

Unique physiology of Lepisosteidae imparts a novel phosphate-water fractionation in their scale biogenic apatite

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Abstract

Determining paleotemperatures in terrestrial environments are much more challenging than those in the ocean because of stratigraphic inconsistencies, strong spatial and temporal variations in temperature, and a paucity of well-tested methods. Here we utilize the ganoine scales ofgars from the family Lepisosteidae to calibrate a new terrestrial paleothermometer. Gars are widespread both in the modern and in the past, as they are a freshwater fish lineage that extends back into the Cretaceous (100 Ma) and have remained relatively unchanged during that

time span. Gars constantly record water temperatures, whose yearly average is closely related to mean annual temperature, in their body tissues, including scales. These scales grow continuously throughout life, are >95% hydroxyapatite and thus are highly resistant to diagenetic alteration.

Oxygen isotopes in both biogenic phosphates and carbonates have been used to reconstruct environments on land with varying degrees of success. Phosphate-oxygen isotopes are more resistant to post-mortem alteration as the phosphorus-oxygen bond is stronger than the carbon-oxygen bond. We investigate the application of phosphate oxygen isotopes to gar scales by collecting scales from modern individuals from a north-south transect across the United States, exploiting the latitudinal temperature gradient in mean annual temperatures, measuring $\delta^{18}\text{O}_{\text{phosphate}}$ of those scales, and comparing these values to the average $\delta^{18}\text{O}_{\text{water}}$ and temperature of each locality. We compare our $\delta^{18}\text{O}_{\text{phosphate}}$ calibration to previously published curves. Our work demonstrates that the $\delta^{18}\text{O}_{\text{phosphate}}$ values of gar scales are robust recorders of temperature and $\delta^{18}\text{O}_{\text{water}}$.

1. Introduction

Our understanding of Earth's climatic history is dominated by the interpretation of oceanic sediments, which lacks sensitivity to the climate of continental interiors. Terrestrial paleoclimate knowledge comes from well-preserved plant remains, fossils, speleothems, paleosols, and lacustrine and riverine sediments. Lakes and streams are known to closely match atmospheric air temperature, as they are about 1 °C warmer over the course of a year (e.g. Fricke and Wing, 2004). As ectotherms with indeterminate growth, fish record water temperatures in their body tissues, such as their teeth and scales, over their lifespan of years to decades. These

tissues are an integrated record of mean annual temperature (MAT), probably the most useful measurement of terrestrial temperature. For this study, we evaluate the $\delta^{18}\text{O}$ values of phosphate-rich non-migrating material from gar fish as an emerging methodology in the pursuit of terrestrial climate history.

Vertebrate bone, tooth dentin, and enamel contain calcium phosphate, primarily in the form of hydroxyapatite $\text{Ca}_5(\text{PO}_4)_3(\text{OH})$. Of the biogenic apatites, enamel is the densest, nearly inorganic, and highly crystalline, and thus is known to be the most resistant to isotopic resetting during fossilization (Ayliffe et al., 1994; Koch et al., 1997). Phosphate is more resistant to isotopic resetting than carbonate (e.g., Grimes et al., 2003) and is therefore more ideal for paleoclimate studies.

The isotopic composition of oxygen in biogenic phosphate ($\delta^{18}\text{O}_\text{p}$) and carbonate ($\delta^{18}\text{O}_\text{c}$) are in equilibrium with body fluid during growth (e.g., Iacumin et al., 1996). The body water in turn is directly related to the isotopic composition of ingested water ($\delta^{18}\text{O}_\text{w}$), either from food, drinking water, or water vapor uptake (Luz et al., 1984). The $\delta^{18}\text{O}_\text{c}$ in modern fish apparently does not relate to either the isotopic composition or the temperature of the water in which the fish lived (Iacumin et al., 1996; Kolodny and Luz, 1991). Where $\delta^{18}\text{O}_\text{c}$ falls short in fish and other marine multicellular organisms, the more robust $\delta^{18}\text{O}_\text{p}$ demonstrates a clear temperature-dependent fractionation between biogenic phosphate oxygen and water. Since the oxygen isotopic exchange among inorganic dissolved phosphate and water is depleted in ^{18}O compared to the fractionation in biogenic phosphates, inorganic calibrations cannot be applied to biological systems (Lécuyer et al., 1999). A single phosphate oxygen equation was originally empirically developed for marine ectotherms, demonstrating that the $\delta^{18}\text{O}_\text{p}$ of bivalves and marine fish

67 closely follow the organism's water source (e.g. Kolodny et al., 1983; Longinelli and Nuti,
68 1973a, b).

69 Several following studies better defined the fractionation factor between $\delta^{18}\text{O}_p$, $\delta^{18}\text{O}_w$,
70 and temperature, culminating in the discovery of a universal fractionation equation between
71 dissolved phosphate and water related to a fundamental biosynthetic pathway using the enzyme
72 inorganic pyrophosphatase (PPase) present in all organisms (Chang and Blake, 2015). PPase
73 hydrolyzes inorganic pyrophosphate to inorganic phosphate, imparting a distinct fractionation
74 between water and dissolved inorganic phosphate (DIP).

75 Interestingly, the fractionation factor (α) between body water, bone, or mineral phosphate
76 in mammals does not follow this pattern and seems to vary according to species (e.g. Ayliffe and
77 Chivas, 1990; Longinelli, 1984; Luz et al., 1990). The further fractionation in mammals indicates
78 that taxon-specific fractionation equations may be necessary for certain species, which cannot be
79 explained by the universal fractionation factor alone. The final phosphate oxygen isotopic
80 composition of hard tissues is the result of the mineralization temperature, the initial oxygen
81 isotopic composition of the dissolved phosphate in body fluids, and the biosynthetic pathway
82 using PPase, and further 'vital effects' if the hard tissue is found in mammals.

83 Accounting for the taxon-specific relationships of some organisms, we provide a new
84 calibration using the ganoine scales found in Lepisosteidae, or gars. Gars are more primitive than
85 the Teleosts used in previous empirical calibrations (Kolodny et al., 1983; Longinelli and Nuti,
86 1973a; Puc  at et al., 2010) and their enamel production is more similar to that of mammals than
87 Teleosts due to underlying genetic factors (Braasch et al., 2016). As a freshwater ectotherm, gar
88 bioapatite records a mineralization temperature as well as the $\delta^{18}\text{O}_w$ of the environment. Gars are

model species because of their widespread occurrence in lacustrine settings in the past and present.

Gars are an ancient ray-fined fish lineage which date back to at the Late Cretaceous at least 100 million years ago (Wright et al., 2012). Fossil gars have been found in Europe, India, South America, and north and central Africa in addition to North America. Modern gars retain a large geographic range, extending throughout North America and into Central America and the Caribbean (Grande, 2010). They are unique in that their ganoid scales are highly mineralized with hydroxyapatite and are not underlain with dentin. The enamel directly overlies bone. Ganoid scales are extremely resistant to degradation, and are abundant in the fossil record.

Puc  at et al. (2010) recalibrates the marine fish equation of Kolodny et al. (1983) and Longinelli and Nuti (1973a) using revised analytical methods. Improved methodology to measure $\delta^{18}\text{O}_\text{p}$ has resulted in the cheaper, safer, and more efficient analysis of its isotopic composition (O'Neil et al., 1994). O'Neil et al. (1994) chemically converts biogenic PO_4^{3-} to Ag_3PO_4 and combusts it with graphite to produce CO_2 that is then measured for $\delta^{18}\text{O}$. The equation of Puc  at et al. (2010) is the most widely used calibration using the Ag_3PO_4 method at present for phosphate minerals (e.g., Finnegan et al., 2011; Joachimski et al., 2012).

