcephalus* (0.11 body lengths s$^{-1}$ and 1.22 km h$^{-1}$) [152], the biggest ectothermic shark inhabiting polar waters [153]. On the contrary, the scenario that assumes regional endothermy seems to be more congruent with all these aspects. In that case, the temperature at which the NCS of *Cretoxyrhina* equates its RMR (∼5˚C; Fig 4A) fits well within the lowest water temperatures inhabited by this taxon according to its northernmost distribution limit (5˚C–10˚C according to [151]).

Interestingly, a validation test performed in 17 living sharks from simultaneous records of their cruise swimming speeds, water temperatures and body masses supports the notion that this approach has a high predictive power, as it is able to correctly predict the thermophysiological strategy of the vast majority of the species (Fig 4B). Energetic viability of ectothermy is supported for all the 14 ectotherm species studied with no exception (inferred NCS lay below

### Table 3. Predicted values of net cost of swimming (NCS) and routine metabolic rate (RMR) in *Cretoxyrhina mantelli* assuming ectothermy and regional endothermy at five different scenarios of water temperature.

| Predicted logNCS (mgO$_2$·kg$^{-1}$·h$^{-1}$) | Scenario | R$^2$ | Predicted logRMR (mgO$_2$·kg$^{-1}$·h$^{-1}$) |
|-----------------------------------------|----------|-----|---------------------------------|
| Cret.                                    |          |     |                                 |
| 5.572                                    | Ectothermy| 0.97| 4.716 ± 0.344                   |
| 5.572                                    | Ectothermy| 0.97| 4.897 ± 0.344                   |
| 5.572                                    | Ectothermy| 0.97| 5.078 ± 0.344                   |
| 5.572                                    | Ectothermy| 0.97| 5.259 ± 0.344                   |
| 5.572                                    | Ectothermy| 0.97| 5.440 ± 0.344                   |
| 5.794                                    | R. endothermy| 0.95| 5.780 ± 0.486                   |
| 5.794                                    | R. endothermy| 0.95| 5.961 ± 0.486                   |
| 5.794                                    | R. endothermy| 0.95| 6.142 ± 0.486                   |
| 5.794                                    | R. endothermy| 0.95| 6.323 ± 0.486                   |
| 5.794                                    | R. endothermy| 0.95| 6.504 ± 0.486                   |

https://doi.org/10.1371/journal.pone.0185185.t003
or within the confidence intervals of the estimated RMRs). On the other hand, regional endothermy is predicted as the only energetically sustainable strategy in two regional endotherms, *Lamna ditropis* and *Carcharodon carcharias* (note that ectothermy cannot be rejected as a sustainable strategy in a third regional endothermic taxon, *Isurus oxyrinchus*). Thus, the
recognition of the ectothermic scenario as a non-energetically viable possibility is highly indicative of regional endothermy, this being the case for *Cretoxyrhina mantelli*.

**Implications of the regional endothermy of otodontids and cretoxyrhinids**

Three independent lines of evidence (i.e., isotopic data, swimming speed estimates and metabolic inferences) strongly support the starting hypothesis of this work, strengthening the evidence for the existence of regional endothermy in otodontids and cretoxyrhinids. The detection of this adaptation in such extinct groups entails a series of implications, some of them with important multidisciplinary significance.

**Implications for sea surface paleotemperature estimations.** Several works have used the oxygen isotope composition of vertebrate remains for paleotemperature inferences of past seawater (e.g., [40–42,69,154–160] and many more). Most of them have been carried out by analyzing the enameloid from teeth and scales of fishes, without an exhaustive selection of the included taxa, assuming a unique fractionation equation applicable to all the species [41]. However, the presence of regional or whole endothermy in some fish lineages could affect the oxygen isotopic fractionation in their biominerals [58]. In fact, a number of studies have empirically demonstrated differences in the $\delta^{18}$O isotopic composition of mineralized tissues of coexisting taxa due presumably to differences in thermoregulatory physiology (e.g., [37,39,62]). In this sense, it is striking that several works aiming to reconstruct paleotemperatures of seawater [39,156,158] or interpret hydrographic changes [64] have ignored this fact, including $\delta^{18}$O isotopic data from taxa belonging to groups with regional endothermy, such as lamnids (e.g., *Carcharodon*, *Isurus*, *Isuralamna*, *Cosmopolitodus* and *Macrorhizodus*) or alopids (e.g., *Usakias*). Likewise, $\delta^{18}$O isotopic composition of teeth of otodontids and cretoxyrhinids has also been used for similar purposes (see for example [38–42,64,154–156,158,161]), without taking into account that these groups are potential regional endotherms given their close phylogenetic relationship with lamnids. The presence of cretoxyrhinids is especially frequent in works dealing with paleoenvironmental reconstructions of the Late Cretaceous—Paleocene as a result of the high abundance of *Cretolamna* teeth in sediments of that age. For instance, among the 108 fossil remains considered by Kocsis et al. [38], 30 are teeth belonging to *Cretolamna* and *Otodus*, constituting 28% of the total sample and up to 50% in some of the studied localities. Similarly, 24% of all teeth analyzed by Pucéat et al. [42] belong to cretoxyrhinids (*Cretolamna* and *Archaelomana*) and otodontids (*Otodus*). Among these works, the study carried out by Amiot et al. [39] is probably one of the most remarkable cases where only two (15%) of all included shark teeth belong unquestionably to ectothermic groups, whereas the remaining 11 teeth belong to otodontids (8%) and lamnids (77%). In addition, some of these studies include shark teeth with poor taxonomic determination meaning that it is not possible to rule out, in most cases, the possibility that they belong to one of the regional endothermic groups.

The evidence provided in the present work in favour of cretoxyrhinids and otodontids as regional endothermic sharks imply that some of the previous paleotemperature reconstructions based partially on these taxa could be overestimated and should be revised. As proof of this, the effect of including representatives of such groups in this kind of studies has been tested here considering the isotopic data provided by Pucéat et al. [42]. ANCOVA analyses support this prediction detecting significant differences both in the $\delta^{18}$O values and temperatures calculated from remains of cretoxyrhinids and ectothermic taxa at different latitudes ($\delta^{18}$O, $p = 0.022$; Temperature, $p = 0.013$; Fig 1E and 1F). Parameter estimation of ANCOVA analysis indicates that temperatures inferred from cretoxyrhinids are 2.3°C above those predicted from ectothermic fishes (Fig 1F). These differences are expected to be more important
in high latitudes where body temperature of regional endothermic animals is much higher than that of the surrounding water (see for example [162,163]). However, no significant differences have been found in the slope of ecothermic taxa and cretoxyrhinids (Interaction, $p = 0.505$; Fig 1F), possibly due to the small sample size of the latter in latitudes above 30°. In any case, the effect of cretoxyrhinids on previous sea paleotemperature reconstructions becomes clear when temperatures are re-estimated removing the isotopic data of such taxa from the analysis. Overestimations of up to 1.9°C have been calculated here for latitudes around 20° where a large number of *Cretolamna* teeth were taken into account (Note that no significant differences have been detected in latitudes around 10° and 40° probably due to the smaller sample size of cretoxyrhinids considered in those cases, Fig 1H). Hence, in view of these results, remains of regional endothermic taxa or closely related groups should be treated more cautiously in future studies dealing with seawater paleotemperature reconstructions.

