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Calcium stable isotopes place Devonian conodonts as first level consumers

V. Balter^{1*}, J.E. Martin¹, T. Tacail², G. Suan¹, S. Renaud³, C. Girard⁴



Abstract

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Conodont animals are an extinct group of jawless vertebrates whose hard parts, also known as conodont elements, represent the earliest evidence of a mineralised skeleton in the vertebrate lineage. Conodont elements are interpreted as parts of a feeding apparatus, which together with the presence of eyes and microwear patterns, support the controversial hypothesis that conodont animals were macrophagous predators and/or scavengers. Here, we explore the trophic position of five conodont genera (*Palmatolepis, Polygnathus, Ancyrodella, Ancyrognathus* and *Icriodus*) from five contemporary Late Devonian sites distributed worldwide (France, Morocco, Vietnam and Australia) by means of calcium (Ca) stable isotope compositions. The seawater Ca isotope composition was calibrated using contemporary Late Devonian brachiopod isotopic values. By comparison with extant marine trophic chain composed of cartilaginous fish, conodont Ca isotope compositions are indicative of a zooplanktivore - primary piscivore niche, with

no genus-specific trophic pattern. The question of active predation or scavenging cannot be resolved definitively but our results strongly suggest that Late Devonian conodonts were first level consumers.

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Introduction

Until the publication of the discovery of the first specimen of the conodont animal in 1983 with conodont elements in situ forming a feeding apparatus, the nature and function of the conodont elements was one of palaeontology's great mysteries (Briggs et al., 1983). Since then, several other specimens have shown similar schemes for this feeding apparatus, in which the anterior elements form a structure allowing them to trap food that is further processed by the posterior elements (Purnell and Donoghue, 1997). Today, despite the advancement of synchrotron microtomography that allows reconstructing virtual movements of the elements forming the feeding apparatus (Goudemand et al., 2011), the dietary behaviour of conodonts remains an open debate but recently Shirley et al. (2018) suggested a predatory or scavenger mode of life. Conodonts possessed sclerotic eye capsule and extrinsic eye musculature (Gabbott et al., 1995; Purnell, 1995a), consistent with conodonts having pattern vision and an active predatory lifestyle. Lastly, microwear patterns were found on conodont elements, which constituted the first direct evidence that they functioned as teeth (Purnell, 1995b).

In the present work, which is a pilot study, we use for the first time calcium (Ca) stable isotopes to infer the feeding habit

of conodont animals. The first studies showing that trophic levels of animals, including fish, could be inferred from the Ca isotope compositions of their shell or inner skeleton lay back in the 2000's (Skulan et al., 1997; Skulan and DePaolo, 1999; Clementz et al., 2003; DePaolo, 2004). These results were recently confirmed on modern and fossil elasmobranchs (Martin et al., 2015), a subclass of cartilaginous fish, including the sharks, rays and skates, and sawfish. Some authors have already measured the Ca isotope composition of conodonts but with the aim to reconstruct variations of the seawater composition (Hinojosa et al. 2012; Jost et al. 2014; Le Houedec et al. 2017). Here, the Late Devonian period, particularly the Frasnian-Famennian boundary (F/F), was chosen because it is accompanied by important variations in the shape of conodont elements, suggestive of changes in the feeding behaviour of several genera (Balter et al., 2008; Girard and Renaud, 2008). The Material and Method sections are described in the Supplementary Information.

Results

All values presented in this work are expressed as $\delta^{44/42}$ Ca and defined as $\delta^{44/42}$ Ca = ($(^{44}$ Ca/ 42 Ca_{sample}) / (44 Ca/ 42 Ca_{smple}) – 1) * 1000. All measured samples were plotted as $\delta^{43/42}$ Ca against

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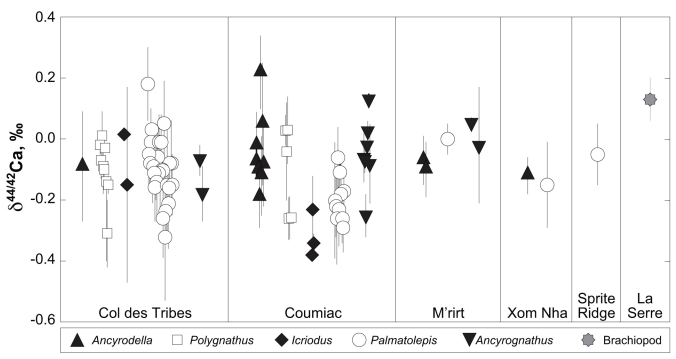


Figure 1 Ca isotope compositions of conodonts ($\delta^{44/42}$ Ca) relative to SRM915a (%) measured in the study.