For this work, we collect modern gar scales from a wide latitudinal range of lakes and rivers in the United States and calibrate their temperature- $\delta^{18}\text{O}_\text{p}$ - $\delta^{18}\text{O}_\text{w}$ relationship with the revised Ag_3PO_4 method. Environmental influences on $\delta^{18}\text{O}_\text{p}$ are evaluated and its usefulness is assessed in reconstructing arguably non-diagenetic, regional $\delta^{18}\text{O}_\text{w}$ and temperature signals.

2. Materials and Methods

2.1. Model Organism, Specimen Acquisition and Site Location

Wild specimens of longnose gar (*Lepisosteus osseus*) and shortnose gar (*Lepisosteus platostomus*) scales were collected over the summer and late fall of 2014 under Yale University Institutional Animal Care and Use Committee Protocol #2012-10681 (Table S1 and Figure 1). Specimens were acquired either personally or with the assistance of the Mississippi Wildlife, Fisheries, & Parks Department, the Illinois Department of Natural Resources, and the Tennessee Wildlife Resources Agency through a combination rod and reel, gill netting, and electrofishing. Further gar specimens were loaned from the Yale Peabody Museum (YPM), Florida Museum of Natural History (UF), Texas Natural Science Center (TNSC), and the Chicago Field Museum (FMNH).

Wild caught fish were used in lieu of those reared from fry in tanks largely because of the extensive amount of time needed to raise these fish into adulthood, as their growth rates only begin to slow at four years of age, or roughly the beginning of sexual maturity (Netsch and Witt, 1962). An extensive, closely monitored tank set-up and feeding regimen is also needed to promote growth (Solomon, 2012). Working with museum collections as well as with state parks and wildlife departments to collect live fish was the most cost and time effective.

Because gars from the *Lepisosteus* and *Atractosteus* genera are able to hybridize (Herrington et al., 2008), any gars from these genera could potentially be used. For the sake of consistency, the paleothermometer is restricted to a single genus. When *Lepisosteus osseus* is not readily available, a sister taxon, *Lepisosteus platostomus*, the shortnose gar, is used.

Lepisosteus osseus, or the longnose gar, is the most widely spread extant gar species in North America. Its range spreads from the tributaries feeding the Mississippi River to the Atlantic coastline. Although the longnose gar has not yet been radio tagged and tracked, the tagging of a closely related sister species, *Atractosteus spatula*, the alligator gar, showed that it does not move more than 12.25 km (Sakaris et al., 2003). The restricted home range of any single individual, tolerance to salinities up to 31 ppt and temperatures from 4° to 31°C (McGrath, 2010), and easy accessibility make it an ideal model species as each specimen can be assigned a specific temperature and $\delta^{18}\text{O}_w$ composition.

2.2. Silver Phosphate Precipitation

For live fish that were caught as well as museum specimens, a 1 to 2 cm² section of scales were cut from the left side of the fish behind the pectoral fin. The right side was left intact for archival purposes. The scales were exposed to dermestid beetles for several weeks to remove overlying tissues.

From the cut section, 0.07 grams of scales were removed and placed in 30% H₂O₂ for a month to remove residual organic matter. Using the methodology of Vennemann et al. (2002) to further remove organics, samples were soaked in 2.5% NaOCl for 24 hours followed by 0.1 M NaOH for 48 hours before dissolution in 2M HNO₃.

Following methods of Liang (2005) and Liang and Blake (2006), modified from Kolodny et al. (1983), dissolved scales were first precipitated as ammonium phosphomolybdate (APM). The APM crystals were dissolved, filtered, and then dissolved phosphate was re-precipitated as magnesium ammonium phosphate (MAP). MAP crystals were dissolved in nitric acid and

cations removed with a resin before trisilver phosphate (Ag_3PO_4) was precipitated. Ag_3PO_4 was vacuum roasted at 550°C to remove residual organics and water.

Oxygen isotope analyses were performed at the YIBS Earth System Center for Stable Isotope Studies (ESCSIS) at Yale University using a Thermo-Chemolysis Elemental Analyzer (TC/EA) coupled to a Delta+ XP continuous flow isotope ratio monitoring mass spectrometer (Thermo-Finnigan, Germany) with a precision of $\pm 0.3\text{‰}$ (1 SD). Ag_3PO_4 crystals were heated to 1450°C in a graphite reactor to release O_2 which reacts with graphite to produce CO. The resultant CO was entrained in a He carrier gas, passed through a gas chromatograph (GC) and introduced into the mass spectrometer. $\delta^{18}\text{O}_{\text{phosphate}}$ values were calibrated against conventional fluorination standards according to Vennemann et al. (2002). All oxygen isotope data are reported as $\delta^{18}\text{O}$ in per mil relative to the VSMOW international reference standard. $\delta^{18}\text{O}_\text{p}$ values are standardized to two internal lab standards, YR1-a and YR3-2, with values of -5.49‰ and 33.64‰ , respectively, for each run.

Water samples from each locality were analyzed on the same Delta+ XP with a GasBench (Thermo-Finnigan, Germany). Yearly $\delta^{18}\text{O}_\text{w}$ values of rivers using the coordinates of each specimen were taken from the Global Network of Isotopes in Rivers (GNIR) database hosted by the International Atomic Energy Agency (IAEA/WMO, 2018). Yearly temperatures were averaged for 15 years before the collection date of the gar specimen (National Centers for Environmental Information, 2022). Temperatures were converted to river temperature using the relationship $T_{\text{atmosphere}} = 1.01 * t_{\text{river}} - 1.13$; river temperatures are approximately 1°C higher than ambient temperature (Fricke and Wing, 2004).

3. Results

Oxygen isotope compositions ($\delta^{18}\text{O}_p$) of gar scales vary from 15.71‰ to 18.76‰ (SE = 0.084, $n = 51$, Table 1). Standard error is calculated from the analytical error for each $\delta^{18}\text{O}_p$ measurement with replicate uncertainty. All errors are at 1σ .

Oxygen isotopes in phosphate are dependent on two variables, temperature (T), and the oxygen isotopic composition of water ($\delta^{18}\text{O}_w$). Of these, the error on $\delta^{18}\text{O}_w$ (SE = 0.320) is larger than that on T (SE = 0.185). Uncertainty for T is \sqrt{n} . For the sake of brevity, $\delta^{18}\text{O}_w - \delta^{18}\text{O}_p$ is shortened to Δ_{p-w} , notation first used in Kolodny et al. (1983) and elucidated in Slater et al. (2001). Following notation used in previous calibrations, T is graphed against Δ_{p-w} and a linear regression model is applied to the data (Figure 2) (Zaarur et al., 2013, supplementary information). Δ_{p-w} contains the largest error ($\delta^{18}\text{O}_w$) so it is placed on the dependent (Y) axis while T, with the smaller error, is placed on the independent (X) axis. All statistical analyses are done with T as the independent variable. The reduced $X^2 = 1.031$ indicates that the best-fit linear regression adequately captures the variance in the dataset.

The Δ_{p-w} residuals have a marginally larger standard deviation ($\sigma = 0.534$) than the standard deviation of the replicates, or multiple fish specimens, from each of ten sites ($n = 39$, $df = 29$, $\sigma = 0.532$). Both of these are smaller than the standard deviation of Δ_{p-w} ($\sigma = 1.133$) and T ($\sigma = 2.663$), indicating that a large source of noise in the dataset is natural variation.

