Refining trace metal temperature proxies in cold-water scleractinian and stylasterid corals

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The Li/Mg, Sr/Ca and oxygen isotopic (δ¹⁸O) compositions of many marine biogenic carbonates are sensitive to seawater temperature. Corals, as cosmopolitan marine taxa with carbonate skeletons that can be precisely dated, represent ideal hosts for these geochemical proxies. However, efforts to calibrate and refine temperature proxies in cold-water corals (<20°C) remain limited. Here we present skeletal Li/Mg, Sr/Ca, δ¹⁸O and carbon isotope (δ¹³C) data from live-collected specimens of aragonitic scleractinian corals (Balanophyllia, Caryophyllia, Desmophyllum, Enallagma, Flabellum, Lophelia, and Vagahuelle), both aragonitic and high-Mg calcitic stylasterid genera (Stylaster and Errina), and shallow-water high-Mg calcite crustose coralline algae (Ulothryphnum, Hydrolithon, and Neogoniolithon). We interpret these data in conjunction with results from previously explored taxa including aragonitic zooxanthellate scleractinia and foraminifera, and high-Mg calcite octocorals. We show that Li/Mg ratios covary most strongly with seawater temperature, both for aragonitic and high-Mg calcitic taxa, making for reliable and universal seawater temperature proxies. Combining all of our biogenic aragonitic Li/Mg data with previous calibration efforts we report a refined relationship between temperature: Li/Mg_{kall Aragonite} = 5.42 \exp(-0.050 \times T(°C)) (R² = 0.97). This calibration now permits paleo-temperature reconstruction to better than ±3.4°C (95% prediction intervals) across biogenic aragonites, regardless of taxon, from 0 to 30°C. For taxa in this study, aragonitic stylasterid Li/Mg offers the most robust temperature proxy (Li/Mg_{Stylasterid (Ar)} = 5.64 \exp(-0.046 \times T(°C)) (R² = 0.95)) with a reproducibility of ±2.3°C. For the first time, we show that high-Mg calcites have a similar exponential relationship with temperature, but with a lower intercept value (Li/Mg = 0.63 \exp(-0.050 \times T(°C)) (R² = 0.92)). This calibration opens the possibility of temperature reconstruction using high-Mg calcite corals and coralline algae. The commonality in the relationship between Li/Mg and temperature transcends phylogeny and suggests a similar abiogenic trace metal incorporation mechanism.

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1. Introduction

Understanding the linkages between ocean circulation, carbon cycling, and rapid climate change on geologic and histori-

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ocal timescales remains a key goal of paleoceanographic research (Broecker, 1982). Paleo-seawater temperature is a hydrographic variable vital to our interpretation of these ocean-climate interactions (Thiagarajan et al., 2014). Much attention has therefore been given to the development of geochemical paleo-temperature proxies in marine carbonates to extend temperature timescales beyond the range of instrumental records (Anand et al., 2003; Case et al., 2010; Spooner et al., 2016). Deep-sea sediment cores remain a popular archive for obtaining ancient biogenic carbonate material (e.g. (Poggemann et al., 2017; Roberts et al., 2016)). However, impaired age control often limits use of these records for understanding the leads and lags of the climate system on millennial timescales or shorter. Therefore, robust temperature proxies from archives with precise age control are much needed.

Corals occupy a wide geographic and bathymetric range (Roberts et al., 2006). They can be accurately and precisely dated by radiometric methods (Cheng et al., 2000) or annual growth band counting (Knutson et al., 1972) making them exciting targets for paleoceanographic research. Selected trace metal and stable isotopic ratios in marine carbonates have been shown to vary as a function of seawater temperature including: Sr/Ca (Beck et al., 1992; Cohen et al., 2001), Li/Ca (Hall and Chan, 2004; Marriott et al., 2004), Mg/Ca (Anand et al., 2003; Case et al., 2010), and δ18O (Cobb et al., 2003; Gagan et al., 2012). Such temperature proxies have been used to reconstruct temperature on sub-annual to millennial timescales (Cohen et al., 2001; McCulloch et al., 1999).

Scleractinian corals (Subclass Hexacorallia; Order Scleractinia) calcify from an extracellular calcifying fluid (ECF) that is semi-restricted from ambient seawater (Sinclair and Risk, 2006). The chemistry of this ECF is modified via the Ca-ATPase pump, raising internal pH and saturation state to promote precipitation of skeletal calcium carbonate (A-Horani et al., 2003). Such changes to internal seawater have the potential to impact the chemistry independently of seawater temperature. For instance, the δ13C and δ18O composition of the coral skeleton is impacted by kinetic isotope effects during hydration and hydroxylisation of CO2 (Guo, 2020; McConnaughey, 1989), pH dependent dissolved inorganic carbon speciation (i.e. HCO3 to 18O-depleted CO2−; Adkins et al., 2003), and carbonate anhydride activity (Chen et al., 2018). Skeletal trace metal compositions can be similarly modified by the balance between carbonate precipitation, direct seawater leakage, and ion pumping into the ECF. Through Rayleigh fractionation and/or growth entrainment, ions can become either depleted or enriched in the ECF (relative to Ca) depending on their partition coefficient (D) into the carbonate lattice as calcification progresses (e.g. DSr = Sr/Ca(calcite)/Sr/Ca(seawater) > 1; (Gaetani et al., 2011; Gagan et al., 2007; Watson, 2004)). Trace element ratios may therefore be impacted by the calcification process, driving variations between coral species, colonies, and even individuals, thus hindering their use as temperature monitors (Chen et al., 2018; Gagan et al., 2012; Gagan et al., 2007). Paired coral U/Ca ratios have been used to correct Sr/Ca values for growth rate effects, yielding a more robust relationship with seawater temperature (DeCarlo et al., 2016). However, this Sr-U approach is only applicable to large, growth-handled, shallow-water corals, from which seasonally resolved records can be obtained to characterise the Sr-U relationship for each individual (Ross et al., 2019). Alternative approaches are therefore required for developing robust temperature proxies in smaller cold-water coral taxa.

