Temperature dependence of clumped isotopes ([?]47) in aragonite

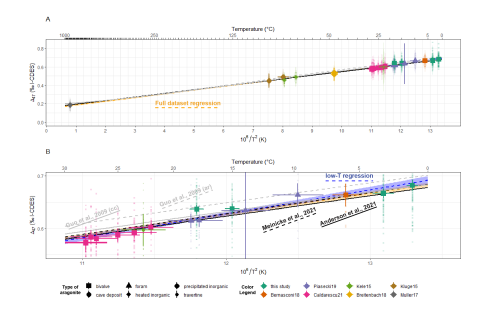
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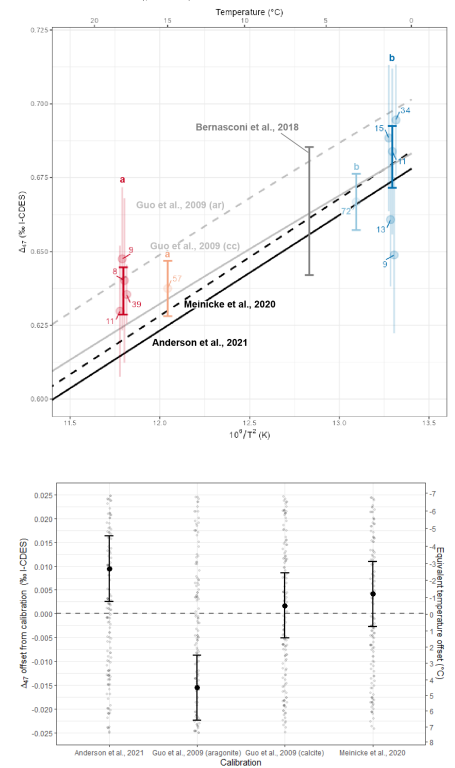
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Abstract

Clumped isotope thermometry can independently constrain the formation temperatures of carbonates, but a lack of precisely temperature-controlled calibration samples limits its application on aragonites. To address this issue, we present clumped isotope compositions of aragonitic bivalve shells grown under highly controlled temperatures (1-18°C), which we combine with clumped isotope data from natural and synthetic aragonites from a wide range of temperatures (1-850degC). We observe no discernible offset in clumped isotope values between aragonitic foraminifera, mollusks, and abiogenic aragonites or between aragonites and calcites, eliminating the need for a mineral-specific calibration or acid fractionation factor. However, due to non-linear behavior of the clumped isotope thermometer, including high-temperature (>100degC) datapoints in linear clumped isotope calibrations causes them to underestimate temperatures of cold (1-18degC) carbonates by 2.7 + 2.0degC (95% confidence level). Therefore, clumped isotope-based paleoclimate reconstructions should be calibrated using samples with well constrained formation temperatures close to those of the samples.



Arctica islandica Δ_{47} vs Temperature



1 Temperature dependence of clumped isotopes (Δ_{47}) in aragonite

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11 Key points

Precise control on carbonate formation temperatures enables more accurate clumped
 isotope-temperature calibrations

Isotopic ordering and acid fractionation in aragonite have a similar temperature
 dependence as in calcite, enabling combined calibrations

16 - The $\Delta_{47} - \frac{1}{T^2}$ relation in carbonates is non-linear, including hot calibration data offsets the 17 calibration in the cold temperature range

18

19 Abstract

Clumped isotope thermometry can independently constrain the formation temperatures of 20 carbonates, but a lack of precisely temperature-controlled calibration samples limits its 21 22 application on aragonites. To address this issue, we present clumped isotope compositions of aragonitic bivalve shells grown under highly controlled temperatures (1-18°C), which we 23 combine with clumped isotope data from natural and synthetic aragonites from a wide range 24 of temperatures (1-850°C). We observe no discernible offset in clumped isotope values 25 between aragonitic foraminifera, mollusks, and abiogenic aragonites or between aragonites 26 and calcites, eliminating the need for a mineral-specific calibration or acid fractionation factor. 27 However, due to non-linear behavior of the clumped isotope thermometer, including high-28 temperature (>100°C) datapoints in linear clumped isotope calibrations causes them to 29 30 underestimate temperatures of cold (1-18°C) carbonates by 2.7 ± 2.0°C (95% confidence level). Therefore, clumped isotope-based paleoclimate reconstructions should be calibrated 31 using samples with well constrained formation temperatures close to those of the samples. 32

33

34 Plain language summary

35 Clumped isotope analysis is a highly accurate method for reconstructing temperatures in 36 Earth's past climate from calcium carbonate fossils of calcifying organisms. Unfortunately, calibration studies so far were predominantly based on samples of calcite, a common calcium 37 38 carbonate mineral. It is therefore unknown whether these clumped isotope calibrations yield accurate temperature reconstructions when applied to aragonite, another carbonate mineral 39 which corals and many shells consist of. Therefore, we grew mollusks that build their shell out 40 of aragonite in a lab at constant water temperatures to test the clumped isotope method on 41 42 aragonitic shells. We find no significant difference in the temperature sensitivity of the method between our aragonites and the previous calibrations and show that the temperature 43 44 calibration can be improved by combining data from different minerals. However, we find subtle differences in the temperature dependence of clumped isotopes between hot (>100°C) 45 46 carbonates and cold (<30°C) carbonates, which cause previous calibrations to underestimate 47 temperatures of colder carbonates. We conclude that using carbonate samples grown at temperatures close to the temperatures of the samples used in climate reconstructions can 48 49 eliminate a bias of 2.7°C, resulting in more accurate reconstructions of past temperatures.

50

51 Keywords

52 Clumped isotopes, aragonite, paleoclimate, mollusk, temperature

53

54 **1. Introduction**

Since its first applications (e.g. Schauble et al., 2003; Wang et al., 2004; Ghosh et al., 2006), carbonate clumped isotope analysis has developed into a valuable tool for paleothermometry in the geosciences. Clumped isotope analysis is based on the thermodynamic principle that molecules with multiple heavy isotopes (so-called "multiply-substituted isotopologues") have lower vibrational energies than molecules containing lighter isotopes (Urey, 1947). Consequently, the increase in system entropy at higher temperatures causes a decrease in

the occurrence of multiply-substituted isotopologues, and "clumping" of heavy isotopes within 61 the same molecule is favored in low-energy systems (Eiler, 2007). In carbonates, this principle 62 causes heavy carbonate ions (e.g. ¹³C¹⁸O¹⁶O₂; mass 63 or ¹²C¹⁸O₂¹⁶O; mass 64) to become 63 more abundant with decreasing calcification temperatures (Ghosh et al., 2006). The 64 distribution of these isotopologues is proportional in the CO₂ gas after reaction of carbonates 65 with acid (e.g. ¹³C¹⁸O¹⁶O; mass 47 and ¹²C¹⁸O₂, mass 48 respectively) and is measured with 66 reference to the distribution of isotopologues in a fully scrambled heated CO₂ gas with the 67 same isotopic composition: 68

69
$$\Delta_{47}[\%_0] = \left(\frac{R^{47}}{R^{47}*} - 1\right) \quad (1)$$

In which R^{47} is the ratio of CO_2 molecules with mass 47 (predominantly ¹³C¹⁸O¹⁶O) relative to CO₂ with the most common mass 44 (¹²C¹⁶O₂) in the sample, and R^{47*} is the same ratio in stochastic equilibrium (Daëron et al., 2016). This Δ_{47} value is a measure for the degree of "clumping" in the sample which depends on its calcification temperature.

74 The main advantage of carbonate clumped isotope analysis over previous paleothermometers is its basis on thermodynamic principles and its independence from the chemistry of the 75 76 precipitation fluid (Eiler, 2007). The latter represents an improvement over the often-used oxygen isotope paleothermometer (δ^{18} O), which requires knowledge of the oxygen isotope 77 composition of the precipitation fluid (δ¹⁸O_w; e.g. Epstein et al., 1953; Kim & O'Neil, 1997). The 78 79 clumped isotope method has many applications, notably to reconstruct absolute temperature 80 variability throughout Earth's history (Rodríguez-Sanz et al., 2017; Henkes et al., 2018; Vickers et al., 2020a; de Winter et al., 2021a; Meckler et al., 2022; Agterhuis et al., 2022). 81

Inter-lab standardization of carbonate Δ_{47} measurements has reconciled former offsets between laboratories using different CO₂ preparation methods and reconciled the clumped isotope temperature calibration of calcites with the results of thermodynamic *ab initio* models (Bernasconi et al., 2018; 2021; Petersen et al., 2019; Jautzy et al., 2020). A unified linear calibration was established through re-standardized Δ_{47} values of carbonates precipitated at a wide range of known temperatures (0.5-1100°C; Anderson et al., 2021). This eliminates
concerns over the confounding effects of differences in the origin of carbonates (e.g. biogenic
vs. inorganic; Henkes et al., 2013), varying mineralization rates (Daëron et al., 2019), different
acid digestion temperatures and different carbonate mineralogies (e.g. dolomite vs. calcite;
Müller et al., 2019) on the clumped isotope thermometer. However, it remains unclear whether
biological process (i.e. "vital effects") influence isotopic ordering in some biogenic carbonates.