Higher latitude sites undergo a wider temperature fluctuation than lower latitude sites. The lowest latitude site, Coldspring, TX has a yearly standard temperature deviation of 6.81°C with a range between 9.86°C in January and 28.07°C in August. The highest latitude site, Gull Lake, MI ranges from -5.88°C in January to 22.69°C in July with a standard temperature

deviation of 10.34°C. This imparts a wider temperature error at the colder endpoint of the temperature- $\delta^{18}\text{O}_w$ - $\delta^{18}\text{O}_p$ relationship.

The distribution of the data is tighter for samples from closed basins (e.g., Tims Ford Lake, TN) and more dispersed for samples from large drainages (e.g., Rice Lake, off Illinois River, IL.). Large variations in $\delta^{18}\text{O}_w$ coincide with large variations in $\delta^{18}\text{O}_p$. The $\delta^{18}\text{O}_w$ of river water is a combination of precipitation and groundwater values. At coastal sites, such as those from Mississippi and Florida, the precipitation contribution is higher than that found in some inland sites, such as Illinois. The isotopic signature of storm tracks from the Gulf of Mexico imparts a higher $\delta^{18}\text{O}_w$ variation at these sites.

Gars are known to experience faster growth during the late spring before spawning (e.g. Love, 2003; Netsch and Witt, 1962), so comparisons were made to May through August averages in addition to yearly temperatures. $\delta^{18}\text{O}_w$ were the averages from the same months.

4. Discussion

4.1 New Paleothermometer for Gar Scales

This study shows that the oxygen isotopic composition of gar scales is correlated to both the temperature and the oxygen isotopic composition of the source water (Figure 2, Table 1). The inverse-regression method was used to calculate the calibration and confidence intervals. Reliability ratios, λ (Carroll et al., 2006), indicate both variables are relatively insensitive to error ($\lambda_{\Delta P-W} = 0.935$ and $\lambda_T = 0.995$). In contrast to other T- $\delta^{18}\text{O}_w$ - $\delta^{18}\text{O}_p$ calibrations, temperature is chosen as the independent variable on the horizontal axis as it has marginally less error than $\Delta P-W$. Standard error (SE) is corrected for small sample size using Student's t distribution.

There is a marked difference between the recent marine bioapatite thermometry calibration, $T(^{\circ}\text{C}) = 124.6 - 4.52(\delta^{18}\text{O}_p - \delta^{18}\text{O}_w)$ (Puc  at et al., 2010) and the two equations derived here (Figure 3). Reduced $\chi^2 = 1.031$ indicates the standard errors are within what is expected from this calibration. $\text{SE}(\Delta_{p-w})$ is reduced by sampling multiple fish specimen across multiple sites. 39 of the 51 samples were from 10 sites (n_s) with 3 or more different specimens (n_r), which directly affected the calculated confidence interval curves. Adequately minimizing error can be done by sampling at least $n_s = n_r = 4$ (Figure S2).

The calculated calibration for gar scales and mean annual river temperatures (MAWT) is (Figure 3):

$$T(^{\circ}\text{C}) = -2.072(\Delta_{p-w}) + 61.973 \quad (R^2 = 0.778), \quad (1)$$

and that for mean-summer river temperatures (MSWT) is:

$$T(^{\circ}\text{C}) = -1.152(\delta^{18}\text{O}_{\text{phosphate}} - \delta^{18}\text{O}_{\text{water}}) + 50.081 \quad (R^2 = 0.6503) \quad (2)$$

where T is in $^{\circ}\text{C}$ and Δ_{p-w} is in SMOW.

These equations differ in both intercept and slope as well as degree of linear fit, which we attribute to indeterminate gar scale growth recording a MAWT signal (Figure 3). Both calibrations (1, 2) have a shallower slope when using gar scales as compared to the Puc  at et al. (2010) calibration using fish teeth. All teleosts and marine invertebrates were previously assumed to follow the same temperature- $\delta^{18}\text{O}_w$ - $\delta^{18}\text{O}_p$ equation (Kolodny et al., 1983; Longinelli

and Nuti, 1973a, b; Puc  at et al., 2010). This may not be the case for the freshwater gar fish, and for Holostei as a whole. Applying the Puc  at et al. (2010) whole fish-tooth calibration to gar scales returns artificially warmer MAWT temperatures of 2 to 11  C (Figure 3), corresponding to a 0.5 to 2.5   difference in $\delta^{18}\text{O}_p$.

Summer sampled water oxygen isotope values are in agreement with summer $\delta^{18}\text{O}_w$ recorded by Kendall and Coplen (2001)(Figure S2, Table S2) and support using their annual $\delta^{18}\text{O}_w$ measurements.

4.2. Environmental influences on recorded $\delta^{18}\text{O}_{phosphate}$

The MAWT calibration has a significantly better fit than the MSRT calibration as it accounts for a larger fraction of the MAWT and $\delta^{18}\text{O}_{phosphate}$ variation. Error, at least in MAWT, is reduced through time averaging. For the sites chosen, daily and monthly ambient temperatures fluctuate widely while yearly temperatures are relatively consistent. Many of these sample localities experience more sunny days and thus higher evaporation in the summer months while other sample localities receive their greatest precipitation in the spring and early summer. Like temperature, precipitation can still vary widely on a monthly basis. These changes in evaporation and precipitation affect $\delta^{18}\text{O}_w$ and change the recorded $\delta^{18}\text{O}_p$.

Marine systems, particularly in large open oceanic basins, have less variation in both temperature and oxygen isotopic values than their terrestrial equivalents (e.g., Bowen and Revenaugh, 2003; LeGrande and Schmidt, 2006). $\delta^{18}\text{O}_w$ in the ocean is closely correlated with evaporation and precipitation, as manifested in salinity. From the tropics to mid-latitudes oceanic $\delta^{18}\text{O}_w$ ranges from ~ -6 to $\sim +2$    (LeGrande and Schmidt, 2006). On land from the equator to the

264 Arctic Circle, $\delta^{18}\text{O}_w$ ranges from ~ -20 to ~ 0 ‰ (IAEA/WMO, 2018). Monthly high and low
265 mean temperatures of the same latitudinal range on land varies from ~ -30 °C to $\sim +40$ °C whereas
266 those from the ocean from ~ 0 °C to $\sim +28$ °C (National Centers for Environmental Information,
267 2022).

268 Large oceanic basins are unaffected by most of the processes that lead to the wide range
269 of $\delta^{18}\text{O}_w$ and temperatures experienced by lacustrine systems. Lake size, amount of mixing,
270 residence time, catchment size, and whether the body of water is open or closed are some of the
271 many factors that affect lacustrine $\delta^{18}\text{O}_w$ (Leng and Marshall, 2004). To illustrate, larger lakes
272 are generally better mixed than their smaller counterparts, so daily, monthly, and seasonal
273 fluctuations in precipitation $\delta^{18}\text{O}_w$ is averaged. Closed lakes, particularly those in arid climates,
274 have heavier $\delta^{18}\text{O}_w$ values as they are more affected by evaporation, .

275 The larger variation in freshwater lakes compared to marine basins is alluded to in
276 Kolodny et al. (1983). The upper 200 m in Lake Baikal in Siberia, Russia is highly stratified by
277 temperature, ranging from 2 to 12 °C. Below the thermocline, temperatures are relatively
278 constant at 3.2 to 4 °C. Kolodny et al. (1983) accounts for temperature differences by only
279 sampling fish who live in a narrow depth range. $\delta^{18}\text{O}_w$ is relatively consistent at all depths at -
280 15.9 ± 0.2 ‰ (Seal II and Shanks III, 1998). As the deepest and largest freshwater lake by
281 volume, water turnover time is slow, taking over ~ 300 years. Evaporation in this lake contributes
282 from 13 to 19% of its total water loss and the remainder is through riverine outflow (Seal II and
283 Shanks III, 1998 and references therein). The isotopic composition of riverine and precipitation
284 influx of -15.2 ‰ for $\delta^{18}\text{O}_w$ reinforces the long residence time of the water, as Lake Baikal has

not reached a steady state where the lake $\delta^{18}\text{O}_w$ is heavier than the influx $\delta^{18}\text{O}_w$ as would be expected from large amounts of evaporation.