 $\delta^{44/42}$ Ca and fall on a line with a slope of 0.557 close to the theoretical 0.507 slope predicted by the exponential approximation of mass dependent fractionation (Tacail et al., 2014; Fig. S-2). Quality control assessment is given in Table S-2, and Ca isotope values of conodonts measured in this study (Table S-3; Fig. 1) range from -0.38 ‰ to 0.22 ‰, with an average value of -0.10 ± 0.22 ‰ (± 2 s.d., n = 80). The average $\delta^{44/42}$ Ca value is -0.10 ± 0.20 ‰ (±2 s.d., n = 39) at Col des Tribes, $-0.12 \pm 0.28 \%$ ($\pm 2 \text{ s.d.}$, n = 39) at Coumiac, $-0.03 \pm$ $0.12 \% (\pm 2 \text{ s.d.}, n = 5)$ at M'rirt, $-0.14 \pm 0.04 \% (\pm 2 \text{ s.d.}, n = 2)$ at Xom Nha and -0.05 ‰ at Sprite Ridge (Table S-3). Comparison between conodonts at Coumiac and Col des Tribes, the two most abundant sites, reveals no significant difference of the $\delta^{44/42}$ Ca value (Student's *t*-test, p = 0.373). Comparisons between conodonts grouped by genus reveal no taxonomic difference (Table S-4). The brachiopod yielded a $\delta^{44/42}$ Ca value of 0.13 ‰ (Table S-1), which represents one of the highest values of the dataset.

Discussion

The present paper focuses on the trophic position of conodont animals based on their Ca isotope compositions, but these could have been affected by diagenetic processes. Discussion on the effects of diagenesis is developed in the Supplementary Information. We conclude, in the absence of any evidence of Ca isotope compositions being diagenetically reworked, that the measured $\delta^{44/42}\text{Ca}$ values of conodonts are biogenic.

The present paper focuses on the trophic position of conodont animals based on their Ca isotope composition, but accurate comparisons with modern analogues first necessitate calibrating the Ca isotopic values of the conodont elements relative to that of the contemporaneous seawater. Based on the existing Ca isotopic fractionation factor between modern brachiopods (br) *Terebratalia* and seawater (sw), $\alpha_{\rm br-sw} = 0.99915$ (Gussone *et al.*, 2005), the $\delta^{44/42}$ Ca value of end-Devonian seawater ($\delta^{44/42}$ Ca_{sw}) was estimated at 0.55 ‰, *i.e.* ~0.4 ‰ lower than that of modern oceans (Blättler *et al.* 2012). With an age estimated slightly younger than the Devonian-Carboniferous boundary of 360 Ma, this value falls in the range, but

in the lower limit, of the reconstructed Phanerozoic seawater Ca isotope composition of Farkaš et al. (2007; Fig. S-3). We can now calculate the Ca isotopic offset between seawater and conodonts, which is equal to 0.65 ± 0.25 ‰, (\pm s.d., n = 80). Conodonts are made up of hydroxylapatite (hap), which is more or less fluorinated, but it is the same mineral phase as that of elasmobranch teeth. This allows comparing the average Ca isotopic offset between Devonian seawater and conodonts with that of modern seawater and extant elasmobranch tooth enameloid (Martin et al. 2015), which is annotated $\Delta^{44/42}$ Ca_{sw-hap} (Fig. 2). Using a modern seawater $\delta^{44/42}$ Ca_{sw} value of 0.92 ‰, an offset of ~0.65 ‰ is observed nowadays between seawater and the zooplanktivore and primary piscivores group, which are characterised by average $\Delta^{44/42}$ Ca_{sw-hap} values of 0.56 \pm 0.27 ‰ (\pm 2 s.d., n = 5) and 0.86 \pm 0.08 ‰ (± 2 s.d., n = 6), respectively (Fig. 2). To fully encompass the $\delta^{44/42}\text{Ca}_{sw}$ variability at that time, which is well described by the study of Farkaš et al. (2007), we can also calculate the $\Delta^{44/42}Ca_{sw-hap}$ with the upper limit of the contemporaneous $\delta^{44/42}$ Ca_{sw} value, *i.e.* ~0.67 ‰. Even with this higher value, the calculated $\Delta^{44/42}\text{Ca}_{\text{sw-hap}}$ offset shows that conodonts are still in the the zooplanktivore - primary piscivores group (Fig. 2). The observation that conodonts fall as first level consumers is in accordance with the macrophagous hypothesis (i.e. feeding on relatively large particles of food), but is at odds with the view that conodont animals had a purely predatory lifestyle, which would have implied a $\delta^{44/42}$ Ca value of conodont elements around 1 ‰. Scavenging of fish cannot be ruled out, but must have involved small fish above all, otherwise the $\delta^{44/42}$ Ca values would have been those of predators.

Another argument in favour of a basal trophic position for conodonts, is that modern piscivore elasmobranchs exhibit a much tighter grouping of the $\delta^{44/42}\text{Ca}$ values than modern zooplanktivore elasmobranchs and conodonts (Figs. 1 and 2). The range of $\delta^{44/42}\text{Ca}$ values for a given trophic level of piscivore elasmobranchs never exceeds 0.1 ‰ while it is higher than 0.2 ‰ for modern zooplanktivore elasmobranchs and conodonts. No definitive explanation can be put forward from the state of the results, but a reasonable hypothesis could be that animals at the bottom of the trophic chain are more likely to sample local isotopic heterogeneities. This variability