**Evolutionary scenario of regional endothermy in lamniforms.** Among living fishes, endothermy is present in several orders and it is thought to have evolved independently at least six times (in lamnids, billfishes, the butterfly mackerel, tunas, alopids and lamprids [18,21,23]); but probably more if taking into account some other possible regional endotherms included within mobulids and labrids [164–166]. Here the presence of regional endothermy is strongly supported in two additional groups of extinct sharks, Cretoxyrhiniidae and Otodontidae, pushing the appearance of this adaptation among fishes back to the Albian (Late Cretaceous) [1] (note that some authors [11,12] have also suggested the possible existence of endothermy in a few Paleozoic groups). Interestingly the evolution of regional endothermy in extant fish families is thought to occur much later, during the Eocene coinciding with a major cooling trend in the global climate ([18]; although see below). The monophyly of some cretoxyrhinids and otodontids is widely accepted (*Cretolamna*-otodontid lineage) [1,9,31,47,167–171], thus a common origin for the regional endothermy of both groups is the most parsimonious option. However, phylogenetic affinities between these extinct families and living lamniform groups are still controversial and, in consequence, it is difficult to establish the evolutionary relationship between the regional endothermy of cretoxyrhinids and otodontids and that of alopids and lamnids. Otodontids have usually been regarded as lamnids (e.g., [6,9,26–29]), as their sister-group, or as a closely related group (e.g., [1,5,30,31–36]). In fact, both groups may have evolved from *Cretolamna* species, sharing a common ancestor within cretoxyrhinids [1,9,26,31]. At present, monophyly of cretoxyrhinids is not sustained and they are considered as a paraphyletic or even polyphyletic group although their systematics is maintained awaiting further results [1,172,173]. Cappetta [1] provisionally included five genera within this family, of which *Cretolamna* and *Cretoxyrhina* are the best known by far. The phylogenetic position of *Cretolamna* among the extant lamniform taxa remains largely unknown [44]. A number of works have proposed *Cretolamna* as being the “ancestor” of otodontids [1,9,31,47,167–171], *Cretoxyrhina* [1] and even of lamnids [1,9,26,31] on the basis of dental characters. Similarly, the phylogenetic position of *Cretoxyrhina* is also debated, having been included within cretoxyrhinids traditionally [1], within alopids following the only cladistical approach on the group [174] and recently, after the discovery of one Cretaceous articulated specimen of *Isurus denticulatus*, *Cretoxyrhina* has been placed among the earliest species of the *Isurus* lineage as a lamnid [175]. In sum, several possible scenarios about the evolution of regional endothermy within lamniforms can be drawn (Fig 5), some of them implying the independent evolution of this adaptation up to three (or possibly more) times. Independent evolution of the regional endothermy of alopids seems to be clear on the basis of their phylogenetic position [176] and some morphological aspects of their orbital retia [17] (Fig 5A). Lower Eocene remains of *Alopias denticulatus* represent the earliest record of Alopidae [173], however both the stratigraphic range and the origin of regional endothermy in alopids could be
Fig 5. Possible evolutionary scenarios for the origin of regional endothermy in lamniforms considering (A) only extant lamniform groups, (B) *Cretoxyrhina* within alopids, and (C) *Cretolamna* representatives as ancestors of both lamnids and otodontids, and *Cretoxyrhina* within lamnids. Stratigraphic ranges taken from [1].

https://doi.org/10.1371/journal.pone.0185185.g005
extended back to the Late Cretaceous if *Cretoxyrhina* is included within the family (according to [174]) (Fig 5B). On the other hand, given the relatively close phylogenetic affinity between lamnids and otodontids, it is rather parsimonious to consider the regional endothermy of both groups to be homologous and have evolved at least during the Late Cretaceous (if including *Cretolamna* representatives as ancestors and/or considering *Cretoxyrhina* within *Isurus*; Fig 5C). However, different scenarios could be also possible and independent evolution of regional endothermy in cretoxyrhinids cannot be ruled out given the possible polyphyly and poorly resolved phylogenetic relationships of these groups. Therefore, the finding of further articulated remains that allow robust cladistic analyses (not only based on dental characters) are needed to unravel the affinities of otodontids and cretoxyrhinids and, in consequence, the origin of regional endothermy in lamniforms.

Several hypotheses have been proposed to explain the evolution of regional endothermy in living groups (see [18] for a detailed review). Among them, the thermal niche expansion hypothesis was supported by Dickson and Graham [18] comparing the thermal niche of extant taxa, their fossil record, and the paleoceanographic conditions during the time that endothermic fishes radiated. According to their ideas, oceanic cooling and tropical compression during the Eocene were the promoters of the almost simultaneous independent evolution of regional endothermy in several groups of fishes. However, the discovery of regional endothermy among cretoxyrhinids could extend the stratigraphic range of some regional endotherm lineages (e.g., lamnids and alopids) back to the Cretaceous thus calling into question this hypothesis. More recently, the elevated cruising speed hypothesis has gained support with the development of modern animal-tracking tools. In this sense, Watanabe et al. [85] have demonstrated in free swimming fishes that regional endotherms have higher cruise swimming speeds and exhibit larger-scale annual migrations than ectotherms. This increases prey encounter rates, enables larger-scale annual migrations and allows potentially greater access to seasonally available resources. The selection pressures that lead to the evolution of regional endothermy in cretoxyrhinids are difficult to ascertain. Current evidence only allows for speculation about this issue and for some possible scenarios to be suggested for testing in future studies. The thermal niche expansion hypothesis seems unlikely for this group as the first reports of both *Cretolamna* and *Cretoxyrhina* from the Albian (Late Cretaceous) [1] are coincident with an extremely warm climate and markedly reduced vertical and latitudinal temperature gradients (see for example [148,177]). On the other hand, the worldwide distributions of cretoxyrhinids [1] and morpho-functional interpretation of the caudal fin morphology of *Cretoxyrhina* (see above) support good migratory capacities and strong swimming capabilities compatible with the elevated cruising speed hypothesis. Also, regional endothermy could have provided cretoxyrhinids with several evolutionary advantages over their ectothermic prey and competitors ([18] and references therein). Interestingly, some groups of marine reptile predators underwent a rapid radiation and diversification during the Late Cretaceous [178,179] a few million years after the appearance of the first cretoxyrhinids [1]. Such evolutionary success has sometimes been explained by the acquisition of endothermy providing them with some advantages during the Late Cretaceous climate cooling [37,62]. Thus, the progressive cooling of the oceans [180] together with increasing competition with marine reptile top-predators might also be suggested as major promoters of the evolution of regional endothermy in cretoxyrhinids (although see [181]). In any case, several selection pressures could have worked together to promote the evolution of regional endothermy in fishes and, at the same time, different evolutionary scenarios could be expected in each separate group.

Regarding the anatomical and physiological aspects of regional endothermy, diverse structures and mechanisms are involved in heat production and maintenance of high body temperatures in extant regional endothermic groups [18]. The physiological and anatomical
adaptations that allowed high body temperatures in cretoxyrhinids and otodontids are difficult to determine. The large size and low surface area-to-volume ratio expected for some otodontids might suggest a passive accumulation of heat (gigantothermy). However, although this phenomenon could contribute to heat retention in these huge taxa, it seems unlikely that it played a major role in smaller forms such as *Cretolamna* where other active mechanisms are expected to be present. Considering then that *Cretolamna* occupies a basal position to (or within) otodontids [47], it is expected that such mechanisms were inherited by all the species of otodontids. On the other hand, the detection of relatively low $\delta^{18}O$ values in teeth of cretoxyrhinids and otodontids implies the existence of high temperatures in the anterior part of their bodies, consistent with the presence of cranial endothermy. Based on the apparent dependence of cranial endothermy on red muscle endothermy, Dickson and Graham [18] argued that the former should have evolved after or at the same time as the latter. Accordingly, regional endothermy could be present in lamniforms even earlier than proposed here. In this sense, performing isotopic analyses on postcranial endoskeletal elements (e.g., vertebra) could be interesting for testing the existence of other sources of heat within the trunk region, compatible with the presence of red muscle endothermy.

**Distribution patterns, extinction and the evolution of gigantism in the megatooth sharks.** The existence of an ectothermic physiology and/or low tolerance to cold waters in otodontids has sometimes been proposed as a constraining factor on their distribution patterns and as an underlying cause of their extinction [6,9,10,18]. Gottfried et al. [6] and Diedrich [10] suggested that differences in prey and water temperature preference in megatooth and white sharks led to the extinction of the former in the Pleistocene due to climate change and ocean cooling. Segregated distribution patterns of both taxa, with otodontids and white sharks inhabiting warmer and colder waters respectively, have been sometimes explained by competitive exclusion of the latter sharks from the preferred warmer areas of giant-toothed species [9]. Similarly, Dickson and Graham [18] interpreted the differential success of white and megatooth sharks as a consequence of the differences in the thermal physiology of both taxa. Accordingly, the regional endothermy of white sharks allowed them to face oceanic cooling during the Pliocene-Pleistocene successfully, whereas some supposed ectothermic taxa such as *O. megalodon* became extinct. On the other hand, shifts in the distribution of large marine mammals to colder high-latitude conditions have also been proposed as possible causes of the extinction of otodontids [6]. However, Pimiento et al. [16] have recently demonstrated that the range of water temperatures inhabited by *O. megalodon* was wider than previously expected being rather consistent with the existence of regional endothermy in this taxon. Furthermore, they noted that the occupancy range of *O. megalodon* was not correlated with climatic changes thus suggesting that extinction of otodontids was not primarily driven by climatic change and ocean cooling. Results presented here strongly support the existence of regional endothermy in the *Cretolamna*-otodontid lineage making it necessary to integrate this phenomenon and its consequences (e.g., tolerance to a wider range of water temperatures and less vulnerability to ocean cooling) in new explanatory hypotheses about the extinction of otodontids. In this sense, hypotheses that involve global habitat loss produced by sea-level oscillations during the Pliocene [182] and/or biotic factors such as the drop in the diversity of potential prey (filter-feeding whales) or the appearance of new competitors (large predatory whales and the great white shark) could be more in agreement with all these aspects ([16,115] and references therein).