93 The unified calibration dataset includes only one aragonitic carbonate, insufficient to test for different clumped isotope temperature dependencies between aragonites and calcites 94 (Anderson et al., 2021). Results of ab initio models suggest that such a difference between the 95 two polymorphs may exist (Schauble et al., 2006; Guo et al., 2009) and experimental studies 96 97 disagree on a difference in acid fractionation factor (AFF) between calcite and aragonite (Guo et al., 2009; Müller et al., 2019; Petersen et al., 2019). These uncertainties are confounded by 98 the fact that most carbonates used in current calibrations are precipitated under natural 99 circumstances with indirectly estimated or else poorly controlled temperature regimes (e.g. 100 Kele et al., 2015; Peral et al., 2018). The potential Δ_{47} offset between aragonite and calcite 101 102 might introduce an unknown bias when using the unified temperature calibration on aragonite data (e.g. Caldarescu et al., 2021); a severe limitation given the common occurrence of 103 aragonite in biogenic calcifiers (e.g. bivalves; Kennedy et al., 1969, gastropods; Taylor and 104 105 Reid, 1990, and foraminifera; Hansen, 1979) as well as inorganic natural carbonates (e.g. 106 speleothems; Frisia et al., 2000, and travertines; Kele et al., 2015).

This study presents new clumped isotope results from precisely temperature controlled, labgrown aragonitic *Arctica islandica* bivalve shells. The bivalve *Arctica islandica* is a highly utilized climate archive, and a promising substrate for clumped isotope-based paleothermometry (e.g. Witbaard et al., 1997; Burchardt and Simonarson, 2003; Schöne et al., 2005; Schöne and Fiebig, 2009; Butler et al., 2013). Combined with preexisting aragonite clumped isotope data (Kluge et al., 2015; Kele et al., 2015; Müller et al., 2017; Breitenbach et al., 2018; Bernasconi et al., 2018; Piasecki et al., 2019; Caldarescu et al., 2021) standardized to the new Intercarb-Carbon Dioxide Equilibrium Scale (I-CDES) reference frame (Bernasconi
et al., 2021), our dataset resolves potential vital effects on clumped isotopes in aragonitic
mollusks by comparing species and specimens grown under the same controlled conditions.
This study aims to offer a detailed investigation of the clumped isotope temperature
dependence in aragonites.

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120 2. Materials and Methods

121 2.1 Lab grown Arctica islandica

122 Arctica islandica bivalves were cultured inside the lab of the Royal Netherlands Institute for 123 Sea Research (NIOZ, Texel, the Netherlands). Specimens used for this study were grown under four different, constant, and monitored temperature regimes: $1.1 \pm 0.2^{\circ}$ C, $3.2 \pm 0.3^{\circ}$ C, 124 $15 \pm 0.4^{\circ}$ C and $18 \pm 0.3^{\circ}$ C (see **Table 1**; **S9**). Details on the culturing setup are provided in **S1** 125 and Witbaard et al. (1998). Aragonite from cleaned and dried Arctica islandica shells was 126 127 sampled using a hand-held Dremel 3000 rotary drill at low speed equipped with a tungsten-128 carbide drill bit (see **S1**). Gathering enough aragonite for reliable Δ_{47} analyses for each temperature treatment (>2 mg; Müller et al., 2017; Fernandez et al., 2017) typically required 129 combining material from multiple (3-5) specimens grown under the same temperature 130 131 conditions. To test potential inter-specimen differences, results were tracked per individual specimen for the $1.1 \pm 0.2^{\circ}$ C and $18 \pm 0.3^{\circ}$ C treatments (see **Table 1**). 132

133 2.2 Clumped isotope analysis

The clumped isotope composition of 278 aliquots of shell aragonite were analyzed over two 6month periods (March – August 2020; May – November 2021) on two Thermo isotope ratio mass spectrometers (one MAT253 and one MAT253 plus) coupled to Kiel IV carbonate preparation devices (see **S1**). After correcting for variability in the pressure baseline (He et al.,2012), clumped isotope results were processed relative to the I-CDES through an empirical transfer function (ETF) based on measurements of ETH standards (ETH-1, -2 and -3) and their

accepted I-CDES values (Bernasconi et al., 2021). Isotopic values were calculated using the 140 latest IUPAC values (Brand et al., 2010; Daëron et al., 2016). No AFF was applied after I-141 CDES standardization because the carbonate standards used for the ETF undergo the same 142 143 acid reaction as the samples (Bernasconi et al., 2021). Long-term accuracy and reproducibility of Δ_{47} results were assessed based on repeated measurements of the IAEA-C2 monitoring 144 standard (Δ_{47} IAEA on MAT253 plus: 0.6382 ± 0.026‰; Δ_{47} IAEA on MAT253: 0.6445 ± 0.046‰; 145 1 σ). Results were indistinguishable from the accepted value for IAEA-C2 (0.6409 ± 0.003‰; 146 147 95% CL; Bernasconi et al., 2021). Full results of all sample aliquots and standards used to standardize the results are provided in S2. 148

149 2.3 Data compilation

The Arctica islandica dataset was augmented with literature Δ_{47} values of aragonites with 150 known calcification temperatures (see S3). The dataset includes samples from mollusks 151 (Bernasconi et al., 2018; aragonitic Megapitaria aurantiaca samples in Caldarescu et al., 2021; 152 this study), foraminifera (Piasecki et al., 2019), travertines (Kele et al., 2015; Bernasconi et al., 153 2018), cave deposits (Breitenbach et al., 2018), lab-grown aragonites (Kluge et al., 2015) and 154 heated aragonites (Müller et al., 2017). Data from several older studies (e.g. Ghosh et al., 155 2006; 2007; Tripati et al., 2010; Wacker et al., 2013; 2014; Zhang et al., 2018; Zhai et al., 2019; 156 Dong et al., 2021) were not included because they were not corrected for the pressure baseline 157 (He et al., 2012; Bernasconi et al., 2013), could not be transferred into the standard reference 158 frame (Dennis et al., 2011), lacked the standardization required to bring Δ_{47} values into the I-159 CDES scale (Bernasconi et al., 2021) or because the aragonite was precipitated out of 160 equilibrium (e.g. Kimball et al., 2015; Chen et al., 2019; S1; S3). Clumped isotope data from 161 the literature was brought to the I-CDES reference frame using the multi-linear correction 162 proposed in Appendix A of Bernasconi et al. (2021) using values of carbonate standards 163 164 reported in the studies (see S1). Uncertainties on the formation temperatures of the non-165 temperature controlled datapoints from previous studies were generally in the order of 1°C (1o; see **S1**). The full dataset including Δ_{47} values and temperatures with their uncertainties used 166

- in this study is provided in S4. Unless stated otherwise, uncertainties are cited at the 95%confidence level.
- All data processing for this study is done in R (R Core Team, 2022) and scripts are provided in **S5** and published on Github (<u>https://github.com/nielsjdewinter/Aragonite_clumped</u>). Details on data processing are provided in **S1**. We compare our data with calibrations by Anderson et al. (2021) and Meinicke et al. (2020) as well as with temperature dependencies of aragonite and calcite clumped isotope compositions from *ab initio* modelling in Guo et al. (2009) brought into the I-CDES reference frame (see **S1**).

175 3. Results

176 3.1 Clumped isotope values in Arctica islandica

Clumped isotope results from A. islandica are summarized in Table 1 and Figure 1. There is 177 178 no significant clumped isotope difference between specimens in the same temperature treatment (F(4,77) = 1.937, p = 0.11 for the 1°C specimens and F(3,63) = 0.377, p = 0.77 for 179 the 18°C specimens; see S6). The number of measurements per specimen was large enough 180 to exclude per-specimen Δ_{47} differences outside the reproducibility standard deviation of the 181 182 clumped isotope analyses (0.046‰; see Table 1 and S6). Differences between all temperature treatments are statistically significant (P(3,274) = 15.68, p < 0.01), except for differences 183 184 between the 15°C and 18°C temperature bin and the difference between 1°C and 3°C (95%CL; 185 S6).