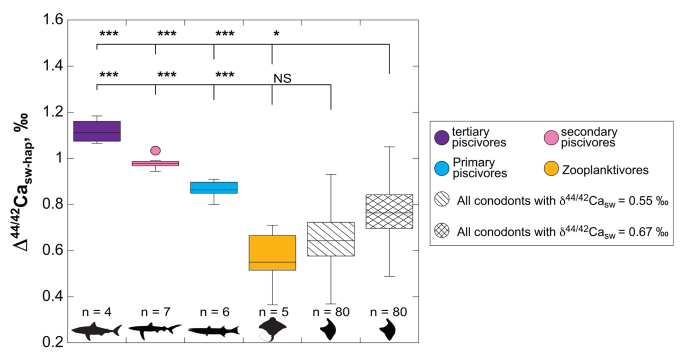


Figure 2 Ca isotopic offset between seawater and hap of modern elasmobranchs compared to that of conodonts ($\Delta^{44/42}$ Ca_{sw-hap}) measured in the study. Modern elasmobranch data are from Martin *et al.* (2015). The $\Delta^{44/42}$ Ca_{sw-hap} offset is calculated with two δ^{44/42}Ca_{sw} values, 0.55 ‰ (diagonal lines) and 0.67 ‰ (diagonal cross hatch) corresponding to the range given by Farkaš *et al.* (2007; Fig. S-3). Boxplots delimit 5, 25, 50, 75 and 95 % percentiles. Depending on the δ^{44/42}Ca_{sw} value, statistics show that conodonts and zooplanktivores have similar δ^{44/42}Ca values. Under the null hypothesis that there is no difference in the distribution of two groups of δ^{44/42}Ca values, the p value of Student's *t*-tests provides the smallest level of significance at which null hypothesis would be rejected (NS, non-significant *p* value; **p* = 0.01–0.05; ***p* = 0.001–0.01; and ****p* < 0.001).

is subsequently contracted at higher trophic level probably thanks to a biopurification process such as in the case of the Sr/Ca and Ba/Ca ratios (Peek and Clementz, 2012).

The present results suggest that no significant difference in trophic level may have existed among conodonts, because genera exhibit indistinguishable $\delta^{44/42}$ Ca values (Table S-2). This overall similarity suggests that competition must have existed between some genera occupying similar trophic levels at the same time, *i.e.* between *Palmatolepis* and *Polygnathus* for instance. It is noteworthy that *Ancyrodella* is the only genus analysed in the study that disappeared at the F/F boundary, questioning the possibility that a distinct ecological trait would have triggered the extinction of this conodont genus.

Using a similar $\Delta^{44/42}\text{Ca}_{\text{sw-hap}}$ for conodont elements and modern elasmobranchs to retrieve the trophic position of conodont animals implies similar vital effects (i.e. isotope fractionation due to biological processes) in both groups. This approach is however probably simplistic. In fish, Ca is taken up along three pathways, (1) directly from the water via the gills, which contain a lot of ion-transporting cells or chloride cells (also known as ionocytes), but also through the intestine from (2) drinking water and (3) food (Flik and Verbost, 1993). No evidence for gills has ever been reported in preserved specimens of conodont animals (Aldridge and Purnell, 1996), which would suggest distinct $\Delta^{44/42}$ Ca_{sw-hap} values between conodont animals and elasmobranchs. Total intestinal absorption of calcium in marine fish represents around 30 % of the total calcium intake (Björnsson and Nilsson, 1985; Sundell and Björnsson, 1988). To our knowledge, relative proportions of drinking water and food in fish have never been determined, but the isotopic results of Martin et al. (2015) in elasmobranchs demonstrate that food must make a sizable proportion, otherwise no trophic effect would have been observed. Indeed, the most likely explanation to account for the depletion of Ca heavy isotopes up trophic chains, being marine or terrestrial, is that preys are wholly ingested along with their skeleton

which is depleted in heavy Ca isotopes. If the three types of Ca uptake described above are characterised by different isotope fractionation intensity, and their relative proportions vary between fish groups, this should in principle result in a different $\Delta^{44/42} \text{Ca}_{\text{sw-hap}}$ fractionation. Analysis of dietary relevant trace elements for marine organisms, such as Sr/Ca and Ba/Ca ratios (Balter and Lécuyer, 2004, 2010; Le Houedec *et al.*, 2013; Peek and Clementz, 2012) would corroborate the present results, but we question whether this would be feasible in light of the difference of vital effects discussed above, and of potential diagenetic effects. Further analysis of Ca isotopes in conodont assemblages will document the diversity of their ecological niches within Palaeozoic oceanic trophic chains.

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Additional Information

Supplementary Information accompanies this letter at http://www.geochemicalperspectivesletters.org/article1912.