On the other hand, previous interpretations regarding the evolution and maintenance of gigantism in the biggest otodontids have been supported almost exclusively by the availability of high energetic blubber-rich marine mammals as prey. Gottfried et al. [6], based on the analysis of bite marks in fossil cetacean bones and previous dietary studies in extant white sharks,
suggested that the giant megatooth sharks could have relied on large blubber-rich marine mammals that may constituted an important part of their diet. Purdy [9] also noted that marine mammals could have constituted an important part of otodontid diet on the basis of positive correlation between the abundances of fossil remains of both groups. Regarding evolutionary intrinsic mechanisms that allowed gigantism in otodontids, Ehret [5] described acceleration of growth rates and delay in the offset timing of somatic growth in some otodontids but proposed again the evolution and diversification of cetaceans as the most plausible driver of these changes. However, although the availability of potential prey played a fundamental role in this regard, all these previous hypothesis obviated some other intrinsic aspects that should also be fundamental for maintaining an active lifestyle with such large body mass.

In this sense, Ferrón et al. [11] have recently provided key ideas to understand the evolution of gigantism in active vertebrate predators from a metabolic perspective. Activity and feeding strategies of living organisms are limited by specific values of mass-specific metabolic rate (i.e., metabolic ceilings). Given that mass-specific metabolic rate decreases when the body mass increases, active macropredation is not sustainable once a given body mass is reached and only less active modes of life and feeding strategies (e.g. filter feeding) are physiologically affordable above this size. However, this limit is reached at different body masses depending on thermoregulatory strategy and, ultimately, metabolic level; endothermic macropredators can attain larger potential body masses than their ectothermic counterparts (Fig 6A). Interestingly, shifts towards higher metabolic levels, promoted by different factors (i.e., increases in ambient temperature and atmospheric oxygen levels, evolution of highly efficient respiratory systems or acquisition of endothermy), can allow the same activity level and feeding strategy to be sustained at larger body masses, offering a suitable explanation for the evolution of gigantism in active macropredators (see Ferrón et al. [11] for further discussion) (Fig 6B). Among these factors, endothermy seems to have played an important role in the evolution of gigantism in many extinct macropredatory groups including some dinosaurs [134,183,184], ichthyosaurs, plesiosaurs and mosasaurs [37,62]. Ferrón et al. [11] also suggested that a number of extinct macropredatory fishes, including some members of the family Otodontidae, could have been regional or whole endotherms considering their presumed active lifestyle and big body sizes (Fig 6C). Now, regional endothermy of otodontids is strongly supported here from multiple approaches, and is confirmed as the key element that promoted the metabolic shift needed to reach huge sizes as macropredators in this group. Therefore, the integration of ecological and physiological triggers (i.e., availability of blubber-rich prey and endothermy inherited from cretoxyrhinids) offers a more holistic hypothesis to explaining the evolution of gigantism and the maintenance of active modes of life in this shark lineage.

The gigantic sizes reached by some otodontids were the result of an evolutionary tendency towards larger body sizes [46]. According to the ideas presented in Ferrón et al. [11], a trend to less active lifestyles in bigger species would be expected if all otodontids shared the same metabolic level. This phenomenon is well documented in the evolution of mysticetes where extinct small basal forms were actively carnivorous whereas more derived giant species are slow filter feeders (and intermediate-sized forms were probably facultative filter feeders) [185] (Fig 6D). However, the existence of similar active macropredatory lifestyles in all species of otodontids implies that more derived and larger forms would have had higher metabolic levels and, therefore, the tendency towards gigantism should be accompanied by a tendency towards a better-developed endothermy (Fig 6E). The factors that promoted this tendency towards gigantism are poorly understood. Pimiento and Balk [46] have recently argued that this phenomenon could be the result of a long-term selective pressure on otodontids favouring bigger forms with a broader range of prey. On the other hand, the fact that otodontids grew in size proportionally to cetaceans during the Neogene [10] could suggest that gigantism coevolved in both groups as
Fig 6. (A) Diagram showing the relationship between body mass, metabolic rate (≈ activity level and feeding strategy) and metabolic level (ecto, meso and endothermy) in aquatic vertebrates. (B) Schematic explanation of how shifts towards higher metabolic levels, promoted by different factors, contribute to maintaining a predatory lifestyle at bigger body sizes. (C) Diversity of body masses, feeding and thermoregulatory strategies of living and extinct aquatic vertebrates (ed, endotherm; ec, ectotherm; me, mesotherm; cross sign denotes an extinct taxon). A-C modified from Ferron et al. [11]. (D-E) Diagrams showing body sizes, feeding and thermoregulatory strategies of mysticete cetaceans and otodontid sharks respectively. Mysticete skull and otodontid tooth outlines modified from Fitzgerald [185] and Pimiento and Balk [46]. Bluish and reddish tones represent lower and higher metabolic levels, respectively.

https://doi.org/10.1371/journal.pone.0185185.g006
a result of an evolutionary race between predators and prey. In either case, the acquisition of
anatomical structures and physiological mechanisms that allowed regional endothermy in cre-
toxyrhinids probably played a key role in the subsequent evolutionary history of the lineage,
acting as a trigger for the evolution of gigantism in more derived otodontids, allowing active
macropredatory lifestyles and helping them in the struggle for survival.

Supporting information

S1 Table. Detailed δ¹⁸O data of cretoxyrhinids-otodontids and associated ectotherms compiled for this study.
(XLSX)

S2 Table. Simultaneous records of internal body organs/tissues and surrounding water temperature in three extant species of regional endothermic sharks. Data from Lowe and Goldman [7].
(XLSX)

S3 Table. δ¹⁸O tooth values of cretoxyrhinids and ectothermic fishes from several Campanian-Maastrichtian localities and latitudes. Data from Pucéat et al. [6].
(XLSX)

S4 Table. Records of cruise and burst swimming speed of living ectothermic and regional endothermic fishes. Standard Lengths (SL) and Fork Lengths (FL) has been transformed into Total Lengths (TL) using equations from Froese and Pauly [8] (equations from Kohler et al. [9] were also used but only for the case of Isurus oxyrinchus). Records denoted with an asterisk were not considered for estimating swimming speeds of cretoxyrhinids and otodontids as these data and the methodology by which they were obtained have been sometimes questioned (see Results and discussion section).
(XLSX)

S5 Table. Cobb’s angle, hypochordal ray angle and aspect ratio measurements of several living lamniform shark species and the fossil taxon Cretosyrhina mantelli (data from Kim et al. [83]).
(XLSX)

S6 Table. Routine metabolic rate (RMR) of several living ectothermic and regional endothermic sharks and rays adjusted to five different temperatures (5˚C, 10˚C, 15˚C, 20˚C, and 25˚C).
(XLSX)

S7 Table. Records of cruise swimming speeds, water temperatures and body masses of living ectothermic and regional endothermic sharks. Data taken from Watanabe et al. [17].
(XLSX)

S1 Fig. (A) Caudal fin variables used for aspect ratio inferences of Cretosyrhina mantelli. Cobb’s angle (Ca) describes the curvature of the vertebral column segment comprised between the anterior-most caudal vertebra (acv) and the posterior caudal vertebra (pcv); Hypochordal Ray angle (HRA) describes the orientation of the longest hypochordal ray of the caudal fin (diagram modified from Kim et al. [83]). (B) Photographs of the well-preserved C. mantelli specimen CMN 40906 from Shimada et al. [84]: figs. 1 and 4; courtesy of the New Mexico Museum of Natural History & Science). Scale bar equals to 30 cm and 10 cm in the complete and enlarged view of the specimen respectively.
(TIF)
S2 Fig. Power-performance curves relating the oxygen consumption (MO2, mgO2 kg⁻¹ h⁻¹) to relative swimming speed (U, l·s⁻¹) of (A) Carcharhinus acronotus (from Carlson et al. [85]) and (B) Isurus oxyrinchus (from Sepulveda et al. [86]), used in this study as models of ectothermic and regional endothermic sharks for metabolic inferences in Cretoxyrhina mantelli. NCS, Net cost of swimming; SMR, Standard metabolic rate; TMR, Total metabolic rate.