Coral Mg/Ca and Li/Ca ratios alone have complex, but contrasting, relationships with calcification temperature. Studies show that skeletal Mg/Ca has a positive correlation with temperature (Hathorne et al., 2013a), while others report this correlation to be poor (Case et al., 2010; Montagna et al., 2014). Conversely, coral wall Li/Ca has a weak negative correlation with temperature (Case et al., 2010). With similar partition coefficients (both DLi and DMg ≈ 1) however, Li/Ca and Mg/Ca are similarly impacted by calcification processes (Marchitto et al., 2018). Thus the ratio of coral Li/Mg has great promise; preserving the temperature dependency of these elements while minimising growth-rate effects (Case et al., 2010). A universal Li/Mg relationship to temperature appears robust across published data for both zooxanthellate and azooxanthellate scleractinia (Andersson et al., 2020; Case et al., 2010; Cuny-Guirriec et al., 2019; Fowell et al., 2016; Hathorne et al., 2013a; Montagna et al., 2014; Raddatz et al., 2013; Ross et al., 2019) and also aragonitic foraminifera (Hoeglundina elegans; Bryan and Marchitto, 2008; Hall and Chan, 2004; Marchitto et al., 2018) (Fig. 1A). This recalculated relationship is described by the equation,

\[
\text{Li/Mg} = \frac{5.27[±0.06]e^{-0.049[±0.001]×T(°C)}}{(R^2 = 0.96; [±SE])}
\]

and is based on Li/Mg data compiled from the literature that we now adjust for interlaboratory analytical offsets (see Methods and Supplementary Information). More than three quarters of the data characterising this Li/Mg relationship to seawater temperature however come from shallow-water coral specimens harbouring zooxanthellae (Fowell et al., 2016; Hathorne et al., 2013a; Montagna et al., 2014; Ross et al., 2019), with few observations obtained from azooxanthellate deep-water taxa (Case et al., 2010; Cuny-Guirriec et al., 2019; Montagna et al., 2014; Raddatz et al., 2013). Fewer still of these samples are from locations with well-characterised temperature (Case et al., 2010). Full characterisation of this relationship across multiple cosmopolitan cold-water coral taxa (<20°C) from localities with well-documented hydrography is therefore much needed. Cold-water corals are known to possess large heterogeneities in microstructural chemistry, with centres of calcification (COCs) being particularly enriched in Li and Mg and depleted δ13C and δ18O (Adkins et al., 2003; Anagnostou et al., 2011; Chen et al., 2018; Correa et al., 2010; Gagnon et al., 2007; Stewart et al., 2016). It is important therefore to ensure that sampling of such corals is reproducible and not unduly biased by microstructures.

A second important, but poorly studied, deep-water coral family is the Stylasteridae (Class Hydrozoa: Order Anthoathecata; Fig. 2). These Hydrozoan corals differ significantly from scleractinian corals in their ability to precipitate calcium carbonate skeletons of either aragonite, high-Mg calcite, or in some cases, both polymorphs (Cairns, 2011). Predicting the response of marine calcifiers to future anthropogenic change (e.g. ocean acidification and/or thermal stress) is critical and requires insight into the biocalcification processes of both aragonitic and high-Mg calcitic organisms from skeletal geochemistry (Donald et al., 2017). A recent study shows that stylasterid coral skeletal δ18O remains relatively close to equilibrium with seawater and is thus minimally impacted by growth-rate effects and well-correlated with seawater temperature (Samperiz et al., 2020). Yet, there remains no study to date characterising trace metal compositions of either aragonitic or calcitic stylasterid corals, hence their paleoceanographic proxy potential and biocalcification mechanisms remain poorly understood.

The temperature dependency of trace metals in high-Mg calcites is less well-constrained than it is in aragonitic calcifiers. Exploratory work into high-Mg calcite crustose coralline algae (CCA) grown in laboratory culture (Clathromorphum compactum; 6 to 13°C) suggest that Li/Mg in high-Mg calcites may also be related to temperature (Fig. 1A; Anagnostou et al., 2019). However, this result contrasts with Li/Mg values measured in wild-collected high-Mg calcitic coral *Corallium rubrum* (Octocorallia) which are suggested to have little coherent relationship with seawater temperature (Fig. 1A; Chaabane et al., 2019). Without high-Mg calcite
samples from a greater range of temperatures it remains unclear if the Li/Mg relationship to temperature is linear (Anagnostou et al., 2019), exponential (e.g. scleractinia; Montagna et al., 2014), or indeed absent (Chaabane et al., 2019) in this group of organisms.

To assess temperature proxies within different marine carbonate mineralogy, we present skeletal Li/Mg, Sr/Ca, and $\delta^{18}$O data obtained from modern cold-water coral calyxes from seawater temperatures ranging from 0 to 16°C. These specimens comprise 7 genera of common deep-water scleractinia as well as two genera of stylasterid corals (Stylaster and Errina) that represent all three modes of stylasterid skeletal mineralogy: (i) solely aragonite, (ii) both aragonite and high-Mg calcite, and (iii) solely high-Mg calcite (Cairns and Macintyre, 1992; Samperiz et al., 2020). We present data from the same stylasterid specimens for which $\delta^{18}$O has been previously characterised (Samperiz et al., 2020) for comparison with our trace metal data. We complement these coral sample sets, with CCA collected from tropical surface waters, expanding the range of high-Mg calcite skeletons to include warmer water specimens ($\sim$27°C). We use skeletal $\delta^{13}$C to assess biocalcification and microstructural variability in replicate samples. We critically assess the reliability of each proxy and its applicability for reconstructing paleo-seawater temperature.