We investigated the $\Delta_{47} - \frac{1}{T^2}$ relationship and how it varies along the temperature domain by 186 performing linear regressions on increasingly large parts of our compilation. Note that the 187 uncertainty on clumped isotope data compared to the range of temperatures of the lab-grown 188 A. islandica leaves relatively high uncertainty on a clumped isotope-temperature regression 189 through these results alone compared to the unified clumped isotope calibration (Anderson et 190 191 al. 2021). We therefore do not advice using this and other regression equations in this section for calibrating clumped isotope results (see Discussion). Firstly, a statistically significant 192 temperature relationship (Δ_{47} - $\frac{1}{T^2}$ slope >0; 95% CL) is found for Δ_{47} exclusively from Arctica 193 islandica samples: 194

$$\Delta_{47}(I - CDES) = 0.0280 \pm 0.0042 * \frac{10^6}{T^2} + 0.304 \pm 0.0524 \text{ (T in K, } \pm 1\sigma; \sigma_{res} = 0.047\%)$$
(1)

Secondly, including other aragonitic mollusk data (Caldarescu et al., 2021) yields a regression
indistinguishable from the Anderson et al. (2021) unified clumped isotope calibration:

$$\Delta_{47}(I - CDES) = 0.0443 \pm 0.0024 * \frac{10^6}{T^2} + 0.097 \pm 0.0291 \text{ (T in K, } \pm 1\sigma; \sigma_{res} = 0.043\%)$$
(2)

198 3.2 Aragonite clumped isotope temperature dependence

When including clumped isotope values of other low-temperature (<30°C) aragonites in the compilation, the regression remains indistinguishable from the calibration of Anderson et al. (2021) and similar to Meinicke et al. (2020; 2021) and the Guo et al. (2009) theoretical temperature relationships (**Fig. 2B**):

$$\Delta_{47}(I - CDES) = 0.0451 \pm 0.0024 * \frac{10^6}{T^2} + 0.0871 \pm 0.0287 \text{ (T in K, } \pm 1\sigma; \sigma_{res} = 0.042\%)$$
(3)

Finally, we included higher temperature (>30°C) datapoints, such as the cave deposits of Breitenbach et al. (2018), travertine samples from Kele et al. (2015), precipitated aragonites from Kluge et al. (2015) and heated aragonites from Müller et al. (2017) in the linear regression. This decreases the slope and increases the intercept (see **Fig. 2**):

$$\Delta_{47}(I - CDES) = 0.0403 \pm 0.0005 * \frac{10^6}{T^2} + 0.1435 \pm 0.0485 \text{ (T in K, } \pm 1\sigma; \sigma_{res} = 0.040\%)$$
(4)

The formation temperatures of our A. islandica data on the very cold end of the calibration 207 domain are significantly underestimated by Anderson et al. (2021; $\Delta\Delta_{47} = +0.009 \pm 0.007$ %; 208 -2.71 ± 2.03 °C; **Fig. 3**; $\Delta\Delta_{47}$ = offset between data and calibration). The theoretical aragonite 209 clumped isotope-temperature relationship (Guo et al., 2009) severely overestimates A. 210 islandica temperatures (-0.016 ± 0.007‰; +4.35 ± 1.88°C; Fig. 3). Contrarily, the Meinicke et 211 al. (2020; 2021) calibration ($\Delta \Delta_{47}$ = +0.004 ± 0.007‰; -1.17 ± 2.00°C; Fig. 3) and the 212 theoretical calcite temperature relationship (Guo et al., 2009; $\Delta\Delta_{47}$ = +0.002 ± 0.007‰; -0.47 213 214 ± 1.98°C; Fig. 4) do not significantly over- or underestimate the formation temperature of our A. islandica shells. 215

216 4. Discussion

217 *4.1 Isotope ordering in aragonitic mollusks*

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Clumped isotope values of our temperature-controlled A. islandica samples consistently plot 218 on a Δ_{47} - $\frac{1}{T^2}$ linear relationship with other low-temperature aragonite datapoints (**Fig. 1 and 2**; 219 see section 4.2). The absence of a consistent offset between A. islandica datapoints and other 220 aragonites (mean Δ_{47} difference of +0.003 ± 0.004‰, see Fig. 2 and S8) and agreement 221 between the linear $\Delta_{47} \frac{1}{T^2}$ dependence of the aragonitic mollusk data in this study and the 222 regression through the complete low-temperature aragonite dataset (Fig. 1 and section 3.1) 223 224 strongly supports a common temperature dependence for all aragonites in this study, biogenic 225 or inorganic, and argues against disequilibrium fractionation in aragonite precipitated 226 inorganically or vital effect in bivalves or foraminifera (see section 3.1; Fig. 1 and 2). Our 227 highly temperature-controlled growth experiments uniquely allow us to exclude variability in the growth environment between specimens from the same growth treatment as a driver of 228 229 shell composition. Strong similarity of Δ_{47} values between individual A. islandica specimens 230 grown at the same temperature thus rules out specimen-specific vital effects on the clumped isotope composition aragonitic bivalve shells outside the uncertainty of our measurements (see 231 section 3.1; Fig. 1, Table 1 and S6). These findings corroborate measurements in calcitic 232 mollusks showing that clumped isotope values in mollusk carbonates adhere to the same 233 234 temperature relationship as other carbonates precipitated in equilibrium (except for juvenile oyster shells; Huyghe et al., 2022). Clumped isotope analyses in (fossil) mollusk shells thus 235 provide an independent temperature proxy, allowing paleoclimatologists to disentangle the 236 effects of variability in temperature and the hydrological cycle (as measured in $\delta^{18}O_w$) 237 throughout geological history down to the seasonal timescale (e.g. Caldarescu et al., 2021; de 238 Winter et al., 2021; Letulle et al., 2022). 239

240 4.2 Mineral-specific acid fractionation factor

Residuals of aragonite clumped isotope data around the low-temperature (<30°C) York regression (0.042‰; 1 σ ; see **section 3.1** and **Fig. 2**) are predominantly explained by analytical uncertainty on Δ_{47} measurements (external precision of 0.026‰ and 0.046‰ on the 253Plus and the MAT253 mass spectrometers; 1 σ ; see **section 2.2**). Uncertainty on formation

temperatures in the low-temperature dataset (±0.8°C; 1o; see S4) would add an additional 245 uncertainty of 0.0024‰ (1o) if applied to the weighted average formation temperature of all 246 low-temperature (<30°C) data points (22.0°C; see S4). Outside these uncertainties in the 247 248 compilation data, there is little uncertainty on the temperature relationship in the lowtemperature domain (<30°C; see section 3.2; Fig. 2). If clumped isotope fractionation during 249 acid digestion was indeed different between aragonite and calcite (as suggested in Müller et 250 al., 2017; Petersen et al., 2019) the difference in AFF would be indicated by the difference 251 252 between our aragonite dataset and the previous calcite-based calibrations (Meinicke et al., 2020; 2021). The close similarity between our A. islandica data and the calcite calibration ($\Delta \Delta_{47}$ 253 = 0.004 ± 0.007 ; Fig. 3; S7) leaves little room for the 0.007 and 0.025 difference in AFF 254 reported in Petersen et al. (2019) and Müller et al. (2017), respectively. We therefore conclude 255 256 that the calcite AFF in Petersen et al. (2019), which are included in the I-CDES reference scale (Bernasconi et al., 2021) can be used for aragonite samples. 257

4.3 Non-linear temperature dependence of clumped isotopes in aragonites

Current clumped isotope calibrations (Meinicke et al., 2020; 2021; Anderson et al., 2021) show 259 subtle differences in the low-temperature end of the calibration (<30°C) that would result in 260 ~1.5°C colder temperatures when applying Anderson et al. (2021) compared to Meinicke et al. 261 (2020). In addition, the cold-water (<30°C) carbonate based Meinicke et al. (2020) calibration 262 more closely resembles the modelled temperature relationship for calcites in Guo et al. (2009). 263 Including high-temperature (>30°C) data in our linear regression leads to overestimation of the 264 265 temperature of warmer (>18°C) datapoints ($\Delta\Delta_{47}$ of -0.005 ± 0.006‰, or +1.8^{+2.1}°C for data 266 precipitated at 30°C), while underestimating colder datapoints ($\Delta\Delta_{47}$ of +0.009 ± 0.008‰, or $-2.0^{+2.0}_{-2.0}$ °C for data precipitated at 0°C; Fig. 2; S7). Point-by-point offsets of all data from the 267 268 calibration lines are provided in S8.