S1 Text. Power performance curves of small sharks as a tool for assessing the metabolic rate of bigger species.

S2 Text. Aspect ratio and swimming speed inferences in Cretoxyrhina.

S3 Text. References of supplementary information.

Acknowledgments
I would like to thank Dr. John Cunningham (School of Earth Sciences, University of Bristol) for his help in the English editing of the manuscript and the New Mexico Museum of Natural History & Science for supplying the pictures of Cretoxyrhina mantelli specimen CMN 40906 (shown in S1B Fig). I acknowledge the comments of Dr. Geerat J. Vermeij (Academic Editor), Dr. John Maisey (referee) and an anonymous reviewer that have considerably improved the final manuscript.

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References
1. Capetta H. Chondrichthyes. Mesozoic and Cenozoic Elasmobranchii: Teeth. Handbook of Paleoichthyology Volume 3E. Verlag Dr Friedrich Pfeil, München; 2012.
2. Leriche M. Les poissons Neogenes de la Belgique. Mémoires du Musée royal d'histoire naturelle de Belgique. 1926; 32: 365–472.
3. Bendix-Almgreen SE. Carcharodon megalodon from the Upper Miocene of Denmark, with comments on elasmobranch tooth enameloid: coronoid. Bulletin of the Geological Society of Denmark. 1983; 32: 1–32.

4. Uyeno T, Sakamoto O, Sekine H. The description of an almost complete tooth set of Carcharodon megalodon from a Middle Miocene bed in Saitama Prefecture, Japan. Bulletin—Saitama Museum of Natural History. 1989; 7: 73–85.

5. Ehret DJ. Paleobiology and taxonomy of extinct lamnid and otodontid sharks (Chondrichthyes, Elasmobranchii, Lamniformes). Ph.D. thesis, University of Florida. 2010.

6. Gottfried MD, Compagno LJ, Bowman SC. Size and skeletal anatomy of the giant megalooth shark Carcharodon megalodon. In: Klimley AP, Ainley DG, editors. Great White Sharks: The Biology of Carcharodon carcharias. 1996. pp. 55–66.

7. Randall JE. Size of the great white shark (Carcharodon). Science. 1973; 181: 169–170. https://doi.org/10.1126/science.181.4095.169 PMID: 17746627

8. Shimada K. The relationship between the tooth size and total body length in the white shark. Journal of Fossil Research. 2002; 35: 28–33.

9. Purdy RW. Paleoecology of fossil white sharks. In: Klimley AP, Ainley DG, editors. Great White Sharks: The Biology of Carcharodon carcharias. 1996. pp. 67–78.

10. Diedrich CG. White and megalooth shark evolution and predation origin onto seals, sirenians and whales. Natural Science. 2013; 5: 1203–1218.

11. Ferrón HG, Martínez-Pérez C, Botella H. The evolution of gigantism in active marine predators. Historical Biology. 2017; https://doi.org/10.1080/08912963.2017.1319829

12. Maisey JG, Bronson AW, Williams RR, McKinzie M. A Pennsylvanian ‘supershark’ from Texas. Journal of Vertebrate Paleontology. 2017; e1325369.

13. Alexander RM. All-time giants: the largest animals and their problems. Palaeontology. 1998; 41: 1231–1245.

14. Ebert D, Fowler S, Compagno L. Sharks of the world. Wild Nature Press, Plymouth; 2013.

15. Jefferson TA, Webber MA, Pitman RL. Marine mammals of the world: a comprehensive guide to their identification. Academic Press, London; 2015

16. Pimiento C, MacFadden BJ, Clements CF, Varela S, Jaramillo C, Velez-Juarbe J, et al. Geographical distribution patterns of Carcharocles megalodon over time reveal clues about extinction mechanisms. Journal of Biogeography. 2016; 43: 1645–1655.

17. Block BA, Finnerty JR. Endothermy in fishes: a phylogenetic analysis of constraints, predispositions, and selection pressures. Environmental Biology of Fishes. 1994; 40: 283–302.

18. Dickson KA, Graham JB. Evolution and consequences of endothermy in fishes. Physiological and Biochemical Zoology. 2004; 77: 998–1018. https://doi.org/10.1086/423743 PMID: 15674772

19. Ezcurra JM, Lowe CG, Mollet HF, Ferry LA, O’Sullivan JE. Captive feeding and growth of young-of-the-year white sharks, Carcharodon carcharias, at the Monterey Bay Aquarium. In: Domeier ML, editor. Global Perspectives on the Biology and Life History of the White Shark. 2012. pp. 17–26.

20. Grady JM, Enquist BJ, Dettweiler-Robinson E, Wright NA, Smith FA. Evidence for mesothermy in dinosaurs. Science. 2014; 344: 1268–1272. https://doi.org/10.1126/science.1253143 PMID: 24926017

21. Wegner NC, Snodgrass OE, Dewar H, Hyde JR. Whole-body endothermy in a mesopelagic fish, the opah, Lampris guttatus. Science. 2015; 348: 786–789. https://doi.org/10.1126/science.aaa8902 PMID: 25977549

22. Weng KC, Block BA. Diel vertical migration of the bigeye thresher shark (Alopias superciliosus), a species possessing orbital retia mirabilia. Fishery Bulletin. 2004; 102: 221–229.

23. Bernal D, Sepulveda CA, Beaupre SJ. Evidence for temperature elevation in the aerobic swimming musculature of the common thresher shark, Alopias vulpinus. Copeia. 2005; 1:146–151.

24. Sepulveda CA, Wegner NC, Bernal D, Graham JB. The red muscle morphology of the thresher sharks (family Alopiidae). Journal of Experimental Biology. 2005; 208: 4255–4261. https://doi.org/10.1242/ jeb.01898 PMID: 16272248

25. Lowe CG, Goldman KJ. Thermal and bioenergetics of elasmobranchs: bridging the gap. In: Tricas TC, Gruber SH, editors. The Behavior and Sensory Biology of Elasmobranch Fishes: An Anthology in Memory of Donald Richard Nelson. 2001. pp. 251–266.