2. Methodology

2.1. Samples and hydrographic data

More than 100 coral samples were collected from the Labrador Sea (RRS Discovery DY081), Northeast Atlantic (RRS James Cook JC136), Equatorial Atlantic (RRS James Cook JC094), Drake Pas-
Annelia, by We Fjord NOAA JR15005), Ross Sage Ocean in Fig. 3. Location 4

(Octocorallia Aplysina, Archipelago includes 14 Hydrolithon samples from (R/V Alucia and (Cruise AL1508) Mexico retrieved (Fig. 3). The compilation of hydrographic data were considerably more proximal than this (-5 km from coral sites and within 5 m depth). Where paired shipboard data were unavailable temperature estimates were taken from nearby sites in the GLODAP database (https://www.glodap.info/; Key et al., 2004). With the exception of the Galápagos site, these measurements were all within 280 km and 90 m water depth of coral sampling sites. First choice (most proximal) seawater bottle temperature estimates are typically within 0.3 °C of GLODAP gridded temperature estimates (all within 1.9 °C; excluding Comau Fjord sites where GLODAP gridded data are unavailable). To further assess uncertainty on temperature estimates caused by natural variability and distance between coral and hydrographic stations, two additional nearby bottle seawater temperature measurements were selected from the compiled GLODAP and research cruise databases (Supplementary Information). For the majority of sites these temperatures


Fig. 2. Phylogeny and skeletal mineralogy (Aragonite, high-Mg calcite or both polymorphs) of carbonates in this study.

Fig. 3. Location of modern coral (circles) and coralline algae (square) samples used in this study with respective cruise identifiers (white text). Base map shows World Ocean Atlas gridded seawater temperature at 1000 m (Locarnini et al., 2013).
were within ±0.5 °C (2σ) of each other with a maximum variation of ±3.3 °C for one site in CINMS. In such cases where temperature was more variable, the most proximal bottle seawater temperature measurement is still preferred over averaging all three temperature estimates.

Surface water temperature measurements are typically subject to stronger seasonal bias than deep waters. Hence, annually averaged, 0.25° gridded, World Ocean Atlas temperature data (years 1955 to 2012, Locarnini et al., 2013) were preferred for Caribbean CCA samples over temperature data taken at the time of collection because these are not representative of the annual average (Crook et al., 2012).

Similar to our compilation of sub-surface temperature, we estimate δ18O of seawater for each site using co-located measurements taken by remotely operated vehicle or CTD during the coral sampling expedition (JC094 and NBP1103); (Samperiz et al., 2020). Where such data are unavailable, δ18O of seawater was taken from nearby sites in the NASA Global Seawater Oxygen-18 Database (Schmidt et al., 1999). No nearby seawater δ18O data exist for Cama Fjord in any database, hence we assume values of mean ocean seawater (0%) for these sites (Supplementary Information).

2.2. Coral sample preparation

Coral samples for this study were air dried following collection. The majority of organic matter was removed through physical scraping and treatment in dilute NaClO for 12 h. Whole S1 septa and attached theca were taken from cup corals (Balanophyllia, Caryophyllia, Desmophyllia, Flabellum, and Vaughanella) while whole calyces were taken from branching Endallopamsia and Lophelia samples using a rotating cutting tool. Cross sectional discs (~1.8 mm thickness) were cut from the central trunk (or widest branch) of stylasterid samples. The δ18O composition of growing tips of stylasterid branches has been shown to be further from seawater equilibrium than bulk samples (Samperiz et al., 2020), hence apical tips were avoided. Similarly, microstructures within deep-sea corals such as COCs are known to exhibit highly contrasting trace metal chemistry (Gagnon et al., 2007). Microsampling techniques (e.g. micromilling) can be time consuming and offer no guarantee that COCs are completely excluded, thus are often unfeasible when generating paleo-records (Stewart et al., 2016). As an alternative, sufficiently large samples were cut (~1 g) that would diminish the influence of single microstructural components. While this sampling style may incorporate sample material grown across multiple decades, this approach is simple to follow and targets a representative average of bulk coral chemistry. To help confirm this, corals were sampled and analyzed in duplicate to characterise intra-sample variability. Initial results revealed greater trace metal variability for bulk-sampled Lophelia (whole calyces) compared to other taxa (Section 3.2). For this reason, calyces from 13 out of the 16 Lophelia specimens were resampled, however this time, septa were discarded by dentist drill leaving just the theca wall. In general, Lophelia theca walls contain proportionally fewer COCs than septa (Jurikova et al., 2019) and have been targeted in previous studies (e.g. Montagna et al., 2014) without the need for time consuming micromilling.

Coral and CCA fragments were finely crushed using a pestle and mortar before 5 to 10 mg of the homogenous powder was taken. While we were not sample limited in this instance, note that analysis could be performed on less than 1 mg of powder if required. A warm 1% H2O2 (80 °C; buffered in NH4OH) oxidative cleaning and weak acid polish (0.0005 M HNO3) were applied to the powders (Boyle, 1981; Rae et al., 2011) before dissolution in distilled 0.5 M HNO3.

2.3. Analytical techniques

Trace metal analyses were carried out at the University of Bristol using a Thermo Element2 ICP-MS, at the University of St Andrews using an Agilent 7500 ICP-MS, or at the National Institute of Standards and Technology (NIST; Hollings Marine Lab) using a Thermo ElementXR ICP-MS. An aliquot of the dissolved sample was analyzed using well-characterised, matrix-matched, synthetic standard solutions to yield Li/Ca, Mg/Ca, and Sr/Ca ratios. The same primary calibrant (bracketing standard) was used at both University of Bristol and NIST. Repeat analysis of NIST RM 8301c (n = 35) and JCp-1 (uncleaned) carbonate reference materials yielded analytical precision of ±2% (1σ) for these elemental ratios (Table 1).

Systematic analytical offsets between the labs were quantified and corrected using NIST RM 8301c results to bring values into line with interlaboratory comparison results reported in Hathorne et al. (2013b) and Stewart et al. (2015). To this end, Li/Ca, Mg/Ca and Sr/Ca ratios of samples measured at Bristol and NIST are corrected by a factor of 1.01, 0.91, and 0.97 respectively, while St Andrews values were corrected by a factor of 1.05, 105, and 1.06 respectively. A similar interlaboratory correction exercise was applied to published data according to reported JCp-1 results (Table 1; compiled data in Supplementary information).