In comparison, the main water source to Lake Kinnereth (Sea of Galilee) in Israel is the Jordan River, not precipitation. The mean annual $\delta^{18}\text{O}_{\text{precipitation}}$ for Safed, Israel is $-6.22 \pm 0.69\text{‰}$ (IAEA/WMO, 2018) and a stable -7.51‰ $\delta^{18}\text{O}_w$ for the nearby Dan tributary to the Jordan River (Gat and Dansgaard, 1972). Lake Kinnereth itself ranges from -2.5 to $+0.5\text{‰}$ $\delta^{18}\text{O}_w$ (Gat, 1984), indicative of heavy evaporation. Kolodny et al. (1983) measure $\delta^{18}\text{O}_p$ of three fish species from this lake, but use a $\delta^{18}\text{O}_w$ of -2.1‰ which is on the lighter side. Zaarur et al. (2016) measure $+0.5\text{‰}$ for the southern shoreline of the lake during the summer months. By adjusting the Lake Kinnereth samples of Kolodny et al. (1983) to the average $\delta^{18}\text{O}_w$ composition of -1‰ and accounting for the error on $\delta^{18}\text{O}_w$, these fish plot in between the MAWT calibration for gar scales and the Puc  at et al. (2010) calibration for marine fish (Figure 3).

The same error that occurs with high evaporation is seen in the seasonally heavy $\delta^{18}\text{O}_w$ measured from localities in Texas, USA (Figure S2). One gar (TNSC 15583) from Hempstead, TX was culled from the final dataset because of the large seasonal variation in $\delta^{18}\text{O}_w$ and the large error on $\delta^{18}\text{O}_p$ ($21.89\text{‰} \pm 1.10$), possibly due to incomplete mineralization as the specimen had not yet reached adulthood. Incomplete mineralization can result in a $\delta^{18}\text{O}_p$ signal that reflects the oxygen isotopic composition of phosphate species other than the phosphate from bioapatite, such as dissolved inorganic phosphate.

Large mixed bodies of water are more robust record holders, such as Lake Baikal or Tims Ford Reservoir, TN, USA (this study) than highly evaporative ones such as Lake Kinnereth or many streams and rivers in Texas where evaporation exceeds precipitation. The isotopic

composition of source waters, either from upstream riverine input or precipitation, can also affect the $\delta^{18}\text{O}_w$ and therefore the $\delta^{18}\text{O}_p$ recorded by the fish, either freshwater or marine. For paleoclimate studies, fish scales and teeth should only be sampled from areas where $\delta^{18}\text{O}_w$ can be well constrained or where temperature can be estimated independently.

4.3. Potential seasonal bias in scale growth

Nuances in growth patterns are recorded in scales. Scales begin mineralizing within the first year and are retained throughout life (Thomson and McCune, 1984) so their isotopic signal is time averaged over the lifetime of the fish. Growth artifacts appear as banding in the underlying bone and ridges in the overlying ganoine (Figure 5). Five to six of these bands are formed within the first two years of life, and formation rates decrease as the fish ages to eventually only one or two a year (Thomson and McCune, 1984). The oldest, largest gars often do not lay down even one band a year. Additionally, band count against standard length in gars gives a straight line. This suggests that instead of correlating to incremental growth, with periods of quiescence, the bands better correlate to increasing scale size so as to maintain the constant standard length growth seen throughout the gar's life (Love, 2003; Netsch and Witt, 1962). The data from this study are considered climatic averages experienced by the fish during life since they are from ground whole scales.

Temperature extremes may adversely impact growth in fry (Solomon, 2012) as they are the most vulnerable at this age with high mortality rates (Haase, 1969). Regardless, gars are capable of thriving in a wide temperature range, from Lake Champlain in Vermont for the Longnose Gar to the Yucatán Peninsula in Mexico for the Tropical Gar. Coupled with

uninterrupted growth rates, temperature extremes only result in torpor, not complete cessation of movement and growth in adults. The spotted gar is the most physically active from March until early June, coincident with preparation before spawning (Snedden et al., 1999). Alligator gar also display the same behavior of increased movement during pre-spawn periods (Buckmeier et al., 2013).

Of note is that females grow at faster rates than males, live longer, and are larger, although they reach sexual maturity later in life (Netsch and Witt, 1962). We did not account for the sex of our gar specimens given that it is difficult to distinguish from external features and sex was assumed to not affect isotopic composition. Gars are in the best condition preceding spawning events with their highest gonadosomatic indices (GSI) (Johnson and Noltie, 1997; Love, 2003). Both male and, even more so, female gar lose a significant amount of body mass after spawning (Johnson and Noltie, 1997), which reduces growth rate. Spawning only occurs after a temperature threshold is reached; this timing may change from year to year but is always in the late spring, from late April to June (Echelle and Riggs, 1972; Haase, 1969; Johnson and Noltie, 1997; Netsch and Witt, 1962).

Growth studies in gars, like most fish, have centered on using either branchiostegal rays (Haase, 1969; Johnson and Noltie, 1997; Klaassen and Morgan, 1974; Netsch and Witt, 1962) or otoliths to better infer age (Ferrara, 2001; Smylie et al., 2016). Distinct banding is present in both structures, and each region between annuli are considered to be equivalent to a year of growth. Generally in the otoliths of fish, opaque and translucent bands alternate and are seasonally related, with opaque bands indicating faster growth and translucent zones forming during periods of slower growth, although conflicting conclusions about timing indicate that otolith formation is

also species and latitude (e.g. tropical, temperate, or subpolar) dependent (Beckman and Wilson, 1995). Longnose gars emplace opaque edges anywhere from February to May (Smylie et al., 2016) and alligator gars in May (Buckmeier et al., 2012) coincident with increase in mass and activity levels in preparation for spawning.

All of the locations in this study appear to have spring temperatures, more specifically the dates associated with the spawning of gars and their highest activity, that are within error to MAT and therefore MAWT. For example, in Benton, IL, the NOAA reference location for the Rend Lake Dam samples recorded MAT from 2000 until 2013 of 13.77 ± 0.74 °C. During the same yearly interval, the average spring temperature, from March, April, and May, was 14.05 ± 1.47 °C. For Grenada, MS, from 2000 to 2013, the MAT was 16.82 ± 0.62 °C and the March, April, and May average from the same time interval was 16.91 ± 1.25 °C.

If there is a growth bias in scales towards the spring as seen in otoliths and branchiostegal rays, it can be assumed that it is equivalent to MAWT. However, the enamel of scales only superficially corresponds to age and it loses its efficacy as an age estimator with time (Thomson and McCune, 1984). In older gars, because age estimates from scales are underestimates (Buckmeier et al., 2012), it is unlikely that gar scale formation can be completely explained by spring seasonal formation as displayed in otoliths and branchiostegal rays. The better correlation between $\delta^{18}\text{O}_p$ and MAWT instead of MSWT confirms this.