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Supplementary Information

The Supplementary Information includes:

- Supplementary Material
- Supplementary Method
- > Supplementary Discussion
- Tables S-1 to S-5
- Figures S-1 to S-3
- Supplementary Information References

Supplementary Material

Conodont elements were collected from Frasnian-Famennian levels of five different localities (Figure S-1): 1) a Famennian succession at the Col des Tribes (France), a newly described outcrop in the Montagne Noire, which exposes a continuous record of the Famennian Stage (Girard et al. 2014); 2) the stratotype area of the F/F boundary, the Coumiac Lower Quarry, located close to the Col des Tribes (Girard and Feist 1997); 3) M'rirt (Central Meseta, Morocco), which shows a similar environmental context to Coumiac, where deposits are a succession of well-oxygenated beds (Lazreq 1999). Conodonts are sampled from the Pa. linguiformis Zone (M9, level just before the Upper Kellwasser event) (Girard et al. 2005); 4) the Xom Nha section (Central Vietnam), which is characterised by a continuous carbonate sequence dated from latest Frasnian to Famennian (here the conodonts are from the Pa. linguiformis Zone XN52, Phuong 1998); 5) Sprite Ridge (Canning Basin, Australia), which is part of an elongated Famennian limestone (Feist and Becker 1997). The sample analysed here is of the Early rhomboidea Zone. In addition, a brachiopod sample was collected from the La Serre section in the Bispathodus ultimus zone corresponding to the CT69 level at Col des Tribes. All correlations are based on conodont biostratigraphy (Flajs and Feist 1988; Girard and Feist 1997; Girard et al. 2010, 2014), and absolute ages are given in Figure S-1 according to Becker et al. (2012). All the samples came from stratigraphic levels around the F/F boundary which is dated of ~ 372 Ma. Calcium isotopes were measured in a total of 80 conodont samples, which are distributed as follows: Palmatolepis, n = 38; Ancyrodella, n = 12; Ancyrognathus, n = 11; Icriodus n = 5; Polygnathus, n = 14. For each studied stratigraphic level and for a given genus, between five and twenty conodont elements (depending on their size and the richness of the stratigraphic layer) were selected and completely dissolved in 1 ml ultrapure concentrated HNO3 overnight. The brachiopod shell was sampled following the protocol of Brazier et al. (2015) and dissolved as for the conodonts.



Supplementary Methods

The details of the purification processes and analytical techniques are given in Tacail *et al.* (2014) but are summarised here. Samples were digested on hotplate using 2 ml concentrated distilled HNO₃. Vials were heated at 120°C during 2 h and regularly degassed. A volume of 2 ml Suprapur 30 % H₂O₂ was added on cooled samples and vials were sealed, regularly degassed at ambient temperature. Finally, vials were sealed and heated on hotplate at 100°C during 2 h and evaporated to dryness. The use of MC-ICPMS requires efficient separation of Ca from samples because of isobaric interferences. Strontium, potassium and to a lesser extent magnesium from samples have to be eliminated to avoid any bias in the measurement of ⁴²Ca⁺ (interfering with ⁸⁴Sr²⁺, ⁴¹K¹H⁺ and ²⁵Mg¹⁶O⁺), ⁴³Ca⁺ (⁸⁶Sr²⁺) and ⁴⁴Ca⁺ (⁸⁸Sr²⁺, ²⁶Mg¹⁸O⁺). The elimination of the remaining matrix components is also required to avoid any bias due to matrix effects. The exact elution procedure is shown in Table S-1.

The first elution aims at discarding K and the majority of matrix elements. Once taken up in 1N HCl, samples were processed on 0.76 cm internal diameter Teflon chromatography columns, filled with 2 ml of Biorad AG50W-X12 cationic resin, 200-400 mesh. The AG50W-X12 columns were reused maximum 5 times, in order to avoid any aging of resin. Strontium was finally eliminated by processing samples on 0.7 ml Eichrom Sr-specific resin, packed in 2 ml Eichrom columns, following a classical Sr elimination protocol (Table S-1). Blank levels never exceed 100 ng, which would represent a contribution of 1/30 of a typical signal. The Sr/Ca ratio, measured as the $^{87}\text{Sr}^{2+/44}\text{Ca}^+$ ratio is always below 10^{-5} . Quality control of the Ca chemical purification is controlled in each session by including a blank and one or two standards which are matrix-matched with the samples (SRM915b "Calcium Carbonate" and SRM1486 "Bone Meal"). Data for the standards are reported in Table S-2. Based on replicates of two standards, the external reproducibility has been estimated at ± 0.066 % (2 SD, n = 130) for SRM1486 and ± 0.048 % (2SD, n=17) for SRM915b in the context of the present study. Since 2014, the overall reproducibility of SRM1486 is ± 0.006 (2 SE, n = 404) with a mean $\delta^{44/42}$ Ca value of ± 0.024 %.

The Ca isotopic compositions were measured using a Neptune plus MC-ICPMS (Thermo Scientific, Bremen, Germany). Standard and sample solutions were prepared to reach a 3 mg.l $^{-1}$ concentration in 0.05N HNO $_3$ medium. Calcium solution was introduced as a dry aerosol with a Cetac Aridus II desolvating system allowing reduction of hydride and oxide formation. The Aridus desolvating system was used with Ar sweep gas flow and an additional N $_2$ gas flow. Aerosols were introduced in a 1200 W plasma with uptake rate of 100 to 150 μ L.min $^{-1}$. The optimised MC-ICPMS instrument operating parameters were : cool gas (15 L.min $^{-1}$), auxiliary gas (0.7-0.8 L.min $^{-1}$) and sample gas (1-1.2 L.min $^{-1}$).