269 This difference between $\Delta_{47} - \frac{1}{T^2}$ regressions through the low-temperature (<30°C) and the full 270 dataset (see **section 3.2**; **Fig 2**) likely highlights non-linear behavior of the $\Delta_{47} - \frac{1}{T^2}$ relationship 271 in aragonites. In fact, previous studies based on both clumped isotope analyses and ab initio modelling have suggested a non-linear $\Delta_{47} - \frac{1}{T^2}$ relationship to be a better fit for both calcites 272 (Guo et al., 2009; Jautzy et al., 2020) and dolomites (Guo et al., 2009; Müller et al., 2019) 273 274 precipitated on a large range of known temperatures. Non-linear behavior is also observed in the Anderson et al. (2021) dataset, where Δ_{47} values of calcites precipitated between 100°C 275 and 1000°C are underestimated by the linear relationship, while the hottest datapoints (calcites 276 heated to 1100°C) fall on the linear regression, mimicking the reduced $\Delta_{47} - \frac{1}{T^2}$ slope at the high 277 temperature end of the polynomial regressions through calcite and dolomite data (Guo et al., 278 2009; Jautzy et al., 2020; Müller et al., 2019). A linear $\Delta_{47} \frac{1}{T^2}$ relationship through a calibration 279 dataset with a large temperature range will thus overestimate temperatures for samples with 280 281 Δ_{47} values between 0.2‰ and 0.4‰ (temperatures of 100°C–1000°C; see residuals in Anderson et al., 2021) and underestimate temperatures of cold (<30°C) samples, as confirmed 282 by regressions through our low-temperature datapoints (see Fig. 2-3 and section 4.4). 283 Therefore, more high-temperature aragonite datapoints are needed to constrain the clumped 284 isotope-temperature relationship for temperatures >100°C. 285

4.4 Calibrating the clumped isotope-temperature relationship in cold (<30°C) carbonates

287 Our lab-grown A. islandica shells offer more control on formation temperature than naturally grown carbonates precipitated under variable temperatures. Ideally, the temperature of these 288 natural samples is monitored so an average temperature can be calculated for the targeted 289 growth period (e.g. Kele et al., 2015; de Winter et al., 2020; 2021b; Huyghe et al., 2021). 290 However, formation temperatures are often indirectly estimated through other proxies (e.g. 291 292 $\delta^{18}O_c$) and/or estimates of the living environment (e.g. water depth) of the carbonate producer, accumulating uncertainty (e.g. Peral et al., 2018; Piasecki et al., 2018; Meinicke et al., 2020). 293 294 These caveats obscure the full uncertainty of the formation temperatures of natural carbonates as well as the effect of this unknown uncertainty on the calibrations. Considering the methods 295 by which the "known" temperatures of natural carbonates are estimated in previous studies, 296

297 part of the ~1.5°C temperature offset between Anderson et al. (2021) and Meinicke et al. (2020; 2021; see Fig. 3) and the 2.7 \pm 2.0°C offset between Anderson et al. (2021) and our A. 298 299 islandica data might be caused by uncertainty on the formation temperatures of the calibration 300 dataset. However, our highly temperature-controlled A. islandica datapoints reveal that, despite uncertainty on formation temperature, the Meinicke et al. (2021) calibration locally 301 approximates the non-linear Δ_{47} - $\frac{1}{T^2}$ relationship in the cold temperature domain with higher 302 accuracy than the Anderson et al. (2021) calibration (Fig. 1, Fig. 3; S8). The non-linear 303 theoretical calcite temperature dependence by Guo et al. (2009) also fits well with the data. 304 Precisely temperature-controlled carbonates thus better constrain the slope of the $\Delta_{47} - \frac{1}{\tau^2}$ 305 relationship for cold carbonates (improving calibration accuracy) while reducing the uncertainty 306 on the calibration (improving calibration precision). 307

308 The ~1.5°C difference in reconstructed temperature between the calibrations in the low 309 temperature range (<30°C) may seem trivial and requires the complete A. islandica dataset (N 310 = 278; see Fig. 4) to resolve. However, in paleoclimate reconstructions (e.g. Petersen et al., 311 2016; de Winter et al., 2017; 2021a; Vickers et al., 2020b; Agterhuis et al., 2021; Meckler et 312 al., 2022), this temperature offset may have significant consequences. A ~1.5°C cold bias in 313 temperature reconstructions may lead to a significant underestimation of climate sensitivity to 314 CO₂ forcing, biasing the physical science basis for informing policymakers about future climate 315 change (e.g. Dennis et al., 2013; Modestou et al., 2020; Westerhold et al., 2020; Tierney et al., 2020; IPCC, 2021). Accurate clumped isotope-based temperature reconstructions 316 317 therefore require calibration datasets with precisely constrained formation temperatures 318 tailored to the temperature range of the samples.

319

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329 **Open Research**

- 330 Supplementary materials are deposited on the open-source repository Zenodo and can be
- accessed through the following link: <u>https://doi.org/10.5281/zenodo.6524705</u>. R scripts are
- 332 uploaded on GitHub (https://github.com/nielsjdewinter/Aragonite_clumped) and archived in
- 333 Zenodo (<u>https://doi.org/10.5281/zenodo.6560188</u>).

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Table 1 Arctica islandica clumped isotope results compared to previous calibrations "mixed" = combined samples from multiple specimens, number codes (e.g. "29" or "6A") = material from one individual. "Ais" = Arctica islandica. Significant Δ_{47} and temperature offsets are labeled in bold. cc = calcite, ar = aragonite

Sample	Culturing temperature		∆ ₄₇ (I-CDES; ± 95%CL)	N	Offset from Anderson	Offset from Meinicke	Offset from Guo (cc)	Offset from Guo (ar)
Ais1	1.1 ± 0.2°C	mixed	0.695 ± 0.019‰	34				
		3	0.661 ± 0.023‰	13				
		29	0.688 ± 0.025‰	15				
		6A	0.684 ± 0.028‰	11				
		6B	0.649 ± 0.026‰	9				
		TOTAL	0.682 ± 0.010‰	82	+0.008‰	+0.002‰	+0.003‰	-0.016‰
					-2.12°C	-0.61°C	-0.75°C	+4.48°C
Ais3	3.2 ± 0.3°C	mixed	0.667 ± 0.010‰	72	+0.001‰	-0.005‰	-0.006‰	-0.023‰
A135	5.2 ± 0.5 C				-0.27°C	+1.24°C	+1.57°C	+6.80°C
Ais15	15 ± 0.4°C	mixed	0.637 ± 0.009‰	57	+0.013‰	+0.008‰	+0.004‰	-0.013‰
AISTO	10 ± 0.4 0	mixeu	0.007 ± 0.005/0		-3.63°C	-2.10°C	-0.99°C	+4.25°C
	18 ± 0.3°C	mixed	0.635 ± 0.010‰	39				
Ais18		67	0.647 ± 0.024‰	9				
		89	0.640 ± 0.028‰	8				
		111	0.630 ± 0.022‰	11				
		TOTAL	0.637 ± 0.005‰	67	+0.021‰ -6.63°C	+0.016‰ -5.10°C	+0.012‰ -3.99°C	-0.004‰ +1.25°C

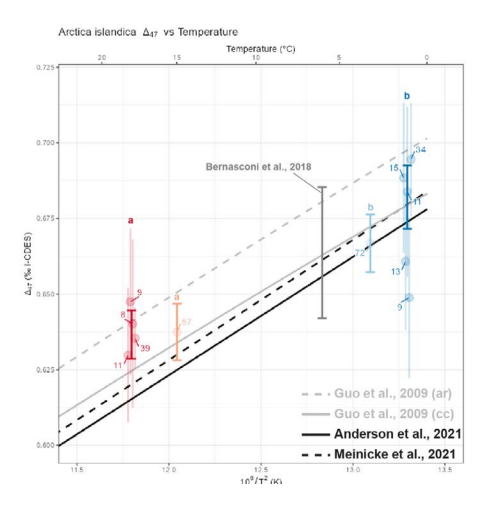


Figure 1: *Arctica islandica* Δ_{47} **results**. Clumped isotope results are aggregated by specimen or multi-specimen sample (round symbols; see **Table 1**). Vertical lines represent 95% CL and number indicate sample size. Data is color-coded per temperature treatment (1°C, 3°C, 15°C and 18°C), with bold error bars indicating 95% CL, pairs of letter labels (a and b) indicate statistically indistinguishable Δ_{47} values (p < 0.05). The grey error bar at 6°C highlights *A. islandica* data from Bernasconi et al. (2018; recalculated to I-CDES). Solid and dashed black lines show calibrations by Anderson et al. (2021) and Meinicke et al. (2020; projected on I-CDES scale; Meinicke et al., 2021), respectively. Grey solid and dashed lines represent, respectively, the theoretical calcite ("cc") and aragonite ("ar") temperature dependencies from Guo et al. (2009; projected on the I-CDES scale, see **S1**). The horizontal axis is scaled to $\frac{10^6}{T^2}$, with T in K, to show the assumed linear relationship with the clumped isotope value.