26. Applegate RW, Espinosa-Arrubarrena L. Fossil history of Carcharodon and its possible ancestor, Cretolemna: a study in tooth identification. In: Klimley AP, Ainley DG, editors. Great White Sharks: The Biology of Carcharodon carcharias; 1996. pp. 19–36.
27. Martin AF. Systematics of the Lamnidae and origination time of mitochondrial DNA sequences. In: Klimey AP, Ainley DG, editors. Great White Sharks: The Biology of Carcharodon carcharias. 1996. pp. 49–53.
28. Gottfried MD, Fordyce RE. An associated specimen of Carcharodon angustidens (Chondrichthyes, Lamnidae) from the Late Oligocene of New Zealand, with comments on Carcharodon interrelationships. Journal of Vertebrate Paleontology. 2001; 21: 730–739.
29. Purdy RW, Schneider VP, Applegate SP, McLellan JH, Meyer RL, Slaughter BH. The Neogene Sharks, Rays, and Bony Fishes from Lee Creek Mine, Aurora, North Carolina. In: Ray CE, Bohaska DJ, editors. Geology and Paleontology of the Lee Creek Mine, North Carolina. 2001. pp. 71–202.
30. Kent BW. Speculations on the size and morphology of the extinct lamnoid shark, Parotodus benedeni (Le Ron). The Mosasaur. 1999; 6: 11–15.
31. Capetta H. Chondrichthyes 2. Mesozoic and Cenozoic elasmobranchii 3B. Gustav Fischer Verlag, Stuttgart; 1987.
32. Ward DJ, Bonavia CG. Additions to, and a review of, the Miocene shark and ray fauna of Malta. Cenozoic Geology. 2008; 20: 191–203.
33. Capetta H, Chondrichthyes 2. Mesozoic and Cenozoic elasmobranchii 3B. Gustav Fischer Verlag, Stuttgart; 1987.
34. Ehret DJ, Hubbell G, MacFadden BJ. Exceptional preservation of the white shark Carcharodon (Lamniformes, Lamnidae) from the early Pliocene of Peru. Journal of Vertebrate Paleontology. 2009; 29: 1–13.
35. Ehret DJ, Macfadden BJ, Jones DS, Devries TJ, Foster DA, Salas-Gismondi R. Origin of the white shark Carcharodon (Lamniformes: Lamnidae) based on recalibration of the Upper Neogene Pisco Formation of Peru. Palaeontology. 2012; 55: 1139–1153.
36. Bernard A, Lécuyer C, Vincent P, Amiot R, Bardet N, Buffetaut E, et al. Regulation of body temperature by some Mesozoic marine reptiles. Science. 2010; 328: 1379–1382. https://doi.org/10.1126/science.1187443 PMID: 20538946
37. Kocsis L, Gheerbrant E, Mouflih M, Cappetta H, Yans J, Amaghzaz M. Comprehensive stable isotope investigation of marine biogenic apatite from the late Cretaceous–early Eocene phosphate series of Morocco. Palaeogeography, Palaeoclimatology, Palaeoecology. 2014; 394: 74–88.
38. Amiot R, Göhlich UB, Lécuyer C, De Muizon C, Cappetta H, Fourel F et al. Oxygen isotope compositions of phosphate from Middle Miocene–Early Pliocene marine vertebrates of Peru. Palaeoecography, Palaeoclimatology, Palaeoecology. 2008; 264: 85–92.
39. Lécuyer C, Grandjean P, O’Neil JR, Cappetta H, Martineau F. Thermal excursions in the ocean at the Cretaceous—Tertiary boundary (northern Morocco): δ18O record of phosphatic fish debris. Palaeoecography, Palaeoclimatology, Palaeoecology. 1993; 105: 235–243.
40. Puceát E, Lécuyer C, Sheppard SM, Dromart G, Reboulet S, Grandjean P. Thermal evolution of Cretaceous Tethyan marine waters inferred from oxygen isotope composition of fish tooth enamels. Palaeoceanography. 2003; 18: 11.11.
41. Puceát E, Lécuyer C, Donnadieu Y, Naveau P, Cappetta H, Ramstein G, et al. Fish tooth δ18O revising Late Cretaceous meridional upper ocean water temperature gradients. Geology. 2007; 35: 107–110.
42. Shimada K, Cumbaa SL, van Rooyen D. Caudal fin skeleton of the Late Cretaceous lamniform shark, Cretaluxuria mantelli, from the Niobrara Chalk of Kansas. In: Lucas GS, Sullivan RM, editors. Late Cretaceous Vertebrates from the Western Interior. 2006. pp. 185–192.
43. Shimada K. Skeletal and dental anatomy of lamniform shark, Cretalamna appendiculata, from Upper Cretaceous Niobrara Chalk of Kansas. Journal of Vertebrate Paleontology. 2007; 27: 584–602.
44. Shimada K, Cumbaa SL, van Rooyen D. Caudal fin skeleton of the Late Cretaceous lamniform shark, Cretalamna appendiculata, from Upper Cretaceous Niobrara Chalk of Kansas. Journal of Vertebrate Paleontology. 2007; 27: 584–602.
45. Pimiento C, Ehret DJ, MacFadden BJ, Hubbell G. Ancient nursery area for the extinct giant shark Megalodon from the Miocene of Panama. PLoS One. 2010; 5: e10552. https://doi.org/10.1371/journal.pone.0010552 PMID: 20479893
46. Pimiento C, Balk MA. Body-size trends of the extinct giant shark Carcharocles megalodon: a deep-time perspective on marine apex predators. Paleobiology. 2015; 41: 479–490. https://doi.org/10.1017/pab.2015.16 PMID: 26321775
47. Shimada K, Chandler RE, Lam OLT, Tanaka T, Ward DJ. A new elusive otodontid shark (Lamniformes: Otodontidae) from the lower Miocene, and comments on the taxonomy of otodontid genera, including the “megatoothed”clade. Historical Biology. 2017; 29: 704–714.
48. Sambilay VC. Interrelationships between swimming speed, caudal fin aspect ratio and body length of fishes. Fishbyte. 1990; 8: 16–20.

49. Mollet HF, Cailliet LJ. Using allometry to predict body mass from linear measurements of the white shark. In: Klimek AP, Ainley DG, editors. Great White Sharks: The Biology of Carcharodon carcharias. 1996. pp. 81–89.

50. Kim SH, Shimada K, Rigsby CK. Anatomy and evolution of heterocercal tail in lamniform sharks. The Anatomical Record. 2013; 296: 433–442. https://doi.org/10.1002/ar.22647 PMID: 23381874

51. Carlson JK, Palmer CL, Parsons JR. Oxygen consumption rate and swimming efficiency of the black-nose shark, Carcharhinus acronotus. Copeia. 1999; 1: 34–39.

52. Sepulveda CA, Graham JB, Bernal D. Aerobic metabolic rates of swimming juvenile mako sharks, Isurus oxyrinchus. Marine Biology. 2007; 152: 1087–1094.

53. Korsmeyer KE, Steffensen JF, Herskin J. Energetics of median and paired fin swimming, body and caudal fin swimming, and gait transition in porpoise (Scarus schlegeli) and triggerfish (Rhinecanthus aculeatus). Journal of Experimental Biology. 2002; 205: 1253–1263. PMID: 11948202

54. William F, Beamish H. Swimming metabolism and temperature in juvenile walleye, Stizostedion vitreum vitreum. Environmental Biology of Fishes. 1990; 27: 309–314.

55. Claireaux G, Couturier C, Groison AL. Effect of temperature on maximum swimming speed and cost of transport in juvenile European sea bass (Dicentrarchus labrax). Journal of Experimental Biology. 2006; 209: 3420–3428. https://doi.org/10.1242/jeb.02346 PMID: 16916977

56. Ohlberger J, Staaks G, Höfler K. Effects of temperature, swimming speed and body mass on standard and active metabolic rate in vendace (Coregonus fallax). Journal of Comparative Physiology B. 2007; 177: 905–916.

57. Dowd WW. Metabolic rates and bioenergetics of juvenile sandbar sharks (Carcharhinus plumbeus). The College of William and Mary Press, Virginia; 2003.

58. Koch PL. Isotopic study of the biology of modern and fossil vertebrates. In: Michener R, Lajtha K, editors. Stable Isotopes in Palaeoecology. The College of William and Mary Press, Virginia; 2003. pp. 99–154.

59. Barrick RE, Showers WJ. Thermophysiology of Tyrannosaurus rex: evidence from oxygen isotopes. Science. 1994; 265: 222–224. https://doi.org/10.1126/science.265.5169.222 PMID: 17750663

60. Barrick RE, Showers WJ. Oxygen isotope variability in juvenile dinosaurs (Hypacrosaurus): evidence for thermoregulation. Paleobiology. 1995; 4: 552–560.

61. Fricke HC, Rogers RR. Multiple taxon–multiple locality approach to providing oxygen isotope evidence for warm-blooded theropod dinosaurs. Geology. 2000; 28: 799–802.

62. Harrell TL, Pérez-Huerta A, Suarez CA. Endothermic mosasaurs? Possible thermoregulation of Late Cretaceous mosasaurs (Reptilia, Squamata) indicated by stable oxygen isotopes in fossil bioapatite in comparison with coeval marine fish and pelagic seabirds. Palaeontology. 2016; 59: 351–363.

63. Lécuyer C, Reynard B, Grandjean P. Rare earth element evolution of Phanerozoic seawater recorded in biogenic apatites. Chemical Geology. 2004; 204: 63–102.

64. Vennemann TW, Hegner E, Cliff G, Benz GW. Isotopic composition of recent shark teeth as a proxy for environmental conditions. Geochimica et Cosmochimica Acta. 2001; 65: 1583–1599.