Faraday cups were set to measure $^{42}\text{Ca}^+$ signal on L4 cup, $^{43}\text{Ca}^+$ on L2 and $^{44}\text{Ca}^+$ on central cup. The use of these three isotopes is sufficient for mass-dependent stable isotopes composition measurements in biological materials. The L1 cup was used to monitor the $^{87}\text{Sr}^{2+}$ corresponding to m/z = 43.5. The 42 and 44 ion beams signals were measured with a $10^{11}\,\Omega$ resistance on faraday cup and 43 signal was measured with $10^{12}\,\Omega$ resistance, because of the low abundance of ^{43}Ca (0.135 % of total Ca). Calcium concentrations were adjusted to be within 10 % of the fixed 3 mg.l-1 concentration. Medium mass resolution was sufficient to resolve polyatomic interferences: including $^{40}\text{Ar}^{1}\text{Hz}^{+}$, $^{12}\text{Ci}^{6}\text{Oz}^{+}$ and $^{14}\text{Nz}^{+}$.

Each analysis consisted of 40 measurements of 4.2 s integrations on m/z ratios 42, 43, 44 and 43.5. Even if Sr levels were very low due to the specific separation on the Sr-Spec resin, we corrected the double charge interferences of Sr on Ca (88Sr²+ on 44Ca+, 86Sr²+ on 43Ca+, 84Sr²+ on 42Ca+) using the 43.5 signal corresponding to 87Sr²+. The correction includes the instrumental mass bias on Sr isotopes, which was monitored using the NBS 987 standard. Corrected and uncorrected ratios were calculated for each measurement and averaged after exclusion of values higher than 1SD from average. Instrumental mass bias was corrected by standard-sample-standard bracketing. A Specpure Calcium plasma standard solution (Alfa Aesar), noted ICP-Lyon, was used as reference and bracketing in-house standard. ICP-Lyon required purification because of the presence of Sr traces responsible for significant interferences on the three measured isotopes.

Supplementary Discussion

Sedimentological analyses (Girard et~al.~2014) at the Col des Tribes and geochemical analyses on bulk sediment (δ^{13} C and δ^{18} O) at the Coumiac Quarry (Joachimski and Buggisch 1993) do not argue for any substantial post-depositional modifications. Regarding more specifically the Ca isotope composition of fossils, its use to unravel paleobiological features is in its infancy and data are still lacking to highlight potential methods to detect diagenesis of the original isotopic composition. Calcium is the major element of endogenous mineralised tissues, bone, dentine and enamel, which are made up of hydroxylapatite, and exogenous mineralised tissues, which are made up of calcite or aragonite. In both cases, Ca represents about 40 % weight of the mineral fraction, rendering Ca hardly prone to diagenesis because diagenetic fluids are incommensurably less Ca concentrated (Martin et~al., 2017). For instance, this led recently Pruss et~al. (2018) to show that 600 Myr old Ediacaran shell-forming organisms had still preserved a Ca isotope signature typical of aragonite. Here, we did not measure the concentration of trace elements that specifically incorporate



hap during diagenetic processes, such as rare Earth elements (REE), due to sample size limitation. However, REE concentrations have been previously measured in conodonts at Coumiac along with the strontium/calcium ratio (Sr/Ca) and the oxygen isotope composition of phosphates (δ^{18} O, Le Houedec *et al.*, 2014). From eight common layers where all the geochemical proxies have been measured, we show that, unless the strong correlation between La/Sm and La/Yb, indicative of substitution mechanisms in the context of 'extensive' or 'late' diagenesis (Reynard and Balter, 2014), no proxy is significantly correlated to any other (Table S-5).

Supplementary Tables

Table S-1 Chart of the purification processes.

1. Matrix elimination							
AG50W-X12 resin (200-400 mesh) ~ 2mL							
Step Eluent Vol. (mL)							
Condition	1N HCl	10					
Load	1N HCl	2+1					
Elution (matrix) 1N HCl 55							
Ca elution (Ca,Sr,Fe) 6N HCl 10							
2. Sr elimination							
Sr-Specific resin (Eichrom) ~ 0.7mL							
Step Eluent Vol. (mL)							
Condition	3N HNO3	5					
Load	3N HNO3	0.5+0.5					
Elution (Ca) 3N HNO3 6							
Remaining on resin: Sr							



 $\textbf{Table S-2} \quad \text{Ca isotope compositions of SRM1486 and SRM915b measured in the study relative to ICP-Lyon.}$

Standard		n	δ ⁴⁴ / ⁴² amu	2 SD	reference
SRM1486	session #				
	1	38	-0.52	0.02	
	2	29	-0.53	0.14	
	3	30	-0.50	0.07	
	4	7	-0.52	0.05	
	5	21	-0.48	0.08	
	6	5	-0.51	0.04	
	literature				
		25	-0.52	0.06	Martin et al. (2015)
		17	-0.48	0.07	Tacail et al. (2014)
		142	-0.51	0.07	Heuser and Eisenhauer (2008)*
		2	-0.50	0.07	Heuser <i>et al.</i> (2011)*
SRM915b	session #				
	1	3	-0.11	0.03	
	2	3	-0.06	0.02	
	3	3	-0.06	0.09	
	4	2	-0.06	0.05	
	5	6	-0.10	0.05	
	literature				
		13	-0.08	0.06	Martin et al. (2015)
		11	-0.06	0.04	Tacail et al. (2014)
		56	-0.08	0.01	Heuser and Eisenhauer (2008)*