Stable oxygen isotope analysis was performed at the National Environmental Isotope Facility based at the British Geological Survey. Approximately 60 to 100 μg of carbonate powder were analyzed using an IsoPrime dual inlet mass spectrometer plus MultiQuad device. Samples were digested in anhydrous H3PO4 at 90 °C. The CO2 released was collected for 15 minutes, cryogenically cleaned and delivered to the mass spectrometer. δ18O is reported in per mil relative to VPDB using a within-run laboratory standard calibrated against NBS-19. A calcite-aragonite and aragonite-calcite fractionation factor (1.00855), was applied to the respective gas-values along with the Craig correction factor to account for δ13C. Average reproducibility of the standard calcite (KCM) was 0.05‰ for δ18O. Note that Lophelia theca samples were not analysed for stable isotopic composition.

3. Results and discussion

3.1. Stylasterid skeletal chemistry

We compare trace metal values measured in bulk aragonitic scleractinia and high-Mg calcite CCA to our (i) solely aragonitic, (ii) both aragonitic and calcitic, and (iii) solely calcitic stylasterids (Fig. 4). Given the range of hydrographic conditions from which these sample sets were taken, we highlight the broadscale geochemical differences between the groups.

We find mineralogy to be a key factor controlling trace metal composition. Aragonitic stylasterids are low in both Li/Ca (~12 μmol/mol) and Mg/Ca (~2.9 mmol/mol) and are therefore similar to their aragonitic counterparts, scleractinia (Fig. 4). However, stylasterids composed of high-Mg calcite have considerably higher Li/Ca (~55 μmol/mol) and Mg/Ca (~82 mmol/mol) ratios that are more similar to other high-Mg calcites such as CCA. Resultant Li/Mg ratios of aragonitic and high-Mg calcite stylasterids are accordingly similar to scleractinia (~4.7 mmol/mol) and CCA (~1 mmol/mol) respectively. Despite only representing a narrow temperature range (0.1 °C), we find that the stylasterid E. gracilis, that are composed of a mixture of aragonite and high-Mg calcite polymorphs, exhibit a wide range of Li/Mg values (0.9 mmol/mol) that lie between these high aragonitic and low calcitic endmembers (Fig. 4).

Aragonitic stylasterids are only slightly higher in Sr/Ca (~11 mmol/mol) compared to scleractinia, whereas these ratios are considerably lower in calcitic stylasterids (~2 mmol/mol) and again
more similar to high-Mg calcitic CCA (Fig. 4). In general, therefore, trace metal compositions in aragonitic stylasterids and calcitic stylasterids are most similar to those of the organisms with similar mineralogy (i.e. scleractinia and CCA respectively), with elemental chemistry of mixed-mineralogy stylasterids falling on a mixing line between these two endmembers.

When our scleractinnian coral stable isotope data are compared to that of stylasterids (Samperiez et al., 2020), the mineralogical dependency is less clear. Aragonitic stylasterid δ⁸⁷⁰⁸ and δ¹³C values are slightly higher, but considerably less variable than aragonitic scleractinia. Furthermore, average high-Mg calcite stylasterid δ¹⁸O is largely indistinguishable from aragonitic Stylasteridae or Scleractinia (Fig. 4).

3.2. Trace metal and stable isotope heterogeneity of corals

It is important to consider the impact of non-uniform internal skeletal chemistry on coral temperature proxies. In all but three specimens, two solid pieces of each coral were analysed. We assess the chemical heterogeneity within each coral specimen by calculating the difference between replicate δ¹⁸O, Sr/Ca, and Li/Mg analyses. In Table 2 we report the median of these differences for each coral taxonomic group.

The mixed-mineralogy stylasterid species E. gracilis (n = 4) showed large median difference for both Li/Mg and Sr/Ca replicates (0.5 and 0.4 mmol/mol respectively). This result is unsurprising given the mixture of aragonitic and calcitic polymorphs that makes up the skeleton of this species. This large range in trace metal composition between replicates is in contrast to the pure high-Mg calcite (E. lateroflora; n = 2) and pure aragonite (E. antarctica, E. boschmai, E. altissipa, Stylaster) stylasterids that have median differences of <0.1 mmol/mol for both of these elemental ratios (i.e. within analytical uncertainty).

The majority of aragonitic scleractinia genera we measured also replicate well (median Sr/Ca and Li/Mg differences of <0.3 and <0.12 mmol/mol respectively; Table 2). This small intra-specimen range in Li/Mg would result in a reconstructed temperature range of less than 0.6°C. However, we find Li/Mg replicates of Balanophyllia and Lophelia vary by more than 0.2 mmol/mol resulting in estimated temperature discrepancies of >1°C. We are unable to assess the significance of poor replication in Balanophyllia with the limited number of specimens available. However, multiple Lophelia specimens replicated poorly for both trace metal compositions and δ¹⁸O (>0.4%), with Li/Mg replicates in one specimen varying by 4.34 mmol/mol (CINMLOp-004, replicate 1 Li/Mg = 7.17 mmol/mol, i.e. higher than all other Li/Mg measurements). While outliers can occur for analytical reasons (particularly for the more challenging Li measurement), numerous studies have suggested that Lophelia are particularly chemically heterogeneous between micro-structures (Jurikova et al., 2019; Montagna et al., 2014; Raddatz et al., 2013). Our results therefore suggest that chemically heterogeneous taxa such as Lophelia should be treated with caution when sampled in bulk for paleo-temperature proxy reconstruction (discussed further in Section 3.3.1).

Scleractinian coral δ¹³C and δ¹⁸O have been observed to positively covary, due to a combination of kinetic isotope effects (McConnaughey, 1989), pH dependent dissolved inorganic carbon speciation (Adkins et al., 2003), and carbonic anhydrase activity (Chen et al., 2018), which can shift coral stable isotope ratios towards values lower than those expected from seawater equilibrium. Biocalcification models suggest that carbon enters the coral ECF via a seawater leak ((δ¹³C ~ +1‰) and from cell-derived CO₂ (δ¹³C ~ −8‰). However, the broadly linear δ¹⁸O vs δ¹³C relationship is predicted to plateau at a δ¹³C value of approximately −8‰, at the point where all of the carbon in the ECF is cell derived (Fig. 5A;
Median earistry suggests contrasts (average in specimens significantly are (Chen al., 2018). These extremely light stable isotope signatures are typical of rapidly acrating coral COCs (Chen al., 2018).