4.4. Genetic basis for unique calibration

Two separate fractionations occur between water and dissolved inorganic phosphate (DIP) and between DIP and biological tissues, including DNA, mineral bioapatite, and

biomolecules and soft tissues (Blake et al., 2005; Blake et al., 2016; Chang and Blake, 2015). The distinct fractionation between DIP and body water is caused by the intracellular enzyme reaction involving inorganic pyrophosphatase (PPase). The amino acid sequence of PPase of prokaryotes (bacteria and archaea) and eukaryotes (fungi, plants, and animals) differs substantially but the active site in the resultant enzyme is remarkably conserved across domains (e.g. Cooperman et al., 1992). The fractionation between DIP and water using PPase hence is likely ubiquitous (Chang and Blake, 2015).

The slopes of the DIP–water oxygen isotope fractionation using the marine teleost data of Puc  at et al. (2010) and yeast data of Chang and Blake (2015) are identical within error. This reaction is the primary process that links the $\delta^{18}\text{O}$ composition of dissolved phosphate within cells and body water to environmental factors such as temperature and river isotopic composition (Blake et al., 2016; Chang and Blake, 2015).

Even though there is no apparent further fractionation in tooth formation in teleosts from DIP to bioapatite, as the $\delta^{18}\text{O}_\text{p}$ values are nearly identical to those predicted from the $\delta^{18}\text{O}_\text{w}$ – $\delta^{18}\text{O}_\text{p}$ fractionation alone, DIP in the body water of an organism and cytoplasm of microbial cells may be further fractionated during the creation of phosphatic biomass. There is evidence that this second fractionation may be species-dependent, due to a combination of natural history, habitat such as humidity and aridity for terrestrial species, and genetic factors. Terrestrial, species-specific phosphate-oxygen isotope calibrations have been calculated for numerous endotherms: rabbits (Huertas et al., 1995), mice, cattle, sheep (D'Angela and Longinelli, 1990), kangaroos (Ayliffe and Chivas, 1990), humans (Daux et al., 2008; Longinelli, 1984) and pigs (Longinelli, 1984). These calibrations all follow the same linear trend with positive slopes. Differences arise

if the species is water dependent or drought tolerant (Kohn and Cerling, 2002), as environmental relative humidity has a strong influence on these calibrations (Ayliffe and Chivas, 1990).

Since humidity should be irrelevant for an aquatic species, feeding patterns could partially explain the difference in slope seen in the gar calibration presented here when compared to previously-observed calibrations using both marine and mammalian species. For example, African ungulates eat a wide range of food with unique $\delta^{18}\text{O}$ and, coupled with varying physiologies (Kohn et al., 1996), have unique $\delta^{18}\text{O}_\text{p}$ - $\delta^{18}\text{O}_\text{w}$ trends. In humans, since vegetables and some meats consist of a large weight percent of water, an offset of 0.6‰ to 0.7‰ per mil can be attributed to diet alone (Daux et al., 2008). Because gars are opportunistic feeders but mainly feed on smaller fish, with only the occasional insect or crustacean (Goodyear, 1967), we assume that diet has a minimal influence on $\delta^{18}\text{O}_\text{p}$ for gar scales as all of these animals experience the same $\delta^{18}\text{O}_\text{w}$.

Oxygen in the phosphate of mammals comes from air in addition to food and drinking water, and the relative portions of these sources ultimately affect recorded $\delta^{18}\text{O}_\text{p}$. If there is complete exchange of all four oxygens in PO_4^{3-} with water via PPase catalysis, the slope of the $\delta^{18}\text{O}_\text{p}$ - $\delta^{18}\text{O}_\text{w}$ trend will be 1 (Blake et al., 2005). A slope less than 1 indicates incomplete oxygen transfer from body water to PO_4^{3-} , and a slope greater than 1 can indicate influence from metabolic water or a large dietary or inhaled air effect, either moisture via humidity or the isotopic signature of O_2 itself. The slopes of $\delta^{18}\text{O}_\text{p}$ - $\delta^{18}\text{O}_\text{w}$ for mammals range from 0.49 to 0.57 for rats (Luz et al., 1984; Navarro et al., 2004), 1.01 for cattle (D'Angela and Longinelli, 1990), 1.34 for foxes (Iacumin and Longinelli, 2002), and 1.54 for humans (Daux et al., 2008; Levinson

et al., 1987; Longinelli, 1984; Luz et al., 1984). In gars, when temperature is held constant, the slope is 1 (Figure S2), confirming that $\delta^{18}\text{O}_w$ is the main source of oxygen.

Most notably, there are differences in biomineralization at the genetic level between the taxonomic group that includes gar fish and that of teleosts. Primitive gars are genetically more similar to mammals as they produce true enamel while more derived teleosts produce enameloid (Braasch et al., 2016). Gars, like mammals, have distinct proteins and enzymes that may cause differences in kinetic fractionation when oxygen in DIP is incorporated into the mineral lattice of biogenic apatite.

Mammalian enamel is produced by several proteins: enamelines, amelogenins, ameloblastins, tuftelins and proteinases (Fincham et al., 1999). Teleostei only employs enamelines, producing enameloid. Both enamel and enameloid are highly mineralized ectodermal tissues but they differ in that enamel consists of monolayers with incremental lines and structured prisms while enameloid consists of crystal bundles and is aprismatic (Fincham et al., 1999; Richter and Smith, 1995). More importantly, in enameloid development, odontoblasts create a collagen-rich organic matrix which is dissolved by dental epithelial cells. These dental epithelial cells promote final crystal growth through inorganic ion supply (Sasagawa et al., 2009).

Enamel has a distinct precursor mineral not found in enameloids, octacalcium phosphate ($\text{Ca}_8\text{H}_2(\text{PO}_4)_6 \cdot 5\text{H}_2\text{O}$), which can hydrolyze to hydroxyapatite ($\text{Ca}_5(\text{PO}_4)_3\text{OH}$), and serves as a template for hydroxyapatite precipitation. Furthermore, amelogenin is important in maintaining the crystal spacing of newly precipitated hydroxyapatite. Enameloid crystals are only shaped by

dentin collagen and dentin phosphoprotein. Enamel crystals utilize these in combination with amelogenin (Simmer and Fincham, 1996) which is absent in enameloid production.

The teeth and scales of both *Lepisosteus sp.* and *Polypterus sp.* produce enamel and enameloid (Sasagawa et al., 2009). During scale formation, the preganoine matrix is collagen-free (Sire et al, 1994). The final ganoine thus lacks the collagen fibers of enameloid and is preceded by dentin deposition, making it a true enamel (Sire et al., 1987). Immunohistochemical studies have determined that the amelogenin protein that forms ganoine and teeth in gars is homologous with the mammalian amelogenin protein because it possesses domains that are shared with those found in mammals (Ishiyama et al., 1999; Sasagawa et al., 2014). Preganoine was found to contain the amelogenin protein, the presumable agent for the oxygen isotope fractionation between body water and the phosphate found in the bioapatite of higher vertebrates.

The enamel link between gars and mammals was further supported with the sequencing of the *L. oculatus* genome (Braasch et al., 2016). Teleosts, mammals, and gars all have different genetic modifications of secretory calcium-binding phosphoproteins (Scpp). Interestingly, gars have orthologs of SCPP genes found only in teleosts and different SCPP genes found only in lobe-finned vertebrates so essentially gars genetically straddle the two groups. Specifically, the gar *ambn* and *enam* genes have a genetic homolog in lobe-finned vertebrates; similar but not identical sequences are identified in Teleosts. None of the SCPP genes that teleosts use to mineralize enameloid directly corresponds to tetrapod SCPPs (Kawasaki et al., 2005). The genetic instructions that gars use to create ganoine are more similar to those that mammals and reptiles use to produce enamel than the ones that teleosts use for enameloid.