^{*} TIMS analysis



Table S-3 Ca isotope compositions measured in the study relative to ICP-Lyon and SRM915a standards. Conversion between ICP-Lyon and SRM915a standards is given by the relationship: $\delta^{44/42}$ Ca_{ICP Lyon} = $\delta^{44/42}$ Ca_{SRM915a} - 0.52 (Martin et al. 2015). Error of the $\delta^{44/42}$ Ca_{SRM915a} value is obtained by adding 0.08 % on the error relative to ICP Lyon.

locality	ID	taxon	shape	n	δ ^{44/42} Ca	2 SD	δ ^{43/42} Ca	2 SD	δ ^{44/42} Ca
					vs. ICP Lyon		vs. ICP Lyon		vs. SRM915a
Col des Tribes	CT22An	Ancyrodella		2	-0.61	0.18	-0.32	0.08	-0.09
	CT22Ag	Ancyrognathus		2	-0.59	0.05	-0.28	0.01	-0.07
	CT23Ag	Ancyrognathus		3	-0.70	0.09	-0.35	0.01	-0.18
	CT22 (IC1)	Icriodus		2	-0.50	0.01	-0.19	0.03	0.02
	CT22 (IC2)	Icriodus		2	-0.67	0.32	-0.28	0.10	-0.15
	CT59-7	Palmatolepis		2	-0.34	0.12	-0.21	0.00	0.18
	CT37	Palmatolepis		2	-0.57	0.10	-0.27	0.07	-0.05
	CT35Pa	Palmatolepis		2	-0.60	0.08	-0.31	0.14	-0.08
	CT33	Palmatolepis		2	-0.53	0.11	-0.28	0.13	-0.01
	CT30Pa	Palmatolepis		2	-0.49	0.01	-0.29	0.02	0.03
	CT23Pa	Palmatolepis	Broad	6	-0.61	0.12	-0.30	0.12	-0.09
	СТ39Ра	Palmatolepis	Broad	2	-0.63	0.07	-0.34	0.00	-0.11
	CT46Pa	Palmatolepis	Broad	6	-0.68	0.11	-0.35	0.08	-0.16
	CT51Pa1	Palmatolepis	Slender	5	-0.63	0.09	-0.33	0.15	-0.11
	CT51Pa2	Palmatolepis	Broad	5	-0.66	0.12	-0.34	0.06	-0.14
	CT54-2Pa	Palmatolepis	Broad	6	-0.58	0.12	-0.28	0.11	-0.06
	CT62Pa_1	Palmatolepis	Slender	4	-0.53	0.09	-0.26	0.05	-0.01
	CT62Pa_2	Palmatolepis	Broad	6	-0.63	0.12	-0.32	0.14	-0.11
	CT69GPa	Palmatolepis	Broad	8	-0.53	0.09	-0.28	0.08	-0.01
	CT56Pa	Palmatolepis	Broad	3	-0.62	0.01	-0.36	0.07	-0.10
	CT33FRPa	Palmatolepis	Broad	3	-0.78	0.13	-0.39	0.14	-0.26
	CT70-2_Pa	Palmatolepis	Broad	3	-0.47	0.14	-0.25	0.07	0.05
	CT35Pa0	Palmatolepis	Broad	3	-0.84	0.21	-0.48	0.13	-0.32
	CT22Pa	Palmatolepis	Broad	3	-0.76	0.10	-0.39	0.09	-0.24
	CT22 (Pa1)	Palmatolepis	Broad	2	-0.67	0.18	-0.40	0.09	-0.15
	CT22 (Pa2)	Palmatolepis	Broad	2	-0.73	0.15	-0.43	0.01	-0.21
	СТ66Ра	Palmatolepis	Broad	2	-0.68	0.07	-0.36	0.03	-0.16
	CT66 (Pa5)	Palmatolepis	Broad	2	-0.60	0.02	-0.36	0.00	-0.08
	CT66 (Pa6)	Palmatolepis	Broad	2	-0.60	0.14	-0.34	0.04	-0.08
	CT66 (Pa7)	Palmatolepis	Broad	2	-0.67	0.10	-0.42	0.10	-0.15
	СТ30Ро	Polygnathus		3	-0.54	0.03	-0.29	0.06	-0.02
	CT46Po	Polygnathus		3	-0.59	0.06	-0.30	0.03	-0.07
	CT69GPo	Polygnathus		4	-0.51	0.08	-0.26	0.10	0.01
	СТ39Ро	Polygnathus		7	-0.61	0.09	-0.33	0.10	-0.09
	CT22Po	Polygnathus		2	-0.62	0.03	-0.29	0.05	-0.10
	CT51Po	Polygnathus		4	-0.55	0.00	-0.28	0.07	-0.03
	CT22 (Po1)	Polygnathus		2	-0.66	0.26	-0.25	0.20	-0.14
	CT22 (Po2)	Polygnathus		2	-0.83	0.11	-0.45	0.01	-0.31
	CT22 (Po3)	Polygnathus		2	-0.67	0.03	-0.31	0.07	-0.15