Our bulk scleractinian coral data fit the predicted general linear trend between $\delta^{18}O$ and $\delta^{13}C$ (Chen al., 2018), with results significantly lower than equilibrium values for the vast majority of specimens (Fig. 5A; average $\delta^{13}C = -3.8\%e$; $\delta^{18}O = +1.0\%e$). This result is consistent with strong modification of carbonate chemistry within the ECF of scleractinian corals. Bulk calyx *Lophelia* $\delta^{13}C$ in particular is at the low end of the range in stable isotope values (average $\delta^{13}C = -6.5\%e$), with many replicates close to the predicted minimum $\delta^{13}C$ value of $-8\%e$. This suggests that close to 100% of the carbon in these specimens is cell derived. This result contrasts with steady-state geochemical modelling (DeCarlo al., 2015) and isotope labelling experiments (Furla al., 2000) that suggest typically only 75% of precipitated carbon in other scleractinian groups is biologically sourced. These *Lophelia* samples with extremely low $\delta^{13}C$ values also have particularly high Mg/Ca ratios (Fig. 5B; >3 mmol/mol). Together, these results suggest that bulk calyx *Lophelia* incorporate material precipitated from an ECF that is more restricted from seawater resulting microstructural chemistry that is variable and highly modified from equilibrium. This is in stark contrast to $\delta^{18}O$ and $\delta^{13}C$ values in aragonitic Stylasterid corals, which occupy a narrow range (Fig. 4) and are generally higher than those of Scleractinia, thus plotting closer to seawater equilibrium (Fig. 5A; Samperiz al., 2020).

3.3. Temperature proxies

3.3.1. $\delta^{18}O$, Sr/Ca and Li/Mg in scleractinia and aragonitic stylasterids

In light of disparity between replicate samples of *Lophelia* we first compare results from bulk calyces to theca-only sub-samples in this genus. We illustrate this by plotting *Lophelia* Li/Ca, Mg/Ca,
Sr/Ca, and Li/Mg results against seawater temperature in Fig. 6. The narrow seawater temperature range of these regression equations mean that they are of little use for paleo-proxy studies. However, the absolute values, range of trace metal results, and \( R^2 \) values of the regressions give a sense of microstructural bias and the broader temperature-proxy-suitability of these sampling approaches for *Lophelia*. Bulk calyx replicates are on average 8 and 9% more enriched in Li and Mg respectively than theca-only samples from the same specimens (Fig. 6A and B). Despite averaging two bulk calyx replicates, the range in both Sr/Ca and Li/Mg results is still larger (30%) than theca sampling alone (Fig. 6C and D). While Sr/Ca results for both sampling approaches show little coherency with seawater temperature (\( R^2 < 0.22 \)), Li/Mg measured in *Lophelia* specimens sampled for just theca wall material yields a tighter relationship with temperature (\( R^2 > 0.7 \)) than bulk calyx sampling. Even if the one anomalous bulk calyx Li/Mg value for sample CinMSloph-004 is removed, the \( R^2 \) value of the bulk calyx exponential relationship remains less than 0.5. Taken together with \( \delta^{13} \)C and Mg/Ca results (Fig. 5B), we infer that high Li and Mg content and inconsistent Li/Mg ratios in bulk *Lophelia* are being driven, at least in part, by the incorporation of greater amounts of skeletal COCs in the septa, precipitated from a highly restricted ECF. Despite Li/Mg being considered minimally impacted by calcification processes (cf. Li/Ca and Mg/Ca), this interpretation is consistent with modelling results that suggest more extreme closed-system calcification does begin to modify skeletal Li/Mg (Marchitto et al., 2018). For this reason, we recommend that future paleo-temperature studies use *Lophelia* sample theca walls and avoid the inclusion of septa. Accordingly, for all temperature regression analysis using trace metal data in subsequent discussion (Fig. 7; Fig. 1B) we use theca samples for *Lophelia* specimens where available.

We assess the individual temperature dependency of each of our cold-water coral \( \delta^{18} \)O, Sr/Ca and Li/Mg proxy datasets (Fig. 7). In line with previous findings, we show that scleractinian coral \( \delta^{18} \)O is poorly correlated with seawater temperature (\( R^2 = 0.34 \); Fig. 7A), with little coherent relationship even if samples are separated at the genus level. Adjustment of these coral \( \delta^{18} \)O values for differences in \( \delta^{18} \)O of seawater at each site (i.e. \( \delta^{18} \)O_coral \( - \delta^{18} \)O_seawater) yields no better correlation with temperature (\( R^2 \) value reduces to 0.22) and is likely hindered by poorly constrained seawater values in the Chilean Fjord sites (Fig. 5A). Nevertheless, our result is consistent with the overwhelming influence of ion pumping and growth rate effects on scleractinian \( \delta^{18} \)O (Chen et al., 2018). Aragonitic stylasterids by contrast show generally higher \( \delta^{18} \)O values, with a stronger correlation with temperature (\( T \)) (Samperiz et al., 2020), described by the equation,

\[
\delta^{18} \text{O}_{\text{Stylerid(Arag)}} = -0.20 \pm 0.02 \times T \left( ^\circ \text{C} \right) + 3.02 \pm 0.14
\]

\( (R^2 = 0.84) \)

This is consistent with the inference from paired \( \delta^{18} \)O and \( \delta^{13} \)C above that stylasterid species precipitate close to seawater equilibrium. For direct comparison to trace metal temperature proxy results, in Equation (2) and Fig. 7A we show only the \( \delta^{18} \)O values for aragonitic stylasterid specimens used in this study and exclude data from other genera measured by Samperiz et al. (2020). These stylasterid \( \delta^{18} \)O are also unadjusted for \( \delta^{18} \)O of seawater since it is a value that is poorly constrained for paleo-temperature reconstructions. Nevertheless, the relationship we show in Equation (2) is similar to the full compiled aragonitic stylasterid \( \delta^{18} \)O-temperature calibration published by Samperiz et al. (2020):

\[
\delta^{18} \text{O}_{\text{Stylerid(Arag)}} = -0.22 \pm 0.01 \times T \left( ^\circ \text{C} \right) + 3.33 \pm 0.06
\]