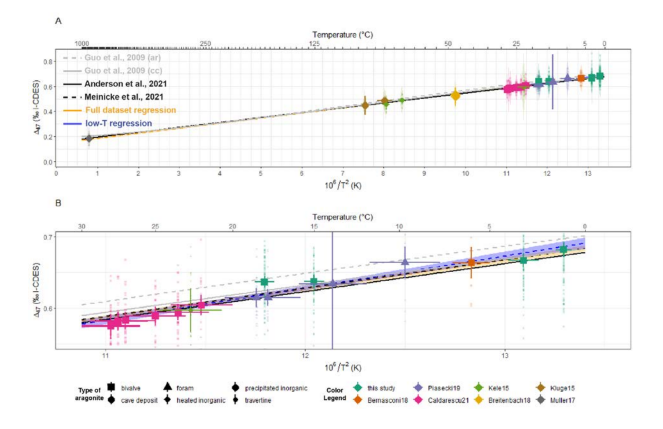


Figure 2: Aragonite Δ_{47} **temperature dependence.** Clumped isotope data of aragonite samples plotted against formation temperature. **A**. All data plotted over the full temperature range (1°C–850°C). Individual datapoints, averages and uncertainty on temperature and Δ_{47} values (95% CL) are color-coded by study. Symbols highlight different types of aragonite. The solid and dashed black lines show calibrations by Anderson et al. (2021) and Meinicke et al. (2020; 2021; plotted for temperatures <30°C). Grey solid and dashed lines represent, respectively, the theoretical calcite ("cc") and aragonite ("ar") temperature dependencies from Guo et al. (2009; projected on the I-CDES scale, see **section 2.5**). Colored dashed lines and shaded envelopes show York regressions through aragonite data and their 95% confidence envelopes, respectively. **B**. Shows a zoom-in of the plot in **A**. for the low-temperature domain (1–30°C). Note that the horizontal axis is scaled to $\frac{10^6}{T^2}$, with T in K, to show the assumed linear relationship with the clumped isotope value.

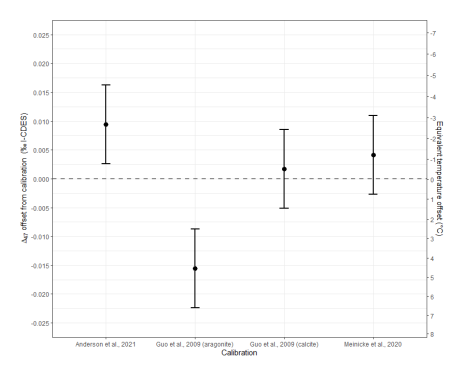


Figure 3: Offset of *A. islandica* data from temperature regressions. Shaded grey points show residual Δ_{47} values relative to four clumped isotope temperature relationships (see horizontal axis). Black symbols with error bars (95% CL) show mean offsets of all *A. islandica* datapoints (grown at 1°C, 3°C, 6°C, 15°C and 18°C) from the calibrations. The vertical axis on the right shows the temperature offset relative to the weighted mean calcification temperature of the full *A. islandica* dataset (8.6°C; see **S4**) based on Anderson et al. (2021).



Geophysical Research Letters

Supporting Information for

Temperature dependence of clumped isotopes (Δ_{47}) in aragonite

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Introduction

This Supplement contains all supplementary text and figures for the manuscript. All data files, scripts and large tables are deposited in the accompanying Zenodo repository (<u>https://doi.org/10.5281/zenodo.6524705</u>, see Open Research chapter).

Text S1: Supplementary Methodology

1. Culture experiments

Living specimens of juvenile *Arctica islandica* (bivalvia) were collected from the Süderfart site in the Kieler Bucht to the north-east of Kiel, Germany (54°34'11"N, 10°51'59"E; see Schaefer et al., 1985). All animals were placed in sand-filled containers in a basin with aerated seawater which

was refreshed two days a week with water from the Marsdiep tidal inlet directly outside the Royal Netherlands Institute for Sea Research (NIOZ, Texel, the Netherlands; 53°00'04"N, 4°47'23"E). Specimens used for this study were grown under four different, constant, and monitored temperature regimes: 1.1 ± 0.2 °C, 3.2 ± 0.3 °C, 15 ± 0.4 °C and 18 ± 0.3 °C and fed *ad libidum* with a suspension of *Isochrysis galbana* and *Dunaliella marina* algae. Specimens were left to acclimatize to the culturing conditions for four weeks. During the experiments, shell growth, temperature, phytoplankton cell counts, chlorophyll concentration, particulate organic carbon content and siphon activity were monitored. The experiment lasted for 95 days, after which the specimens were euthanized, and soft tissue was removed from the shells. Clean shells were rinsed with fresh water, dried at room temperature (~20°C), and stored under cool and dry conditions after the growth experiment. Additional details on the growing conditions of the specimens and the setup of the experiments can be found in Witbaard et al. (1998).

2. Sampling strategy

Aragonite from cleaned and dried shells from these lab-grown *Arctica islandica* specimens was sampled using a hand-held Dremel 3000 rotary drill at low speed equipped with a tungstencarbide drill bit. Minimal pressure was applied to carefully flake off parts of the shell and prevent heating of the samples due to friction, which may alter the Δ_{47} value of the aragonite (Staudigel and Swart, 2016). Care was taken to only sample the part of the shells that grew during the experiment. Pre-experimental shell material was easily avoided due to a clear growth line and color change that separated natural from lab-grown shell material (see **Fig. S1**).

3. Clumped isotope analyses

Aragonite aliquots were reacted with nominally anhydrous (103%) phosphoric acid at 70°C. The produced CO₂ gas was led through two liquid nitrogen-cooled (-196°C) cryogenic traps and a

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PoraPakTM Q trap (Merck KGaA, Darmstadt, Germany) kept at -40°C or (after October 2020) -50°C through a custom-built external cooling unit (Dennis and Schrag, 2010). The purified CO₂ gas was analyzed in micro-volume mode using the LIDI workflow with 400 s integration time against a clean CO₂ working gas (δ^{13} C = -2.82‰; δ^{18} O = -4.67‰) and corrected for pressure baseline effects (Bernasconi et al., 2013; Meckler et al., 2014; Müller et al., 2017). Clumped isotope values were corrected to the Intercarb-Carbon Dioxide Equilibrium Scale (I-CDES) by creating an empirical transfer function (ETF) using measurements of ETH standards (ETH-1, -2 and -3) and their accepted Intercarb values (Bernasconi et al., 2021). We applied the ETF in a moving window of 200 analyses before and after the sample, considering standards measured within a time window of 2–3 weeks surrounding the sample for its correction. This typically resulted in 13 ETH-1, 13 ETH-2 and 70 ETH-3 measurements to constrain the ETF for each sample aliquot. Higher amounts of ETH-3 standards were analyzed to better constrain uncertainties around the expected Δ_{47} values of samples (Kocken et al., 2019). Measurements of the ETH-3 standard were also included every 3–5 samples to check for measurement drift at shorter timescales than the moving window of the ETF.

4. Data compilation

All included datasets except for Kluge et al. (2015) reported measured values for the ETH-1, -2 and -3 standards, yielding three anchor points for the reference scale calibration. The Kluge et al. (2015) data was calibrated to I-CDES using the ETH-3 and Carrara marble standard using the simplified linear correction in Appendix A of Bernasconi et al. (2021). To obtain an I-CDES Δ_{47} value for the Carrara marble standard (which is not reported in Bernasconi et al., 2021), we averaged the 19 replicates of Carrara marble reported in Bernasconi et al. (2018) after correcting their values to the I-CDES scale through the ETH standards, resulting in a value of 0.325 ± 0.008‰ (95% confidence level).

Where possible, we used means and standard deviations of the calcification temperature of the samples as reported in the literature. Measurement uncertainty was often reported at the level of standard errors on the mean of multiple aliquots of the same sample. The standard deviation of Δ_{47} values on individual aliquots was estimated from repeated measurements of reference materials not involved in the calibration (e.g. ETH-4 or Carrara marble) or, if these were not reported, by back-calculating the standard deviation from the standard error through multiplication by the square root of the sample size. In absence of uncertainties on the calcification temperature, the standard deviation of the temperatures was assumed to be 1°C for samples in the low-temperature domain (<100°C) and 10°C for high-temperature samples (the 850°C heated aragonites from Müller et al., 2017).

The Guo et al. (2009) temperature dependencies were brought into the I-CDES reference frame by updating the Δ_{47} - Δ_{63} fractionation factor to 0.268‰ (following Dennis et al., 2011), applying the Δ_{47} -dependent scaling of the Δ_{47} - Δ_{63} fractionation factor of 35 ppm/‰ (cited in Guo et al., 2009 and implemented in Jautzy et al., 2020) and calibrating to the I-CDES scale using a linear calibration through the Δ_{47} values of ETH-1 and ETH-3. For this calibration, the "new" I-CDES values for ETH-1 and ETH-3 were retrieved from Bernasconi et al. (2021) and the "old" CDES25 (CDES with reference to a reaction temperature of 25°C) values were obtained by solving the polynomial functions for the formation temperatures of the ETH standards (assumed to be 20°C for ETH-3 and 600°C for ETH-1, following Meckler et al., 2014).