	1	1						
Coumiac	CUQ31cAn	Ancyrodella	2	-0.54	0.11	-0.29	0.02	-0.02
	CLQ25aAn	Ancyrodella	2	-0.59	0.01	-0.25	0.07	-0.07
	CUQ28cAn	Ancyrodella	2	-0.62	0.01	-0.32	0.02	-0.10
	CLQ25bAn	Ancyrodella	3	-0.71	0.10	-0.37	0.06	-0.19
	CLQ23An	Ancyrodella	3	-0.30	0.12	-0.16	0.15	0.22
	CLQ24An	Ancyrodella	3	-0.64	0.13	-0.32	0.02	-0.12
	CLQ25aAn	Ancyrodella	2	-0.47	0.20	-0.32	0.12	0.05
	CLQ25bAn	Ancyrodella	3	-0.60	0.14	-0.31	0.19	-0.08
	CLQ25aAg	Ancyrognathus	3	-0.58	0.08	-0.28	0.03	-0.06
	CLQ23bAg	Ancyrognathus	2	-0.59	0.04	-0.30	0.02	-0.07
	CLQ25bAg	Ancyrognathus	2	-0.77	0.07	-0.40	0.11	-0.25
	CUQ31cAg	Ancyrognathus	1	-0.54		-0.29		-0.02
	CUQ28cAg	Ancyrognathus	2	-0.49	0.03	-0.24	0.03	0.03
	CLQ23Ag	Ancyrognathus	3	-0.39	0.02	-0.14	0.14	0.13
	CLQ24Ag	Ancyrognathus	4	-0.60	0.13	-0.33	0.12	-0.08
	CLQ28Ic	Icriodus	2	-0.90	0.01	-0.50	0.03	-0.38
	CLQ29aIc	Icriodus	2	-0.75	0.11	-0.39	0.01	-0.23
	CLQ33Ic	Icriodus	2	-0.86	0.03	-0.48	0.06	-0.34
	CLQ23Pa	Palmatolepis	3	-0.72	0.19	-0.38	0.14	-0.20
	CLQ24Pa	Palmatolepis	3	-0.74	0.10	-0.39	0.10	-0.22
	CLQ25aPa	Palmatolepis	3	-0.78	0.15	-0.41	0.11	-0.26
	CLQ25bPa	Palmatolepis	6	-0.58	0.10	-0.31	0.10	-0.06
	CLQ26aPa	Palmatolepis	3	-0.75	0.12	-0.38	0.08	-0.23
	CLQ26bPa	Palmatolepis	2	-0.63	0.04	-0.33	0.10	-0.11
	CLQ27Pa	Palmatolepis	3	-0.70	0.12	-0.36	0.13	-0.18
	CLQ28Pa	Palmatolepis	3	-0.78	0.08	-0.39	0.11	-0.26
	CLQ29aPa	Palmatolepis	3	-0.81	0.08	-0.43	0.01	-0.29
	CLQ33Pa	Palmatolepis	2	-0.69	0.06	-0.40	0.07	-0.17
	CLQ25aPo	Polygnathus	3	-0.49	0.05	-0.25	0.02	0.03
	CLQ25bPo	Polygnathus	3	-0.56	0.16	-0.29	0.07	-0.04
	CLQ29aPo	Polygnathus	3	-0.49	0.11	-0.22	0.08	0.03
	CLQ26bPo	Polygnathus	3	-0.78	0.07	-0.43	0.09	-0.26
	CLQ28Po	Polygnathus	2	-0.78	0.07	-0.39	0.08	-0.26
M'rirt	M9-1An	Ancyrodella	2	-0.62	0.09	-0.32	0.01	-0.10
	M9-2An	Ancyrodella	2	-0.59	0.08	-0.29	0.04	-0.07
	M9-1Ag	Ancyrognathus	2	-0.47	0.02	-0.21	0.02	0.05
	M9-2Ag	Ancyrognathus	2	-0.54	0.19	-0.29	0.06	-0.02
	М9Ра	Palmatolepis	2	-0.52	0.05	-0.25	0.05	0.00
Xom Nha	XNAn	Ancyrodella	3	-0.64	0.06	-0.33	0.03	-0.12
	XNPa	Palmatolepis	2	-0.67	0.14	-0.34	0.09	-0.15
Sprite Ridge	SR1	Palmatolepis	2	-0.57	0.10	-0.33	0.08	-0.05
La Serre	LSbrach	brachiopod	3	-0.39	0.07	-0.20	0.03	0.13



Table S-4 Statistical results (p-value) of Student's t-tests between conodont genera. Under the null hypothesis that there is no difference in the distribution of two groups of $\delta^{44/42}$ Ca values, the p-value provides the smallest level of significance at which null hypothesis would be rejected. (NS, non significant p value; *p = 0.01–0.05; **p = 0.001–0.01; and ***p < 0.001).