Teleosts have much faster evolutionary rates than do gars and their allies, and it appears as though they lost the true enamel protein genes along their evolutionary path (Braasch et al., 2016), likely after their total genome duplication event. The archaic structure of cap enameloid and collar enamel is retained in *Polypterus* and *Lepisosteus* (Sasagawa et al., 2013) and their biomineralization pathway similarity to the mammalian one implies that their temperature- $\delta^{18}\text{O}_w$ - $\delta^{18}\text{O}_p$ relationship is more similar to that found in mammals than to marine fish and invertebrates. Further study is needed to determine any kinetic isotopic effects associated with the enamelin or amelogenin proteins during enamel formation.

The observation that gar scales show banding is similar to the growth striations present in mammalian teeth, termed the striae of Retzius. These striations are mandated by amelogenesis and ultimately circadian rhythms. Growth intervals can range from days to weeks to years (Boyde, 1964; Bromage, 1991; Dean, 1987). Because the frequency of ridges in scales decrease as gars age, striations are presumably more a growth artifact than discrete intervals of growth such as otoliths or tree rings.

5. Conclusions

Gar scales have a relationship between MAWT, $\delta^{18}\text{O}_p$, and $\delta^{18}\text{O}_w$ that is significantly different from the one for marine fish postulated by Puc  at et al. (2010). Genomics indicates that ganoine formation in gars is more closely matches that of mammals than that of teleosts (Braasch et al., 2016) as both gars and mammals use enamelin and amelogenin proteins. Terrestrial mammals have species-specific $\delta^{18}\text{O}_p$ - $\delta^{18}\text{O}_w$ relationships and gars appear to have a species-specific calibration as well.

Scales do not form in the same manner as otoliths or branchiostegal rays as the frequency of growth bands decreases with age. While not expected, any bias in recorded temperatures in scales would reflect spring temperatures preceding spawning events when gars are the most active. Riverine spring temperatures at all of the localities are within error to MAWT.

Gars that swim in large bodies of water with little evaporation instead of in smaller bodies of water with large precipitation or river influxes record relatively invariable $\delta^{18}\text{O}_p$ because of more constrained $\delta^{18}\text{O}_w$. For paleoclimate studies, the best gar scales are those from environments that fit these criteria. Gar scales have tentatively been used in paleoclimate research (Fricke et al., 1998; Fricke and Pearson, 2008) and with this new fractionation equation, their future in unraveling the past is very promising. By applying the temperature- $\delta^{18}\text{O}_w$ - $\delta^{18}\text{O}_p$ relationship found in modern gar scales to fossil gar scales, terrestrial climate can be elucidated. If the temperature variable is constrained by other means, then the $\delta^{18}\text{O}_w$ value can be calculated, thus providing information about the hydrological cycle.

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Tables and Figures

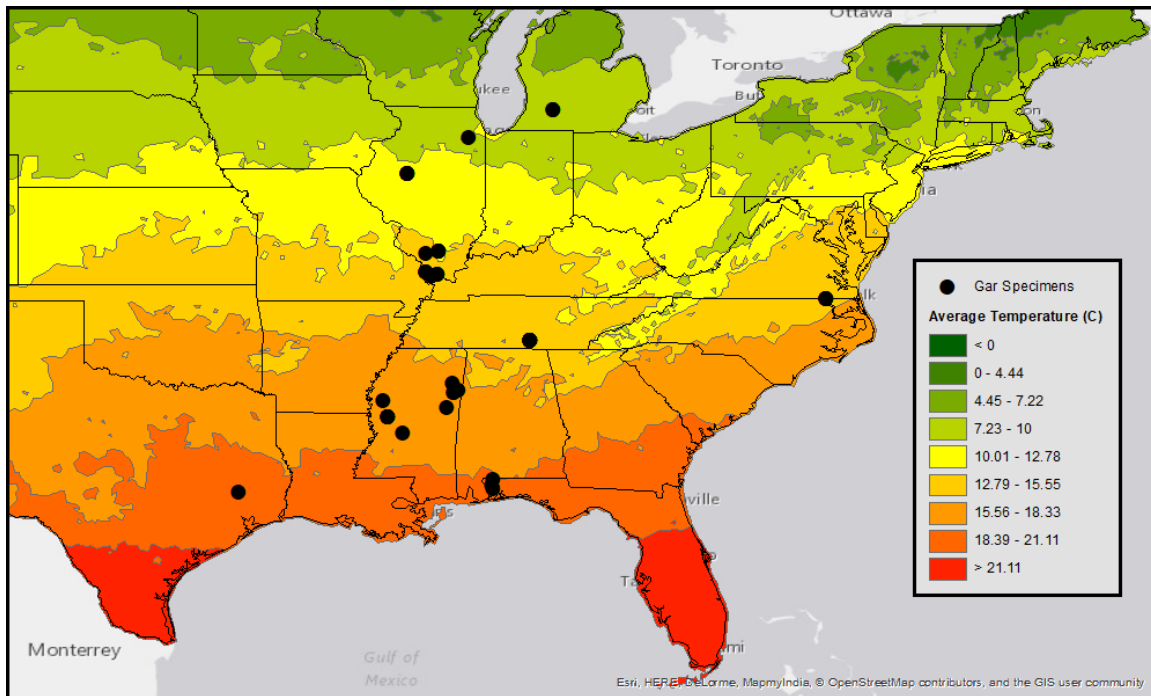


Figure 1. Map of specimen localities superimposed on a map of mean annual air temperature (MAT) (°C) for the continental United States.