5. Statistical evaluation

Differences between temperatures and between specimens within the same temperature treatment were tested using a one-way ANOVA and post-hoc Tukey multiple pairwise comparisons (see **S5**). Uncertainties on Δ_{47} values and calcification temperatures on the aliquot level are propagated through all statistical procedures, and uncertainty-weighted means and 95% confidence levels of samples are used for plotting throughout the manuscript (see section 6 below). The difference in

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reproducibility between aliquots measured on our MAT253 (σ of reproducibility of IAEA-C2 of 0.046‰) and our MAT253 plus (σ of reproducibility of IAEA-C2 of 0.026‰) was considered in our error propagation by initially grouping aliguots by instrument and summarizing statistics (means and σ). The weighted mean and uncertainty of all aliquots from the same specimen or treatment group, but measured on different instruments, was then calculated by combining the statistics of groups of aliquots from the two machines and weighting the contribution of the two instrument groups by the factor $\frac{N}{\sigma^2}$, in which N is the number of aliquots in the instrument group and σ is the standard deviation representing the reproducibility within the group. For the value of σ , either the external standard deviation based on IAEA-C2 measurements on that instrument was used or the standard deviation between the Δ_{47} values of the aliguots within the group, whichever yielded the largest (and therefore most conservative) estimate of the uncertainty. This approach was based on work by Tatebe (2005) and Kirchner (2006) and the calculations are worked out in detail below (derivation) and **S5** (R script). To incorporate uncertainty on both Δ_{47} values and calcification temperatures at the aliquot level into our regressions, we apply a York regression (York, 1966) that takes into account errors on the independent variable as implemented in the "bfsl" R package by Patrick Sturm (Sturm, 2018). This linear York regression is repeated for the full dataset and for the dataset excluding high temperature (>30°C) datapoints to test the effect of these measurements on the regression. Throughout these statistical evaluations, it is assumed that uncertainties on Δ_{47} values and calcification temperatures are normally distributed.

6. Calculating weighted means and standard deviations

Suppose we have a dataset of measurements that measure the same quantity with different uncertainty. This difference in uncertainty can be taken into account when determining the mean and standard deviation of the dataset. In this document we derive equations for the weighted mean and standard deviation for a dataset of raw measurements, and for a dataset consisting pre-binned data in two or more bins.

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6.1 Dataset of individual measurements

Suppose we have a dataset consisting of several raw measurements, all with their own uncertainty. The mean and standard deviation of the dataset can be calculated by weighting the measurements, and the optimal choice of weights is based on the inverse of the measurement variance (Kirchner, 2006).

To put this in mathematical terms: we have a dataset of *n* measurements $x_1, ..., x_n$ each with known variance $\sigma_1^2, ..., \sigma_n^2$ (note that we express the variance as the square of the standard deviation σ). In order to calculate the weighted mean μ and standard deviation σ , we define weights $w_i = 1/\sigma_i^2 = \sigma_i^{-2}$ for i = 1, ..., n. Following equations (1) and (2) of Kirchner, (2006), we find:

$$\mu = \frac{\sum_{i=1}^{n} w_i \cdot x_i}{\sum_{i=1}^{n} w_i} \tag{1}$$

$$\sigma = \sqrt{\frac{n}{n-1} \cdot \frac{\sum_{i=1}^{n} w_i \cdot (x_i - \mu)^2}{\sum_{i=1}^{n} w_i}}$$
(2)

Note that in case of equal weights for all measurements (i.e. $w_i = 1$ for i = 1, ..., n), Equations (1) and (2) coincide with the well-known formulas for unweighted mean and standard deviation:

$$\mu = \frac{\sum_{i=1}^{n} 1 \cdot x_i}{\sum_{i=1}^{n} 1} = \frac{\sum_{i=1}^{n} x_i}{n}$$

$$\sigma = \sqrt{\frac{n}{n-1} \cdot \frac{\sum_{i=1}^{n} 1 \cdot (x_i - \mu)^2}{\sum_{i=1}^{n} 1}} = \sqrt{\frac{n}{n-1} \cdot \frac{\sum_{i=1}^{n} (x_i - \mu)^2}{n}} = \sqrt{\frac{\sum_{i=1}^{n} (x_i - \mu)^2}{n-1}}$$

6.2 Binned datasets

Consider now the situation where the raw measurements have been binned in two or more bins. We know the mean, standard deviation, and bin size of the bins. From this information, we can estimate the weighted mean and standard deviation of the combination of bins. In this case, we weight the measurements using the inverse of the bin variance.

6.3 Dataset of two bins

First suppose we have two bins. Let's put this in mathematical terms again. We have two bins: B_1 with mean μ_1 , standard deviation σ_1 , and size n_1 , and B_2 with mean μ_2 , standard deviation σ_2 , and size n_2 . Bin B_1 consists of (unknown) measurements $y_1, ..., y_{n_1}$, and B_2 consists of (unknown) measurements $z_1, ..., z_{n_2}$. The combination of the bins has size $n = n_1 + n_2$ and consists of (unknown) measurements $x_1, ..., x_n = y_1, ..., y_{n_1}, z_1, ..., z_{n_2}$. We want to estimate the weighted mean and standard deviation using weights $w_1 = 1/\sigma_1^2$ for the measurements of B_1 , and $w_2 = 1/\sigma_2^2$ for the measurements of B_2 . The derivations in this section were inspired by the analysis of Tatebe (2005).

For the weighted mean, we apply Equation (1) to find:

$$\mu = \frac{\sum_{i=1}^{n} w_i \cdot x_i}{\sum_{i=1}^{n} w_i}$$
Apply Equation (1)
$$= \frac{\sum_{i=1}^{n} w_1 \cdot y_i + \sum_{i=1}^{n} w_2 \cdot z_i}{\sum_{i=1}^{n} w_1 + \sum_{i=1}^{n} w_2}$$
Split sums into the two bins
$$= \frac{w_1 \cdot \sum_{i=1}^{n} y_i + w_2 \cdot \sum_{i=1}^{n} z_i}{n_1 \cdot w_1 + n_2 \cdot w_2}$$
The weights in the sums are constants, so the sums can be simplified
$$= \frac{w_1 \cdot n_1 \cdot \mu_1 + w_2 \cdot n_2 \cdot \mu_2}{n_1 \cdot w_1 + n_2 \cdot w_2}$$
For B_1 we have $\sum_{i=1}^{n_1} y_i = n_1 \cdot \mu_1$,

(3)

and a similar relation holds for B_2 .

Note that in case of equal weighting (i.e., $w_1 = w_2 = 1$), Equation (3) comes down to a weighted mean that only takes the difference in bin size into account.

The derivation for the weighted standard deviation is more elaborate. First, we apply Equation (2). Using the same line of reasoning as for the mean, we find:

$$\sigma = \sqrt{\frac{n}{n-1}} \cdot \frac{\sum_{i=1}^{n} w_i \cdot (x_i - \mu)^2}{\sum_{i=1}^{n} w_i}$$

$$= \sqrt{\frac{n}{n-1}} \cdot \frac{\sum_{i=1}^{n1} w_1 \cdot (y_i - \mu)^2 + \sum_{i=1}^{n2} w_2 \cdot (z_i - \mu)^2}{\sum_{i=1}^{n1} w_1 + \sum_{i=1}^{n2} w_2}$$

$$= \sqrt{\frac{n}{n-1}} \cdot \frac{w_1 \cdot \sum_{i=1}^{n1} (y_i - \mu)^2 + w_2 \cdot \sum_{i=1}^{n2} (z_i - \mu)^2}{n_1 \cdot w_1 + n_2 \cdot w_2}}$$
(4)

We can rewrite the terms $\sum_{i=1}^{n_1} (y_i - \mu)^2$ and $\sum_{i=1}^{n_2} (z_i - \mu)^2$ to eliminate the unknown measurements y_i and z_i from the equation. We consider the first term and note that the same line of reasoning holds for the second term. The keys to rewriting the terms are first adding $(\mu_1 - \mu_1) = 0$ to the sum, and then writing out the product of sums in separate sums:

$$\Sigma_{i=1}^{n_1} (y_i - \mu)^2 = \Sigma_{i=1}^{n_1} ((y_i - \mu_1) + (\mu_1 - \mu))^2$$