	Ancyrodella	Ancyrognathus	Icriodus	Palmatolepis	Polygnathus
Ancyrodella		NS 0.849	NS 0.092	NS 0.092	NS 0.327
Ancyrognathus			NS 0.080	NS 0.068	NS 0.253
Icriodus				NS 0.250	NS 0.190
Palmatolepis					NS 0.586
Polygnathus					

Table S-5 Statistical results (correlation coefficient and associated p-value) of Student's t-tests between geochemical proxies used in the study (NS, non significant p value; *p = 0.01–0.05; **p = 0.001–0.01; and ***p < 0.001).

	δ ^{44/42} Ca	δ180	Sr/Ca	La/Yb	La/Sm
δ ^{44/42} Ca		-0.676, NS 0.066	0.730, NS 0.062	0.518, NS 0.233	0.483, NS 0.273
δ ¹⁸ O			-0.578, NS 0.174	0.048, NS 0.918	0.139, NS 0.767
Sr/Ca				0.139, NS 0.767	0.152, NS 0.745
La/Yb					0.958, *** <10-4
La/Sm					



Supplementary Figures

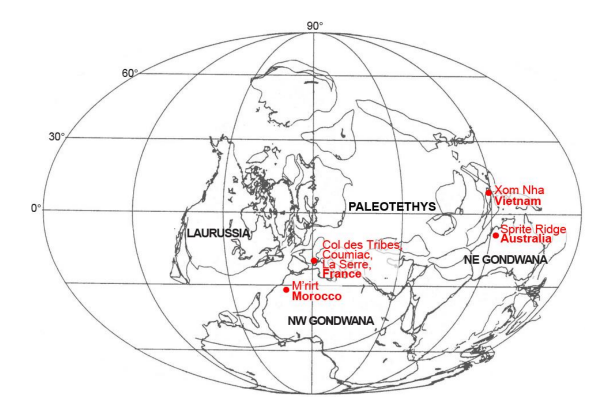


Figure S-1 Localisation of the different sites discussed in the text. Absolute ages are for Col des Tribes, -374 to -360 My; Coumiac, -374 to -371 My; Mrirt and Xom Nha: around -373 My; Sprite Ridge, around 367.5 My. La Serre, -360 My. Ages are from Becker et al. (2012). Conodonts are distributed as following: Col des Tribes, n = 39 (*Palmatolepis*, n=25; *Ancyrodella*, n=1; *Ancyrognathus*, n=2; *Icriodus* n=2; *Polygnathus*, n=9); Coumiac, n = 32 (*Palmatolepis*, n=10; *Ancyrodella*, n=8; *Ancyrognathus*, n=7; *Icriodus* n=2; *Palmatolepis*, n=1); Xhom Nha, n = 2 (*Ancyrodella*, n=1; *Palmatolepis*, n=1); Sprite Ridge, n = 1 (*Palmatolepis*).



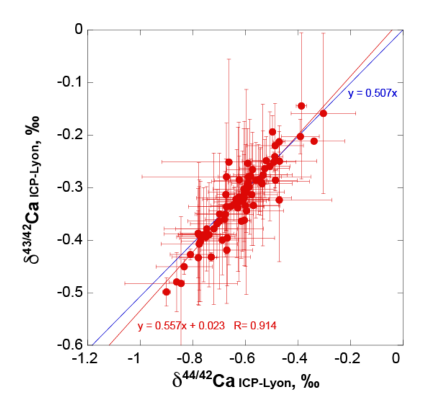


Figure S-2 Three isotopes plot: $\delta^{43/42}$ Ca values as a function of the $\delta^{44/42}$ Ca values (‰, reference standard ICP Ca-Lyon) for all samples and standards analysed in this study. Ca isotope compositions fall on a line with a y-axis intercept of 0.023 indistinguishable from theoretical 0 ‰ intercept. The slope value of this line is 0.557 (in red) very similar from the 0.507 slope (in blue) predicted by the exponential mass-dependent fractionation law.



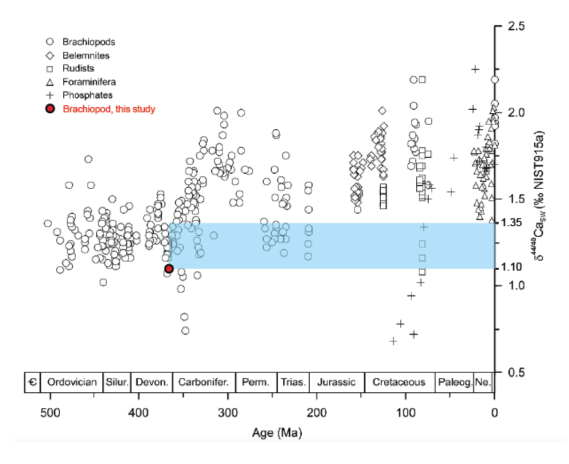


Figure S-3 Value of the $\delta^{44/40}$ Ca of seawater reconstructed from the brachiopod analysed in this study (in red) in the context of the Phanerozoic variations published by Farkaš *et al.* (2007).



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