\( (R^2 = 0.91) \)

Cold-water coral Sr/Ca values have a similar pattern to the \( \delta^{18} \)O data when comparing Scleractinia with aragonitic stylasterids (Fig. 7B). Scleractinian Sr/Ca is poorly correlated with seawater temperature (\( R^2 = 0.29 \)); a result that is again consistent with a dominant growth-rate effect in this group (Gagnon et al., 2007). However, aragonitic stylasterid Sr/Ca correlates with seawater temperature according to the equation,
Aragonitic stylasterid corals fit a similar, exponential relationship, that explains considerably more of the variance (95%) and with less overall scatter,

$$\text{Li/Mg}_{\text{Stylasterid(Arag)}} = 5.64 \pm 0.09 \times 10^{-0.046 \pm 0.003} \times T \left( ^{\circ}C \right)$$

$$\left( R^2 = 0.95 \right)$$

Regression analyses such as these are open to bias by a small number of data points if the ambient seawater temperatures for the sample set are unevenly distributed. The aragonitic stylasterid calibrations might therefore be strongly weighted by the one specimen collected from warmer waters ($\sim 16^{\circ}C$). In this case however, exclusion of this sample has little impact on slope and intercept of the regressions outlined above (Aragonistic stylasterids $< 10^{\circ}C$:

$$\text{Li/Mg} = 5.67 \ e^{-0.047} \times T \left( ^{\circ}C \right); \ R^2 = 0.90.$$  

These results open the possibility of $\delta^{18}O$, Sr/Ca and Li/Mg based temperature proxy reconstructions using aragonitic stylasters. However, the strong geochemical dependence on mineralogy requires that sample mineralogy must be known so that specimens containing a mixture of aragonitic and high-Mg calcite are avoided (Fig. 4; Fig. 7B and E).

Through combining our scleractinia and aragonitic stylasterid results with published Li/Mg measurements in biogenic aragonites (including foraminifera and zooxanthellate corals) we find the following relationship with temperature for all aragonitic corals and foraminifera across the range 0 to 30 $^{\circ}C$ (Fig. 1B),

$$\text{Li/Mg}_{\text{All aragonite}} = 5.42 \pm 0.04 \times 10^{-0.050 \pm 0.0004} \times T \left( ^{\circ}C \right)$$

$$\left( R^2 = 0.97 \right)$$

This refined calibration is within uncertainty of the original exponential fit by Montagna et al. (2014), however, the addition of these new cold-water coral specimens means that we can now explain more of the variance in Li/Mg. We also greatly expand the calibration dataset that was previously dominated by shallow-water zooxanthellate corals. The inclusion of aragonitic stylasters in this calibration further supports that this relationship to temperature is universal to all biogenic aragonites and Li/Mg is minimally impacted by growth rates and organism specific vital effects. This same calibration can therefore be used to generate paleo-temperature records from aragonitic foraminifera collected from deep-sea sediment cores (e.g. Marchitto et al., 2018), individual coral cores (e.g. Fowell et al., 2016), and radiometrically dated solitary cold-water coral samples of any species (e.g. Case et al., 2010).

3.3.2. Li/Mg in high-Mg calcite stylasterids

*E. gracilis* samples (both aragonitic and calcitic) from this sample set vary along a clear mixing line between high Li/Mg values in solely aragonitic stylasterids and low Li/Mg values in solely calcitic. Until further data are generated for this group and other mixed mineralogy carbonates (e.g. Molluscs; Dellingler et al., 2018) across a wide range of hydrographic conditions, we must assume that mixed aragonitic and calcitic stylasterids have little apparent paleo-temperature proxy potential and we exclude them from further discussion.

There have been few studies exploring the Li/Mg temperature relationship in biogenic high-Mg calcites. However, Li/Mg data measured in CCA specimens grown in culture, across a 7 $^{\circ}C$ temperature range, suggest a linear relationship (Fig. 1A; Table 3; Anagnostou et al., 2019). The nature of a Li/Mg-temperature relationship in high-Mg calcite organisms is questioned however by calcitic coral data from across a 3.5 $^{\circ}C$ temperature range that show little temperature sensitivity (Fig. 1A; Chaabane et al., 2019). Yet, over a narrow temperature ranges such as these, it remains impossible to ascertain if the relationship is in fact exponential as it is in aragonitic organisms, and further data are required.

\[ \text{Sr/Ca}_{\text{Stylasterid(Arag)}} = -0.11 \pm 0.02 \times \ T \left( ^{\circ}C \right) + 11.61 \pm 0.09 \quad (R^2 = 0.79) \]

$c_{\text{Li/Ca}}$ and $c_{\text{Mg/Ca}}$ results show respectively weak negative and weak positive relationships with temperature for scleractinia, aragonitic stylasterids, and also high-Mg calcite (Stylasterids and CCA) groups (Fig. 7C and D). While correlations are too weak to determine if these relationships are exponential (e.g. Montagna et al., 2014), $c_{\text{Li/Ca}}$ and $c_{\text{Mg/Ca}}$ ratios both change on the order of 2.5% per $^{\circ}C$ for both aragonitic and high-Mg calcites. Similar to $\delta^{18}O$ and $c_{\text{Sr/Ca}}$ results, we find aragonitic stylasterid $c_{\text{Li/Ca}}$ and $c_{\text{Mg/Ca}}$ ratios to have slightly stronger correlations with temperature than scleractinia, however $R^2$ values remain lower than 0.63.