= $\Sigma_{i=1}^{n_1} (y_i - \mu_1)^2 + 2 (y_i - \mu_1) \cdot (\mu_1 - \mu) + (\mu_1 - \mu)^2$
= $\Sigma_{i=1}^{n_1} (y_i - \mu_1)^2 + \Sigma_{i=1}^{n_1} 2 (y_i - \mu_1) \cdot (\mu_1 - \mu) + \Sigma_{i=1}^{n_1} (\mu_1 - \mu)^2$ (5)

We rewrite each of the three sums in Equation (5) separately. For the first sum, we use the formula of the (unweighted) standard deviation applied to bin B_1 :

$$\sum_{i=1}^{n_1} (y_i - \mu_1)^2 \qquad = (n_1 - 1) \cdot \sigma_1^2 \tag{6}$$

For the second sum, we note that $(\mu_1 - \mu)$ is a constant, so we can take it out of the summation:

$$\sum_{i=1}^{n_1} 2 (y_i - \mu_1) \cdot (\mu_1 - \mu) = 2 \cdot (\mu_1 - \mu) \cdot \sum_{i=1}^{n_1} (y_i - \mu_1)$$

$$= 2 \cdot (\mu_1 - \mu) \cdot (\sum_{i=1}^{n_1} y_i - \sum_{i=1}^{n_1} \mu_1)$$

$$= 2 \cdot (\mu_1 - \mu) \cdot (n_1 \cdot \mu_1 - \sum_{i=1}^{n_1} \mu_1)$$

$$= 2 \cdot (\mu_1 - \mu) \cdot (n_1 \cdot \mu_1 - n_1 \cdot \mu_1)$$

$$= 0$$
(7)

For the third sum, we again use that $(\mu_1 - \mu)$ is a constant:

$$\sum_{i=1}^{n_1} (\mu_1 - \mu)^2 \qquad = n_1 \cdot (\mu_1 - \mu)^2 \tag{8}$$

Plugging Equations (6), (7) and (8) in Equation (5), we find:

$$\sum_{i=1}^{n_1} (y_i - \mu)^2 = \sum_{i=1}^{n_1} (y_i - \mu_1)^2 + \sum_{i=1}^{n_1} 2 (y_i - \mu_1) \cdot (\mu_1 - \mu) + \sum_{i=1}^{n_1} (\mu_1 - \mu)^2$$

= $(n_1 - 1) \cdot \sigma_1^2 + n_1 \cdot (\mu_1 - \mu)^2$ (9)

The same derivation yields:

$$\sum_{i=1}^{n_2} (z_i - \mu)^2 = (n_2 - 1) \cdot \sigma_2^2 + n_2 \cdot (\mu_2 - \mu)^2$$
(10)

Plugging Equations (9) and (10) in Equation (4), the formula for the weighted standard deviation becomes:

$$\sigma = \sqrt{\frac{n}{n-1} \cdot \frac{w_1 \cdot (n_1-1) \cdot \sigma_1^2 + w_1 \cdot n_1 \cdot (\mu_1-\mu)^2 + w_2 \cdot (n_2-1) \cdot \sigma_2^2 + w_2 \cdot n_2 \cdot (\mu_2-\mu)^2}{n_1 \cdot w_1 + n_2 \cdot w_2}}$$
(11)

This equation consists of all known values, making use of Equation (3) for the mean μ . Note that in case of equal weighting (i.e., $w_1 = w_2 = 1$), Equation (11) comes down to an unweighted

standard deviation of all measurements in the two bins in relation to the mean of Equation (3). Equation (11) can also be written in terms of only the characteristics of the two bins, by plugging Equation (3) in Equation (11) and rewriting the outcome. Consider the term $(\mu_1 - \mu)^2$. We can write:

$$\begin{aligned} (\mu_1 - \mu)^2 &= \left(\mu_1 - \frac{w_1 \cdot n_1 \cdot \mu_1 + w_2 \cdot n_2 \cdot \mu_2}{n_1 \cdot w_1 + n_2 \cdot w_2}\right)^2 \\ &= \left(\frac{n_1 \cdot w_1 + n_2 \cdot w_2}{n_1 \cdot w_1 + n_2 \cdot w_2} \cdot \mu_1 - \frac{n_1 \cdot w_1}{n_1 \cdot w_1 + n_2 \cdot w_2} \cdot \mu_1 - \frac{n_2 \cdot w_2}{n_1 \cdot w_1 + n_2 \cdot w_2} \cdot \mu_2\right)^2 \\ &= \left(\frac{n_2 \cdot w_2}{n_1 \cdot w_1 + n_2 \cdot w_2} \cdot \mu_1 - \frac{n_2 \cdot w_2}{n_1 \cdot w_1 + n_2 \cdot w_2} \cdot \mu_2\right)^2 \\ &= \left(\frac{n_2 \cdot w_2}{n_1 \cdot w_1 + n_2 \cdot w_2} \cdot (\mu_1 - \mu_2)\right)^2 \\ &= \frac{n_2^2 \cdot w_2^2}{(n_1 \cdot w_1 + n_2 \cdot w_2)^2} (\mu_1 - \mu_2)^2 \end{aligned}$$

In the same way we can rewrite:

$$(\mu_1 - \mu)^2 = \frac{n_1^2 \cdot w_1^2}{(n_1 \cdot w_1 + n_2 \cdot w_2)^2} (\mu_2 - \mu_1)^2$$

Now, we can use the fact that $(\mu_2 - \mu_1)^2 = (\mu_1 - \mu_2)^2$ to simplify part of Equation (11):

$$w_{1} \cdot n_{1} \cdot (\mu_{1} - \mu)^{2} + w_{2} \cdot n_{2} \cdot (\mu_{2} - \mu)^{2}$$

$$= w_{1} \cdot n_{1} \cdot \frac{n_{2}^{2} \cdot w_{2}^{2}}{(n_{1} \cdot w_{1} + n_{2} \cdot w_{2})^{2}} (\mu_{1} - \mu_{2})^{2} + w_{2} \cdot n_{2} \cdot \frac{n_{1}^{2} \cdot w_{1}^{2}}{(n_{1} \cdot w_{1} + n_{2} \cdot w_{2})^{2}} (\mu_{2} - \mu_{1})^{2}$$

$$= \frac{w_{1} \cdot n_{1} \cdot n_{2}^{2} \cdot w_{2}^{2} + w_{2} \cdot n_{2} \cdot n_{1}^{2} \cdot w_{1}^{2}}{(n_{1} \cdot w_{1} + n_{2} \cdot w_{2})^{2}} (\mu_{1} - \mu_{2})^{2}$$

$$= \frac{w_{1} \cdot n_{1} \cdot w_{2} \cdot n_{2} \cdot (n_{1} \cdot w_{1} + n_{2} \cdot w_{2})}{(n_{1} \cdot w_{1} + n_{2} \cdot w_{2})^{2}} (\mu_{1} - \mu_{2})^{2}$$

$$= \frac{w_{1} \cdot n_{1} \cdot w_{2} \cdot n_{2}}{n_{1} \cdot w_{1} + n_{2} \cdot w_{2}} (\mu_{1} - \mu_{2})^{2}$$
(12)

Plugging Equation (12) into Equation (11), we find:

$$\sigma = \sqrt{\frac{n}{n-1} \cdot \frac{w_1 \cdot (n_1-1) \cdot \sigma_1^2 + w_1 \cdot n_1 \cdot (\mu_1-\mu)^2 + w_2 \cdot (n_2-1) \cdot \sigma_2^2 + w_2 \cdot n_2 \cdot (\mu_2-\mu)^2}{n_1 \cdot w_1 + n_2 \cdot w_2}}$$
$$= \sqrt{\frac{n_1 + n_2}{(n_1 + n_2 - 1) \cdot (n_1 \cdot w_1 + n_2 \cdot w_2)}} \cdot \left(w_1 \cdot (n_1 - 1) \cdot \sigma_1^2 + w_2 \cdot (n_2 - 1) \cdot \sigma_2^2 + \frac{w_1 \cdot n_1 \cdot w_2 \cdot n_2}{n_1 \cdot w_1 + n_2 \cdot w_2}} (\mu_1 - \mu_2)^2\right)$$
(13)