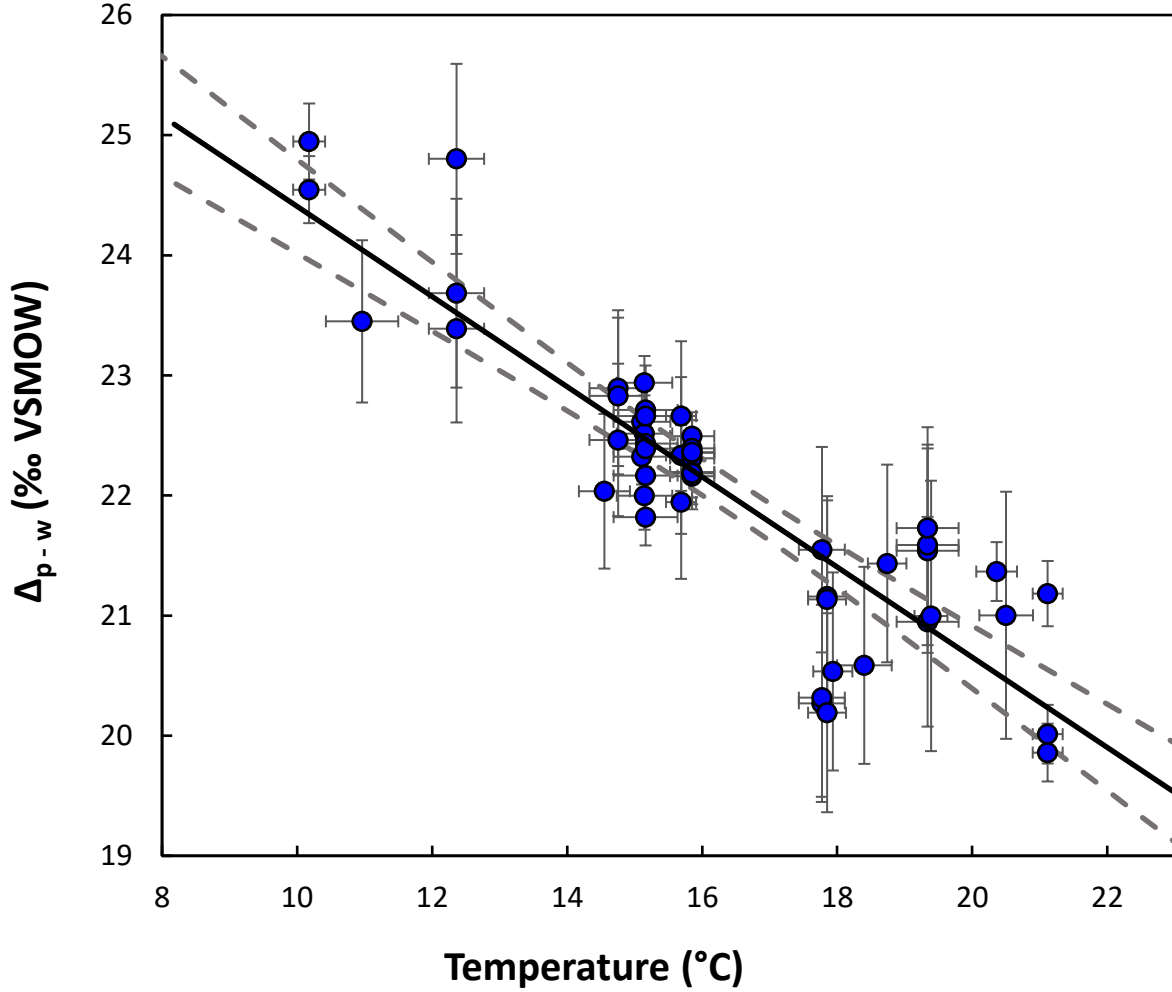


Figure 2. $\Delta_{p-w} - T$ calibration line (solid black line) from oxygen-phosphate isotopes in gar scales ($\delta^{18}O_p$) collected from river water with known oxygen isotopic compositions ($\delta^{18}O_w$) (collectively $\delta^{18}O_p - \delta^{18}O_w$ or Δ_{p-w}), and temperature (T , °C). Error bars are 2σ .

The dashed gray curves are the 95% confidence interval for the calibration line. Temperatures are mean annual water averages over the lifespan of the gar specimens, approximately fourteen to fifteen years. $\delta^{18}O_w$ are three- to four-year averages (Kendall and Coplen, 2001).

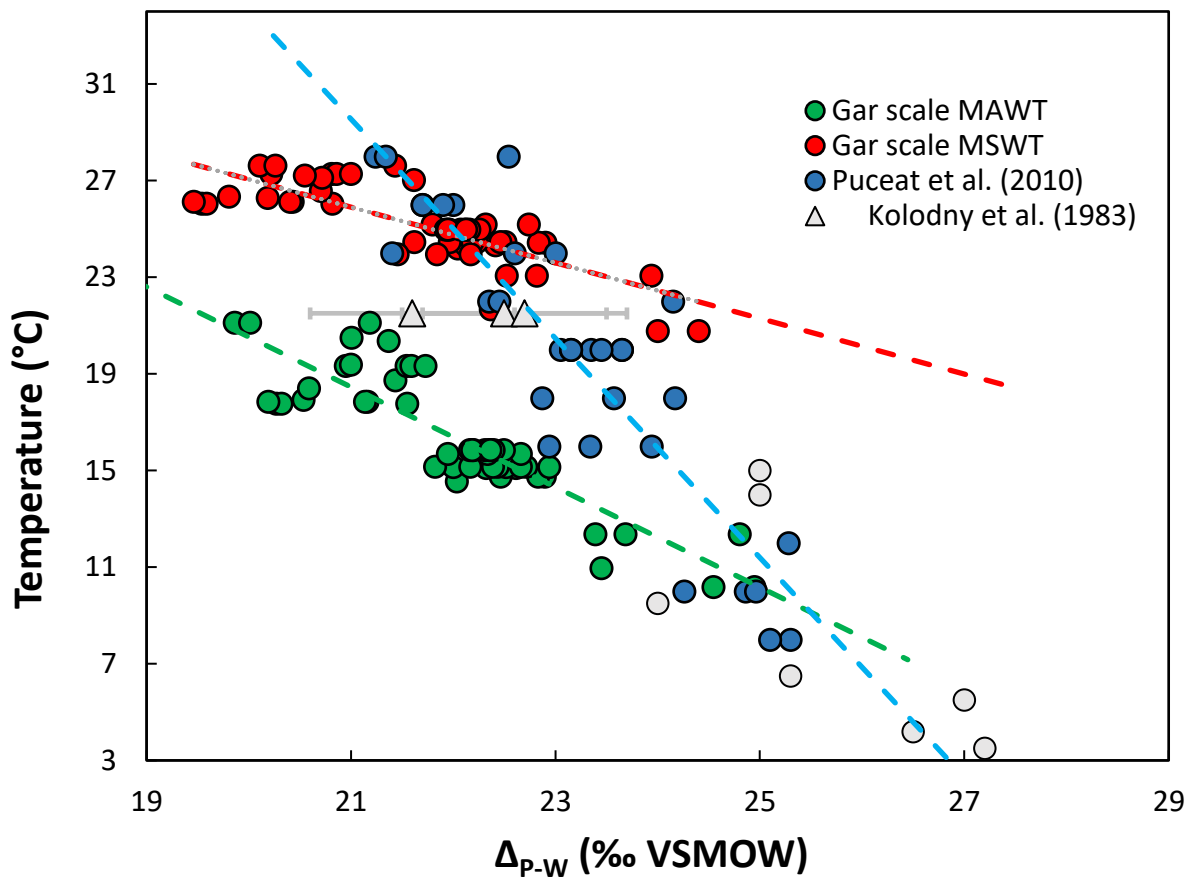


Figure 3. Data from this study (red and green circles) is compared to the values obtained by Pucéat et al. (2010) in their sea bream tank experiment (blue circles). Temperatures used are both mean annual water (green) and mean summer water (red). Gray triangles are from Kolodny et al. (1983), corrected by 2.1 ‰ to account for differences in lab methodology as outlined in Pucéat et al. (2010). Gray triangles and error bars are associated with Lake Kinneret (Sea of Galilee) samples. See section 4.2.