By contrast, $c_{\text{Li/Mg}}$ ratios show by far the most coherent relationships with temperature across the coral groups in this study (Fig. 7E). Consistent with previous studies (Case et al., 2010; Montagna et al., 2014) scleractinian coral $c_{\text{Li/Mg}}$ data are well correlated with seawater temperature, fitting an exponential relationship.

$$\text{Li/Mg}_{\text{Scleractinia}} = 5.57 \pm 0.12 \times 10^{-0.053 \pm 0.003} \times T \left( ^{\circ}C \right)$$

$$\left( R^2 = 0.77 \right)$$

![Fig. 6. Impact of sampling technique on *Lophea* Li/Ca, Mg/Ca, Sr/Ca, and Li/Mg ratios plotted against temperature. We compare the average of two bulk calyx replicates and theca only samples (presumed lower DCO content). The Li measurement discussed in the main text as potentially anomalous is highlighted by a red circle. Analytical uncertainty is smaller than the marker size.](image-url)
Fig. 7. New coral and coralline algae δ¹⁸O, Sr/Ca, Li/Ca, Mg/Ca, and Li/Mg temperature calibration data in this study. δ¹⁸O data for stylasterid corals from Samperiz et al. (2020). Averages of replicates are plotted, however, where available for trace metal data, Lophelia samples are replaced by the theca sample (see Fig. 6). Black and red trend lines are for respectively aragonitic stylasterids and scleractinia data. In all plots analytical uncertainty is smaller than the marker size.

Table 3
Assessment of temperature reconstruction uncertainty using δ¹⁸O, Sr/Ca and Li/Mg proxy calibrations. Confidence and prediction intervals are assessed at the 95% confidence level. Temperature uncertainties are read from confidence and prediction intervals at Li/Mg = 4 mmol/mol and 0.4 mmol/mol for aragonitic and calcitic calibrations respectively. Sr/Ca and δ¹⁸O intervals are read at mean values for data in this study: 11 mmol/mol and 1.5‰. Note, these values are illustrative for comparison between proxies and prediction uncertainty will change across the temperature range. For temperature reconstruction we do not recommend the use of cold-water scleractinia Sr/Ca and δ¹⁸O calibrations (greyed out) owing to their large prediction intervals (>±6°C).

<table>
<thead>
<tr>
<th>Calibration</th>
<th>Confidence interval temperature uncert. (± °C)</th>
<th>Prediction interval temperature uncert. (± °C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Li/Mg</td>
<td>Li/Mg = 5.27 exp(-0.049 x T) (Equation (1))</td>
<td>0.35</td>
</tr>
<tr>
<td>Li/Mg</td>
<td>Li/Mg = 5.42 exp(-0.050 x T) (Equation (7))</td>
<td>0.25</td>
</tr>
<tr>
<td>Li/Mg</td>
<td>Li/Mg = 5.64 exp(-0.046 x T) (Equation (6))</td>
<td>0.65</td>
</tr>
<tr>
<td>Sr/Ca</td>
<td>Sr/Ca = -0.08 x T + 11.06 (Equation (4))</td>
<td>2.16</td>
</tr>
<tr>
<td>Aragonitic stylasterids (this study)</td>
<td>Sr/Ca = -0.11 x T + 11.61 (Equation (6))</td>
<td>1.33</td>
</tr>
<tr>
<td>Scleractinia (this study)</td>
<td>δ¹⁸O = -0.28 x T + 2.49 (Equation (8))</td>
<td>1.16</td>
</tr>
<tr>
<td>Aragonitic stylasterids (samples in this study)</td>
<td>δ¹⁸O = -0.20 x T + 3.02 (Equation (2))</td>
<td>1.45</td>
</tr>
<tr>
<td>Aragonitic stylasterids (Samperiz et al., 2020)</td>
<td>δ¹⁸O = -0.22 x T + 3.33 (Equation (3))</td>
<td>3.5</td>
</tr>
<tr>
<td>Scleractinia (this study)</td>
<td>Scleractinia (this study)</td>
<td></td>
</tr>
<tr>
<td>Aragonitic stylasterids (samples in this study)</td>
<td>Scleractinia (this study)</td>
<td>1.16</td>
</tr>
<tr>
<td>Li/Mg</td>
<td>Li/Mg = -0.02 x T + 0.57 (Equation (8))</td>
<td>0.51</td>
</tr>
<tr>
<td>Li/Mg</td>
<td>Li/Mg = 0.63 exp(-0.050 x T) (Equation (8))</td>
<td>0.51</td>
</tr>
</tbody>
</table>
When the Li/Mg results of high-Mg calcite stylasterids are combined with our data from shallow-water CCA collected from the Caribbean, published values from Anagnostou et al. (2019), as well as Corallium rubrum data from Chaabane et al. (2019), we find the following exponential relationship for high-Mg calcite organisms (Fig. 1B).

\[
\text{Li/Mg}_{\text{all high-Mg calcite}} = 0.63 \pm 0.02 \times 10^{-0.050 \pm 0.002} \times T(\degree C)
\]

\[(R^2 = 0.92)\] (8)

While the higher Mg content of these organisms determines that the intercept of this trendline is lower compared to biogenic aragonites, we find that the exponent of Equation (7) and Equation (8) is remarkably similar. This encouraging result invites further study to characterise the Li/Mg relationship to temperature in multiple high-Mg calcitic organisms to fully assess their paleo-oceanographic utility.

3.4. Biological controls on trace metal incorporation

Our compiled data show that Li/Mg ratios in both biogenic aragonite and high-Mg calcites decrease by 5% per °C change in seawater temperature. Given the phylogenetic separation of groups within each temperature calibration as high as Kingdom level (Fig. 2), this result strongly suggests an abiogenic incorporation mechanism exists for these elements. While \(D_{\text{Li}}\) and \(D_{\text{Mg}}\) differ greatly between aragonites and high-Mg calcites in absolute terms, Li/Ca and Mg/Ca values in this study respectively increase and decrease on the order of 2.5% per °C in both of these types of carbonate (Fig. 7C and D). It is likely therefore that similarity between exponents of Li/Mg temperature relationships in our aragonite and high-Mg calcite samples is driven by a largely equal abiogenic temperature dependency on both partition coefficients. Further study into the controls on Li/Mg ratios in low-Mg calcites is required to ascertain if this temperature sensitivity is universal to all calcium carbonates.