Equation (3) and (13) give the weighted mean and standard deviation in terms of the characteristics of the two bins. Filling in the weights $w_i = \sigma_i^{-2}$, we arrive at

$$\mu = \frac{\sigma_1^{-2} \cdot n_1 \cdot \mu_1 + \sigma_2^{-2} \cdot n_2 \cdot \mu_2}{\sigma_1^{-2} \cdot n_1 + \sigma_1^{-2} \cdot n_2}$$
(14)

$$\sigma = \sqrt{\frac{n_1 + n_2}{(n_1 + n_2 - 1) \cdot (n_1 \cdot \sigma_1^{-2} + n_2 \cdot \sigma_2^{-2})} \cdot \left(\sigma_1^{-2} \cdot (n_1 - 1) \cdot \sigma_1^2 + \sigma_2^{-2} \cdot (n_2 - 1) \cdot \sigma_2^2 + \frac{\sigma_1^{-2} \cdot n_1 \cdot \sigma_2^{-2} \cdot n_2}{n_1 \cdot \sigma_1^{-2} + n_2 \cdot \sigma_2^{-2}} (\mu_1 - \mu_2)^2\right)}$$

$$= \sqrt{\frac{n_1 + n_2}{(n_1 + n_2 - 1) \cdot (n_1 \cdot \sigma_1^{-2} + n_2 \cdot \sigma_2^{-2})} \cdot \left(n_1 + n_2 - 2 + \frac{\sigma_1^{-2} \cdot n_1 \cdot \sigma_2^{-2} \cdot n_2}{n_1 \cdot \sigma_1^{-2} + n_2 \cdot \sigma_2^{-2}} (\mu_1 - \mu_2)^2\right)}$$
(15)

6.4 Dataset of more than two bins

Equation (3) gives the formula for a weighted mean of two bins. It can straightforwardly be generalized to hold for any number of bins B_j with means μ_j , standard deviations σ_j , sizes n_j , and weights $w_j = 1/\sigma_j^2$:

$$\mu = \frac{\sum w_j \cdot n_j \cdot \mu_j}{\sum w_j \cdot n_j} = \frac{\sum \sigma_j^{-2} \cdot n_j \cdot \mu_j}{\sum \sigma_j^{-2} \cdot n_j}$$
(16)

In the same way, Equation (11) can be generalized to hold for any number of bins B_j , using Equation (16) for the mean μ :

$$\sigma = \sqrt{\frac{\sum n_j}{(\sum n_j) - 1}} \cdot \frac{\sum w_j \cdot (n_j - 1) \cdot \sigma_j^2 + w_j \cdot n_j \cdot (\mu_j - \mu)^2}{\sum w_j \cdot n_j}$$
$$= \sqrt{\frac{\sum n_j}{(\sum n_j) - 1}} \cdot \frac{\sum \sigma_j^{-2} \cdot (n_j - 1) \cdot \sigma_j^2 + \sigma_j^{-2} \cdot n_j \cdot (\mu_j - \mu)^2}{\sum \sigma_j^{-2} \cdot n_j}$$

$$=\sqrt{\frac{\sum n_j}{(\sum n_j)-1} \cdot \frac{\sum (n_j-1) + \sigma_j^{-2} \cdot n_j \cdot (\mu_j-\mu)^2}{\sum \sigma_j^{-2} \cdot n_j}}$$
(17)

Note that Equations (13) and (15) for the standard deviation cannot readily be generalized to hold for any number of bins. In case of more than two bins, the weighted mean is first calculated using Equation (16), and then the standard deviation can be calculated using this mean and Equation (17).

References

Kirchner, J. (2006). *Data Analysis Toolkit #12: Weighted averages and their uncertainties*. http://seismo.berkeley.edu/~kirchner/Toolkits/Toolkit_12.pdf

Tatebe, K. (2005). *Combining Multiple Averaged Data Points And Their Errors*. https://docplayer.net/33088897-Combining-multiple-averaged-data-points-and-their-errors.html

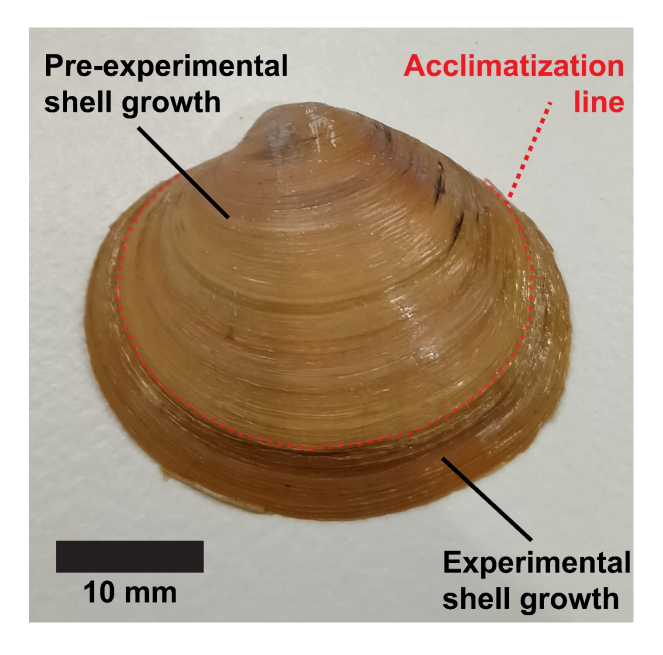


Figure S1. Showing a shell of cultured Arctica islandica with a clearly visible growth mark highlighting transplantation into the culturing conditions.

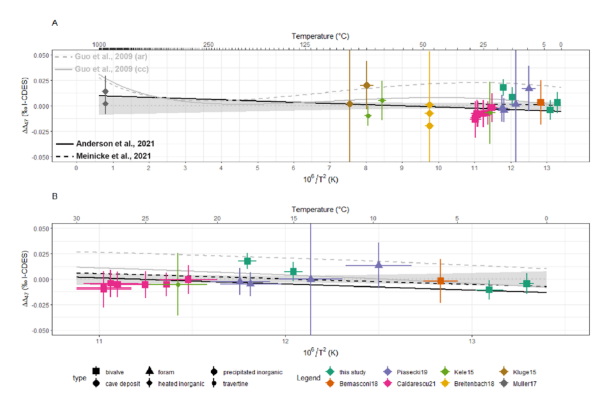


Figure S2. Residuals of aragonite \Delta47-temperature relationships. Showing the difference in clumped isotope value (Δ 47) between aragonite Δ 47 data and the two Δ 47-temperature regressions through the aragonite data compilation (see Fig. 2): A) Residuals relative to the York regression through the full dataset. B) Residuals relative to the York regression through only the low-temperature data (<30°C). Results are colored based on the study from which they originated and symbols code for the type of aragonite that was measured (see legend in A, following Fig. 2). Vertical and horizontal bars on symbols indicate uncertainty on Δ 47 and formation temperature at the 95% confidence level. Note that some vertical error bars are cropped by the restricted extent of the vertical axis. The solid black line shows the unified clumped isotope calibration by Anderson et al. (2021) while the dashed black line represents the temperature relationship by Meinicke et al. (2020; 2021; only plotted for temperatures below 100°C). Grey solid and dashed lines represent, respectively, the theoretical calcite ("cc") and aragonite ("ar") temperature dependencies from Guo et al. (2009; projected on the I-CDES scale, see section 2.5 in the main text).

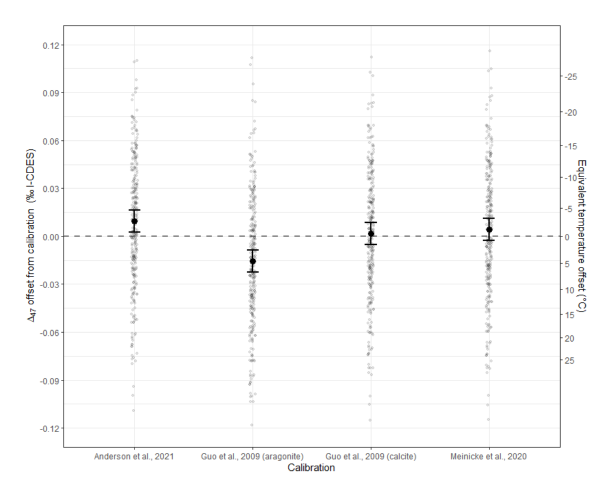


Figure S3. Zoomed out version of Figure 3 in the main text showing residuals of all *A. islandica* aliquots used in this study relative to the four previous clumped isotope temperature calibrations.

Sample	Temperature range $\pm 1\sigma$	Number of Δ_{47}	Reference		
type		aliquots			
Precipitated aragonite	80 ± 2°C – 91 ± 0.5°C	11	Kluge et al., 2015		
Heated aragonite	850 ±10°C	35	Müller et al., 2017		
Cave deposit	47 ± 1°C	16	Breitenbach et al., 2018		
Travertine	22.7 ± 1°C – 79.2 ± 1°C	78	Kele et al., 2015		
Foraminifera	9.7 ± 1°C – 18.5 ± 1°C	63	Piasecki et al., 2019		
Mollusk	6 ± 0.5°C	12	Bernasconi et al., 2018		
Mollusk	22 ± 1°C – 28 ± 1°C	165	Caldarescu et al., 2021		
Mollusk	1.1 ± 0.2°C – 18 ± 0.3°C	278	This study		
TOTAL		8			

Overview of datasets included in aragonite compilation

Table S1. Overview of aragonite data from previous studies sued for the compilation in this study.