65. Carey FG, Teal JM, Kanwisher JW, Lawson KD, Beckett JS. Warm-bodied fish. American Zoologist. 1971; 11: 137–143.
72. Carey FJ, Kanwisher JW, Brazier O, Gabrielson G, Casey JG, Pratt HL Jr. Temperature and activities of a white shark, *Carcharodon carcharias*. *Copeia*. 1982; 2: 254–260.
73. Bernal D, Smith D, Lopez G, Weitz D, Grimminger T, Dickson K, et al. Comparative studies of high performance swimming in sharks II. Metabolic biochemistry of locomotor and myocardial muscle in endothermic and ectothermic sharks. *Journal of Experimental Biology*. 2003; 206: 2845–2857. PMID: 12847128
74. Bernal D, Donley JM, Shadwick RE, Syme DA. Mammal-like muscles power swimming in a cold-water shark. *Nature*. 2005; 437: 1349–1352. https://doi.org/10.1038/nature04007 PMID: 16251963
75. Sepulveda CA, Dickson KA, Frank LR, Graham JB. Cranial endothermy and a putative brain heater in the most basal tuna species, *Allothunnus fallai*. *Journal of Experimental Biology*. 2007; 206: 2845–2857. PMID: 12847128
76. Tubbesing VA, Block BA. Orbital rete and red muscle vein anatomy indicate a high degree of endothermy in the brain and eye of the salmon shark. *Acta Zoologica*. 2000; 81: 49–56.
77. Radtke RL, Williams DF, Hurley PC. The stable isotopic composition of bluefin tuna (*Thunnus thynnus*) otoliths: evidence for physiological regulation. *Comparative Biochemistry and Physiology Part A: Physiology*. 1987; 87: 797–801.
78. Bainbridge R. The speed of swimming of fish as related to size and to the frequency and amplitude of the tail beat. *Journal of Experimental Biology*. 1958; 35: 109–133.
79. Wardle CS. Limit of fish swimming speed. *Nature*. 1975; 255: 725–727. PMID: 1134569
80. Webb PW. The effect of size on the fast-start performance of rainbow trout (*Salmo gairdneri*), and a consideration of piscivorous predator-prey interactions. *Journal of Experimental Biology*. 1976; 65: 157–177. PMID: 993700
81. Woodward KK, Kostecki PT, Stevens ED. The effect of size and swimming speed on locomotor kinematics of rainbow trout. *Journal of Experimental Biology*. 1984; 109: 77–95.
82. Videler JJ. Fish swimming. Vol. 10. Chapman & Hall, London; 1993.
83. Altringham JD, Johnston IA. Scaling effects on muscle function: power output of isolated fish muscle fibres performing oscillatory work. *Journal of Experimental Biology*. 1990; 151: 453–467.
84. Altringham JD, Young IS. Power output and the frequency of oscillatory work in mammalian diaphragm muscle: the effects of animal size. *Journal of Experimental Biology*. 1991; 157: 381–389. PMID: 2061706
85. Watanabe YY, Goldman KJ, Caselle JE, Chapman DD, Papastamatiou YP. Comparative analyses of animal-tracking data reveal ecological significance of endothermy in fishes. *Proceedings of the National Academy of Sciences*. 2015; 112: 6104–6109.
86. Dewar H, Graham J. Studies of tropical tuna swimming performance in a large water tunnel-Energetics. *Journal of Experimental Biology*. 1994; 192: 13–31. PMID: 9317243
87. Dewar H, Graham J. Studies of tropical tuna swimming performance in a large water tunnel-Kinematics. *Journal of Experimental Biology*. 1994; 192: 45–59. PMID: 9317308
88. Knower T, Shadwick RE, Katz SL, Graham JB, Wardle CS. Red muscle activation patterns in yellowfin (*Thunnus albacares*) and skipjack (*Katsuwonus pelamis*) tunas during steady swimming. *Journal of Experimental Biology*. 1999; 202: 2127–2138. PMID: 10409484
89. Sepulveda CA, Dickson KA. Maximum sustainable speeds and cost of swimming in juvenile kawakawa tuna (*Euthynnus affinis*) and chub mackerel (*Scomber japonicus*). *Journal of Experimental Biology*. 2000; 203: 3089–3101. PMID: 11003820
90. Dewar HJ, Sepulveda CA, Graham JB, Dickson KA. Swimming performance studies on the eastern Pacific bonito *Sarda chiliensis*, a close relative of the tunas (family Scombridae) I. Kinematics. *Journal of Experimental Biology*. 2003; 206: 2749–2758. PMID: 12847120
91. Sepulveda CA, Dickson KA, Graham JB. Swimming performance studies on the eastern Pacific bonito *Sarda chiliensis*, a close relative of the tunas (family Scombridae) I. Energetics. *Journal of Experimental Biology*. 2003; 206: 2739–2748. PMID: 12847119
92. Dickson KA. Tunas as small as 207 mm fork length can elevate muscle temperatures significantly above ambient water temperature. *Journal of Experimental Biology*. 1994; 190: 79–93. PMID: 9317376
93. Dickson KA, Johnson NM, Donley JM, Hoskinson JA, Hansen MW, Tessier J. Ontogenetic changes in characteristics required for endothermy in juvenile black skipjack tuna (*Euthynnus lineatus*). *Journal of Experimental Biology*. 2000; 203: 3077–3087. PMID: 11003819
94. Graham BS, Grubb D, Holland K, Popp BN. A rapid ontogenetic shift in the diet of juvenile yellowfin tuna from Hawaii. *Marine Biology*. 2007; 150: 647–658.
95. Beamish FWH. Swimming capacity. In: Hoar WS, Randall DJ, editors. *Fish Physiology*, Volume VII Locomotion. Vol. 7. 1979. pp.101–187.
96. Brill RW, Dizon AE. Effect of temperature on isotonic twitch of white muscle and predicted maximum swimming speeds of skipjack tuna, *Katsuwonus pelamis*. Environmental Biology of Fishes. 1979; 4: 199–205.
97. Magnuson JJ. Locomotion by scrombrid fishes: hydromechanics, morphology, and behavior. In: Hoar WS, Randall DJ, editors. Fish Physiology. Volume VII Locomotion. Vol. 7. 1979. pp. 240–315.
98. Webb PW. Hydrodynamics: nonscombrid fish. In: Hoar WS, Randall DJ, editors. Fish Physiology. Volume VII Locomotion. Vol. 7. 1979. pp. 190–239.
99. Iosilevskii G, Weihs D. Speed limits on swimming of fishes and cetaceans. Journal of The Royal Society Interface. 2008; 5: 329–338.
100. Wardle CS. Effects of temperature on the maximum swimming speed of fishes. In: Ali MA, editor. Environmental Physiology of Fishes. 1980. pp. 519–531.
101. Jacoby DM, Siriwat P, Freeman R, Carbone C. Correction to ‘Is the scaling of swim speed in sharks driven by metabolism?’ Biology Letters. 2016; 12.
102. Domenici P, Blake R. The kinematics and performance of fish fast-start swimming. Journal of Experimental Biology. 1997; 200: 1165–1178. PMID: 9319004
103. Domenici P. The scaling of locomotor performance in predator–prey encounters: from fish to killer whales. Comparative Biochemistry and Physiology Part A: Molecular & Integrative Physiology. 2001; 131: 169–182.
104. Martin JE, Tacail T, Adnet S, Girard C, Balter V. Calcium isotopes reveal the trophic position of extant and fossil elasmobranchs. Chemical Geology. 2015; 415: 118–125.
105. Diedrich CG, Felker H. Middle Eocene shark coprolites from shallow marine and deltaic coasts of the pre-north sea basin in central Europe. In: Hunt AP, Milana J, Lucas SG, Spielmann JA, editors. Vertebrate Coprolites. Vol. 57. 2012. pp. 301–318.
106. Shimada K. Paleoecological relationships of the Late Cretaceous lamniform shark, *Cretxyrhina mantelli* (Agassiz). Journal of Paleontology. 1997; 71: 926–933.
107. Everhart MJ. Evidence of feeding on mosasaurs by the Late Cretaceous lamniform shark, *Cretxyrhina mantelli*. Journal of Vertebrate Paleontology. 1999; 17: 43A–44A.
108. Everhart MJ. Oceans of Kansas -A Natural History of the Western Interior Sea. Indiana University Press, Indiana; 2005.
109. Everhart MJ. Bite marks on an elasmosaur (Sauropodactyla; Plesiosaursia) paddle from the Niobrara Chalk (Upper Cretaceous) as probable evidence of feeding by the lamniform shark, *Cretxyrhina mantelli*. PalArch, Vertebrate paleontology series. 2005; 2. 14–24.
110. Aguilera OA, de Aguilera DR. Giant-toothed white sharks and wide-toothed mako (Lamnidae) from the Venezuela Neogene: their role in the Caribbean, shallow-water fish assemblage. Caribbean Journal of Science. 2004; 40: 368–382.
111. Shimada K, Everhart MJ. Shark-bitten Xiphactinus audax (Teleostei: Ichthyodectiformes) from the Niobrara Chalk (Upper Cretaceous) of Kansas. The Mosasaur. 2004; 7: 35–39.
112. Shimada K, Hooks GE. Shark-bitten protostegid turtles from the Upper Cretaceous Moureville Chalk, Alabama. Journal of Paleontology. 2004; 78: 205–210.
113. Aguilera OA, García L, Cozzuol MA. Giant-toothed white sharks and cetacean trophic interaction from the Pliocene Caribbean Paraguán Formation. Paläontologische Zeitschrift. 2008; 82: 204–208.
114. Kallal RJ, Godfrey SJ, Ortner DJ. Bone reactions on a pliocene cetacean rib indicate short-term survival of predation event. International Journal of Osteoarchaeology. 2012; 22: 253–260.
115. Collareta A, Lambert O, Landini W, Di Celma C, Malinverno E, Varas-Malca R, et al. Did the giant extinct shark *Carcharocles megalodon* target small prey? Bite marks on marine mammal remains from the late Miocene of Peru. Palaeogeography, Palaeoclimatology, Palaeoecology. 2017; 469: 84–91.
116. Pimiento C, Clements CF. When did *Carcharocles megalodon* become extinct? A new analysis of the fossil record. PLoS One. 2014; 9: e111086. https://doi.org/10.1371/journal.pone.0111086 PMID: 25338197
117. Van Damme R, Van Dooren TJ. Absolute versus per unit body length speed of prey as an estimator of vulnerability to predation. Animal Behaviour. 1999; 57: 347–352. https://doi.org/10.1006/anbe.1998.0980 PMID: 10049474
118. Ponganis PJ, Ponganis EP, Ponganis KV, Kooyman GL, Gentry RL, Trillmich F. Swimming velocities in otariids. Canadian Journal of Zoology. 1990; 68: 2105–2112.
119. Gerstein E. Manatees, Bioacoustics and Boats Hearing tests, environmental measurements and acoustic phenomena may together explain why boats and animals collide. American Scientist. 2002; 90: 154–163.
120. Rohr JJ, Fish FE, Gilpatrick JW. Maximum swim speeds of captive and free-ranging delphinids: Critical analysis of extraordinary performance. Marine Mammal Science. 2002; 18: 1–19.