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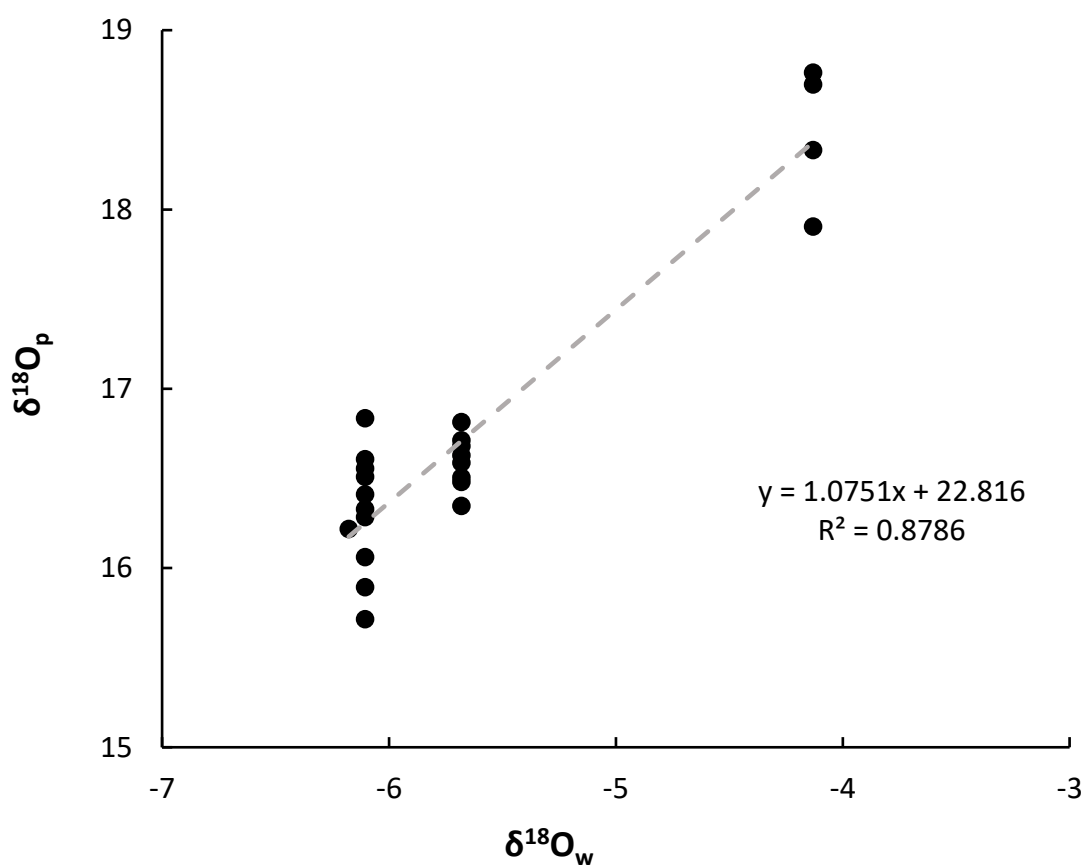


Figure 4. Comparison between $\delta^{18}\text{O}_{\text{water}}$ and $\delta^{18}\text{O}_{\text{phosphate}}$ when mean annual water temperature is held constant at $\sim 24.7^\circ\text{C}$, showing essentially a 1:1 ratio; values are from three different sites; Estill Springs, TN; Ullin, IL; and Peoria, IL.

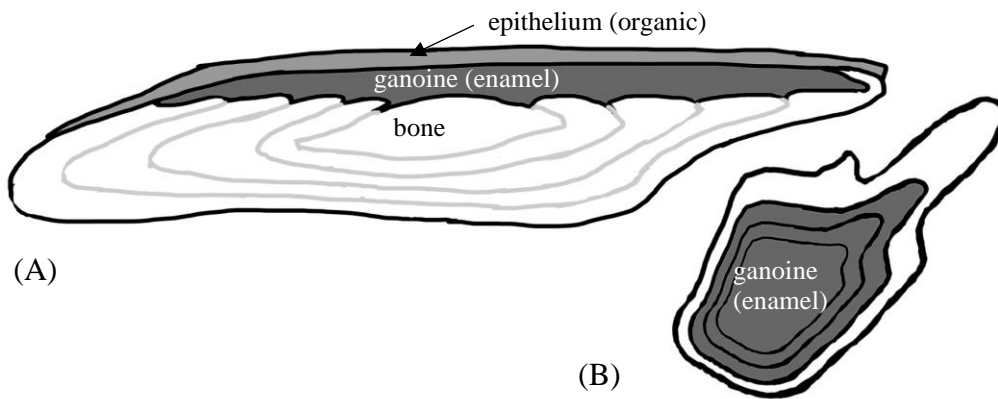


Figure 5. (A) Drawing of a cross section of a gar scale. Ganoine (enamel) directly overlies bone. Banding is seen in the bone with corresponding ridges in the ganoine. In vivo, an organic epithelium covers the scale, giving it color and pattern. (B) One of the three, and most common types of scales seen in gar fish, the dermal scale. Ganoine layering begins in the center (oldest) and extends outwards (youngest). Layering is for illustrative purposes only as banding is most prominent in the bone, not the enamel, and does not represent seasonal growth.

806 **Table 1.** Oxygen isotope values of gar scales. $\delta^{18}\text{O}_p$ is averaged from four replicates. $\delta^{18}\text{O}_w$ are
807 from USGS measurements (Kendall and Coplen, 2001); see Table S2 for water isotope stations.

Specimen Name	Annual Temperature (°C)	$\delta^{18}\text{O}_p$ (‰SMOW)	1 σ	Annual $\delta^{18}\text{O}_w$ (‰SMOW)	Δ_{p-w} (‰SMOW)
YPM 12016	9.15± 0.45	16.60	0.19	-7.94	24.55
YPM 8878		17.00	0.21		24.81
FMNH 85897	9.94± 0.82	16.53	0.33	-6.92	23.45
YPM 27219	11.35± 0.74	15.84	0.09	-7.55	23.39
YPM 27220		17.26	0.24		24.80
YPM 27221		16.14	0.18		23.68
YPM 27211	13.77± 0.77	18.76	0.23	-4.13	22.89
YPM 27212		18.33	0.03		22.83
YPM 27216		18.70	0.26		22.46
UF 129480	13.57± 0.72	17.90	0.19	-4.13	22.04
IC-05-01	14.17± 0.74	16.83	0.09	-6.10	22.94
IC-05-02		15.89	0.31		22.00
IC-05-03		16.41	0.39		22.52
ADY-01-01	14.13± 0.76	16.22	0.14	-6.10	22.32
ADY-01-02		16.51	0.36		22.61
IX-03-01	14.17± 0.74	16.33	0.19	-6.10	22.43
IX-03-05		15.71	0.16		21.82
IX-03-06		16.61	0.28		22.71
IX-03-09		16.06	0.06		22.17
IX-03-10		16.56	0.61		22.66
IX-03-11		16.28	0.13		22.39
YPM 11983	14.71± 0.44	17.44	0.33	-4.89	22.33
YPM 12006		17.06	0.26		21.95
YPM 12007		17.77	0.07		22.66
YPM 027686	14.92± 0.59	16.63	0.06	-5.68	22.31
YPM 027687		16.52	0.51		22.03
YPM 027688		16.67	0.19		22.27
YPM 027689		16.81	0.23		22.49
YPM 027690		16.48	0.17		22.16
YPM 027691		16.71	0.31		22.39
YPM 027692		16.50	0.44		22.19
YPM 027693		16.68	0.18		22.36
MMNS 59029	17.16 ± 0.21	17.23	0.14	-3.91	21.13
YPM 27218		16.36	0.05	-3.91	20.27
YPM 27223		17.64	0.41		21.55
YPM 27224		16.41	0.16		20.32
MMNS 51497	17.29 ± 0.31	16.63	0.15	-3.91	20.37
MMNS 58721	17.16 ± 0.21	16.29	0.19	-3.91	20.19
MMNS 58880	17.16 ± 0.21	17.25	0.26	-3.91	21.16
CFM 109203	17.87 ± 0.54	17.53	0.12	-3.91	21.43
MMNS 54500	17.82 ± 0.68	16.68	0.04	-3.91	20.61
MDF 1	18.40 ± 0.84	17.64	0.38	-3.64	21.54

MDF 2		17.68	0.26		21.59
MDF 3	18.40 ± 0.84	17.82	0.31	-3.64	21.73
MDF 4		17.04	0.50		20.95
BARNRES	18.45 ± 0.45	18.23	0.64	-2.77	21.00
UF 120183	19.44 ± 0.55	17.47	0.13	-3.90	21.45
UF 119897	20.20 ± 0.42	15.96	0.15	-3.90	20.96
UF 119908	20.20 ± 0.42	17.28	0.23	-3.90	21.27
UF 150169	19.83 ± 0.66	16.11	0.21	-3.90	19.43
TNSC 31670	19.39 ± 0.75	17.19	0.64	-3.81	21.00

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