121. Aoki K, Amano M, Mori K, Kourogi A, Kubodera T, Miyazaki N. Active hunting by deep-diving sperm whales: 3D dive profiles and maneuvers during bursts of speed. Marine Ecology Progress Series. 2012; 444: 289–301.

122. Shimada K. Skeletal anatomy of the Late Cretaceous lamniform shark, *Cretoxyrhina mantelli* from the Niobrara Chalk in Kansas. Journal of Vertebrate Paleontology. 1997; 17: 642–652.

123. Reif WE. Morphogenesis and function of the squamation in sharks. Neues Jahrbuch für Geologie und Paläontologie. Abhandlungen. 1982; 164: 172–183.

124. Reif WE. Squamation and ecology of sharks. Courier Forschungsinstitut Senckenberg, Frankfurt am Main; 1985.

125. Reif WE, Dinkelacker A. Hydrodynamics of the squamation in fast swimming sharks. Neues Jahrbuch für Geologie und Paläontologie, Abhandlungen. 1982; 164: 184–187.

126. Ferrón HG, Pla C, Martínez-Pérez C, Escudero-Mozo MJ, Botella H. Morphometric Discriminant Analysis of isolated chondrichthyan scales for palaeoecological inferences: the Middle Triassic of the Iberian Chain (Spain) as a case of study. Journal of Iberian Geology. 2014; 40: 87–97.

127. Raschi W, Tabit C. Functional aspects of placoid scales: a review and update. Marine and Freshwater Research. 1992; 43: 123–147.

128. Thomson KS, Simanek DE. Body form and locomotion in sharks. American Zoologist. 1977; 17: 343–354.

129. Martin RA, Hammerschlag N. Marine predator–prey contests: ambush and speed versus vigilance and agility. Marine Biology Research. 2012; 8: 90–94.

130. Godfrey SJ, Altman J. A Miocene cetacean vertebra showing a partially healed compression fracture, the result of convulsions or failed predation by the giant white shark, *Carcharodon megalodon*. Jeffersoniana. 2005; 16: 1–12.

131. Jefferson TA, Stacey PJ, Baird RW. A review of killer whale interactions with other marine mammals: predation to co-existence. Mammal review. 1991; 21: 151–180.

132. Compagno LJ. Alternative life-history styles of cartilaginous fishes in time and space. Environmental Biology of Fishes. 1990; 28: 33–75.

133. Pontzer H, Allen V, Hutchinson JR. Biomechanics of running indicates endothermy in bipedal dinosaurs. PLoS One. 2009; 4: e7783. https://doi.org/10.1371/journal.pone.0007783 PMID: 19911059

134. Sundström LF, Gruber SH. Using speed-sensing transmitters to construct a bioenergetics model for subadult lemon sharks, *Negaprion brevirostris* (Poey), in the field. Hydrobiologia. 1998; 371: 241–247.

135. Semmens JM, Payne NL, Huveneers C, Sims DW, Bruce BD. Feeding requirements of white sharks may be higher than originally thought. Scientific reports. 2013; 3: 1471. https://doi.org/10.1038/srep01471 PMID: 23503585

136. Peterson CC, Nagy KA, Diamond J. Sustained metabolic scope. Proceedings of the National Academy of Sciences. 1990; 87: 2324–2328.

137. Willmer P, Stone G, Johnston I. Environmental physiology of animals. Blackwell, Carlton; 2009.

138. Root T. Energy constraints on avian distributions and abundances. Ecology. 1988; 69: 330–339.

139. Hall CA, Stanford JA, Hauer FR. The distribution and abundance of organisms as a consequence of energy balances along multiple environmental gradients. Oikos. 1992; 65: 377–390.

140. Humphries MM, Thomas DW, Speakman JR. Climate-mediated energetic constraints on the distribution of hibernating mammals. Nature. 2002; 418: 313–316. https://doi.org/10.1038/nature00828 PMID: 12124621

141. Cairns DK, Gaston AJ, Huettmann F. Endothermy, ectothermy and the global structure of marine vertebrate communities. Marine Ecology Progress Series. 2008; 356: 239–250.
146. Block BA, Jonsen ID, Jorgensen SJ, Winship AJ, Shaffer SA, Bograd SJ, et al. Tracking apex marine predator movements in a dynamic ocean. Nature. 2011; 475: 86–90. https://doi.org/10.1038/nature10082 PMID: 21697831

147. Buckley LB, Hurlbert AH, Jetz W. Broad-scale ecological implications of ectothermy and endothermy in changing environments. Global Ecology and Biogeography. 2012; 21: 873–885.

148. Huber BT, Hodell DA, Hamilton CP. Middle–Late Cretaceous climate of the southern high latitudes: stable isotopic evidence for minimal equator-to-pole thermal gradients. Geological Society of America Bulletin. 1995; 107: 1164–1191.

149. Siversson M, Lindgren J. Late Cretaceous sharks Cretoxyrhina and Cardabiodon from Montana, USA. Acta Palaeontologica Polonica. 2005; 50: 301–314.

150. Cumbaa SL, Shimada K, Cook TD. Mid-Cenomanian vertebrate faunas of the Western Interior Seaway of North America and their evolutionary, paleobiogeographical, and paleoecological implications. Palaeogeography, Palaeoclimatology, Palaeoecology. 2010; 295: 199–214.

151. Cook TD, Wilson MV, Murray AM. A middle Cenomanian euselachian assemblage from the Dunvegan Formation of northwestern Alberta. Canadian Journal of Earth Sciences. 2008; 45: 1185–1197.

152. Watanabe YY, Lydersen C, Fisk AT, Kovacs KM. The slowest fish: swim speed and tail-beat frequency of Greenland sharks. Journal of Experimental Marine Biology and Ecology. 2012; 426: 5–11.

153. McClain CR, Balk MA, Benfield MC, Branch TA, Chen C, Cosgrove J, et al. Sizing ocean giants: patterns of intraspecific size variation in marine megafauna. PeerJ. 2015; 3: e715. https://doi.org/10.7717/peerj.715 PMID: 25649000

154. Kolodny Y, Raab M. Oxygen isotopes in phosphatic fish remains from Israel; paleothermometry of tropical Cretaceous and Tertiary shelf waters. Palaeogeography, Palaeoclimatology, Palaeoecology. 1988; 64: 59–67.

155. Kolodny Y, Luz B. Oxygen isotopes in phosphates of fossil fish—Devonian to Recent. In: Taylor HP, O’Neil JR, Kaplan RL, editors. Stable Isotope Geochemistry: A Tribute to Samuel Epstein. Vol. 3. 1991. pp. 105–119.

156. Lécuyer C, Grandjean P, Paris F, Robardet M, Robineau D. Deciphering “temperature” and “salinity” from biogenic phosphates: the δ18O of coexisting fishes and mammals of the Middle Miocene sea of western France. Palaeogeography, Palaeoclimatology, Palaeoecology. 1996; 126: 61–74.

157. Dera G, Pucéat E, Pellenard P, Neige P, Delsate D, Joachimski MM, et al. Water mass exchange and variations in seawater temperature in the NW Tethys during the Early Jurassic: evidence from neodymium and oxygen isotopes of fish teeth and belemnites. Earth and Planetary Science Letters. 2009; 286: 198–207.

158. Pelc A, Halas S, Niedźwiedzki R. Oxygen isotope analysis of shark teeth phosphates from Bartonian (Eocene) deposits in Mangyshlak peninsula, Kazakhstan. Mineralogia. 2011; 42: 53–61.

159. Fischer J, Voigt S, Franz M, Schneider JW, Joachimski M, Tichomirowa MM, et al. Palaeoenvironments of the late Triassic Rhaetian Sea: implications from oxygen and strontium isotopes of hybodont shark teeth. Palaeogeography, Palaeoclimatology, Palaeoecology. 2012; 353: 60–72.

160. Fischer J, Schneider JW, Voigt S, Joachimski MM, Tichomirowa M, Tütken T, et al. Oxygen and strontium isotopes from fossil shark teeth: environmental and ecological implications for Late Palaeozoic European basins. Chemical Geology. 2013; 342: 44–62.

161. Balter V, Lécuyer C. Determination of Sr and Ba partition coefficients between apatite and water from 5 C to 60 C: a potential new thermometer for aquatic paleoenvironments. Geochimica et Cosmochimica Acta. 2004; 68: 423–432.

162. Anderson SD, Goldman KJ, Gatten RE Jr. Temperature measurements from salmon sharks, Lamna ditris, in Alaskan waters. Copeia. 2001; 3: 794–796.

163. Goldman KJ, Anderson SD, Latour RJ, Musick JA. Homeothermy in adult salmon sharks, Lamna ditris. Environmental Biology of Fishes. 2004; 71: 403–411.

164. Alexander RL. Evidence of a counter-current heat exchanger in the ray, Mobula tarapacana (Chondrichthyes: Elasmobranchii: Batoidea: Myliobatiformes). Journal of Zoology. 1995; 237: 377–384.

165. Alexander RL. Evidence of brain-warming in the mobulid rays, Mobula tarapacana and Manta birostris (Chondrichthyes: Elasmobranchii: Batoidea: Myliobatiformes). Zoological Journal of the Linnean Society. 1996; 118: 151–164.

166. Welsh JQ, Bellwood DR. Regional endothermy in a coral reef fish? PLoS one. 2012; 7: e33187. https://doi.org/10.1371/journal.pone.0033187 PMID: 22403736

167. Siversson M. Biology, dental morphology and taxonomy of lamniform sharks from the Campanian of the Kristianstad Basin, Sweden. Palaeontology. 1992; 35: 519–554.
168. Zhelezko VI, Kozlov VA. Elasmobranchii and Paleogene biostratigraphy of Transurals and central Asia. Materials on stratigraphy palaeontology of the Urals. 1999; 3: 403–404.

169. Zhelezko VI. The evolution of the tooth system of the shark genus of *Pseudosirus* Glikman, 1957. Materials for the stratigraphy and paleontology of the Urals. 2000; 4: 136–141.

170. Ward DJ. Sharks and rays. In: Young JR, Gale AS, Knight RI, Smith AB, editors. Fossils of the Gault Clay. 2009. pp. 275–299.

171. Siversson M, Lindgren J, Newbrey MG, Cederström P, Cook TD. Cenomanian—Campanian (Late Cretaceous) Mid-Palaeolatitude Sharks of *Cretalamna appendiculata* Type. Acta Palaeontologica Polonica. 2015; 60: 339–384.

172. Siversson M. A new large lamniform shark from the uppermost Gareile Siltstone (Cenomanian, Late Cretaceous) of Western Australia. Transactions of the Royal Society of Edinburgh: Earth Sciences. 1999; 90: 49–66.

173. Maisey JG. What is an “elasmobranch”? The impact of palaeontology in understanding elasmobranch phylogeny and evolution. Journal of Fish Biology. 2012; 80: 918–951. https://doi.org/10.1111/j.1095-8649.2012.03245.x PMID: 22497368

174. Shinoda K. Phylogeny of lamniform sharks (Chondrichthyes: Elasmobranchii) and the contribution of dental characters to lamniform systematics. Paleontological Research. 2005; 9: 55–72.

175. Diedrich CG. Skeleton of the fossil shark *Isurus denticulatus* from the Turonian (Late Cretaceous) of Germany—Ecological Coevolution with Prey of Mackerel Sharks. Paleontology Journal. 2014; 93423.

176. Bowden DL, Vargas-Caro C, Ovenden JR, Bennett MB, Bustamante C. The phylogenomic position of the grey nurse shark *Carcharias taurus* Rafinesque, 1810 (Lamniformes, Odontaspididae) inferred from the mitochondrial genome. Mitochondrial DNA Part A. 2016; 27: 4328–4330.

177. Giorgioni M, Weissett H, Bernasconi SM, Hochuli PA, Keller CE, Coccioni R, et al. Paleoceanographic changes during the Alban–Conomanian in the Tethys and North Atlantic and the onset of the Cretaceous chalk. Global and Planetary Change. 2015; 126: 46–61.

178. Everhart MJ. Rapid evolution, diversification and distribution of mosasaurs (Reptilia; Squamata) prior to the KT Boundary. 11th Annual Symposium in Paleontology and Geology. 2005; 16–27.

179. Motani R. The evolution of marine reptiles. Evolution: Education and Outreach. 2009; 2: 224–235.

180. Tabor CR, Poulsen CJ, Lunt DJ, Rosenbloom NA, Otto-Bliesner BL, Markwick PJ, et al. The cause of Late Cretaceous cooling: A multimodel-proxy comparison. Geology. 2016; 44: 963–966.

181. Myers CE, Lieberman BS. Sharks that pass in the night: using Geographical Information Systems to investigate competition in the Cretaceous Western Interior Seaway. Proceedings of the Royal Society of London B: Biological Sciences. 2010; 278: rspb20101617.

182. Pimiento C, Griffin JN, Clements CF, Silvestro D, Varela S, Uhen MD, et al. The Pliocene marine megafauna extinction and its impact on functional diversity. Nature Ecology & Evolution. 2017; https://doi.org/10.1038/s41559-017-0223-6

183. Amiot R, Lécuyer C, Buffetaut E, Escarguel G, Fluteau F, Martineau F. Oxygen isotopes from biogenic apatites suggest widespread endothermy in Cretaceous dinosaurs. Earth and Planetary Science Letters. 2006; 246: 41–54.

184. Eagle RA, Tütken T, Martin TS, Tripati AK, Fricke HC, Connely M, et al. Dinosaur body temperatures determined from isotopic (13C,18O) ordering in fossil biominerals. Science. 2011; 333: 443–445. https://doi.org/10.1126/science.1206196 PMID: 21700837

185. Fitzgerald EM. A bizarre new toothed mysticete (Cetacea) from Australia and the early evolution of baleen whales. Proceedings of the Royal Society of London B: Biological Sciences. 2006; 273: 2955–2963.