While we find temperature to be an important factor driving both Sr/Ca and Li/Mg in scleractinian and stylasterid corals, the secondary biological controls on incorporation of these elements have long been debated (Marchitto et al., 2018; Montagna et al., 2014). We find that the residuals for the Sr/Ca and Li/Mg relationships with temperature (Fig. 7B and Fig. 7E) are positively correlated for scleractinian corals according to the equation:

\[
\text{Sr/Ca residuals} = 0.55 \times \text{Li/Mg residuals} + 0.01 \quad (R^2 = 0.35) \] (9)

Although correlation is weak, this relationship is similar to that reported for scleractinian corals by Ross et al. (2019). Yet, we find no such correlation \((R^2 = 0.01)\) for the residuals for aragonitic stylasterid coral temperature relationships:

\[
\text{Sr/Ca residuals} = 0.10 \times \text{Li/Mg residuals} + 0.01 \quad (R^2 = 0.01) \] (10)

Li/Mg is thought to be unaffected by [Ca] of the ECF, however, biogenic carbonate precipitation experiments suggest that the partition coefficient of Li/Mg is sensitive to carbonate ion concentration (Holcomb et al., 2016; Ross et al., 2019). Sr/Ca may also be indirectly influenced by [CO\(_3\)\(^{2-}\)] in the ECF if kinetic effects and/or Rayleigh fractionation become more dominant as [CO\(_3\)\(^{2-}\)] increases (Ross et al., 2019). The absence of correlation between residuals in aragonitic stylasterid corals that calcify close to equilibrium with seawater suggests that modification of ECF carbonate chemistry exerts a secondary control on both Sr/Ca and Li/Mg ratios in scleractinian corals.

3.5. Temperature prediction uncertainty

When interpreting any paleoceanographic record it is important to consider the fidelity with which a proxy is able reproduce the hydrographic parameter of choice. To this end, it has become the convention in many proxy calibration studies to quote confidence intervals on the regression (e.g. Anand et al., 2003). This approach however merely gives a measure of how well the “true” position of the regression line is known and not how accurately seawater temperature can be predicted from a single geochemical measurement. Therefore, in Table 3 we summarise the temperature uncertainty using confidence intervals and prediction intervals (2σ; assuming that future observations have the same error variance as those used for fitting) that better characterises proxy reliability and discern the results below.

3.5.1. All biogenic aragonites

Combination of our aragonitic stylasterid and scleractinian coral Li/Mg data with published biogenic aragonite Li/Mg temperature calibrations (Equation (7)) offers an improvement in confidence intervals to ±0.25 °C (for a Li/Mg value of 4 mmol/mol) over Li/Mg calibration data from previous studies alone (Fig. 1). Despite the exponential fit now explaining more of the variance, addition of our data only slightly reduces prediction interval uncertainty for “All aragonite” Li/Mg, which is now ±3.4 °C. This result suggests that our data, and that of previous studies, are adequately capturing the variance about the exponential relationship, thus giving a representative estimate of the ability of the Li/Mg proxy to reproduce past seawater temperatures.

3.5.2. Aragonitic stylasterid corals

As this is the first study of stylasterid trace metal chemistry, we have only the 16 observations in this study with which to characterise the aragonitic stylasterid relationships with temperature (Equations (4) and (6)). With fewer observations, these temperature calibrations on their own have wider confidence intervals than that of “All aragonites” at more than ±0.6 °C. Despite this, the prediction intervals of ±5.3 °C show the aragonitic stylasterid Sr/Ca linear temperature calibration (Equation (4)) to be a useful paleo-temperature monitor. The narrower prediction interval of aragonitic stylasterid Li/Mg however, suggests that this has the most potential as a precise seawater temperature proxy with a reconstruction uncertainty of just ±2.3 °C (Equation (6)). This result further implies that the removal of growth rate effects by ratioing Li to Mg is more robust for organisms that precipitate their skeletal carbonate closer to equilibrium with seawater.

3.5.3. High-Mg calcites

We propose a new exponential relationship of Li/Mg to temperature in high-Mg calcites that includes calcitic corals and CCA (Equation (8)). Prediction intervals of ±3.4 °C for this relationship suggest an improvement in temperature reproducibility over the linear relationship proposed for CCA grown in culture alone (Anagnostou et al., 2019). This “all high-Mg calcites” relationship is determined using data from nine discrete temperature conditions, without overlap between biological groups (i.e. stylasterids low temperature, CCA and Corallium high temperature). Further interrogation of this relationship is required to test its robustness across the full seawater temperature range and in particular the robustness of inter-species offsets between high-Mg calcite organisms.

3.6. Conclusions

Depending on their skeletal calcium carbonate polymorph, we find stylasterid coral geochemistry to be similar to other biogenic
aragonites or high-Mg calcite counterparts. If composed of just one carbonate polymorph (i.e. solely aragonite or high-Mg calcite) stylasterid corals have both δ18O values close to equilibrium with seawater and little variability in δ18O, Sr/Ca, and Li/Mg between single sample replicates from a single specimen. This strongly suggests that stylasterids perform little modification of local/internal seawater. Such open system calcification means that stylasterid corals offer a wide array of accurate temperature proxy capabilities beyond that of scleractinia.

Our new data from modern cold-water coral specimens redress the balance of previous temperature calibrations biased towards shallow-water zoanthellate corals. In accordance with previous studies, we find poor correlation between scleractinian δ18O and Sr/Ca with seawater temperature. With the exception of stylasterid corals, we therefore do not advocate the use of δ18O and Sr/Ca ratios for paleo-temperature reconstructions using multiple cold-water coral individuals.

Our coral Li/Mg data show the strongest relationship with temperature; thus, we are better able to characterise the Li/Mg exponential relationship in biogenic aragonites for seawater temperatures below 16°C. Our data further suggest that this calibration is universal to all biogenic aragonites, demonstrating Li/Mg to be an extremely accurate and versatile temperature proxy, particularly in aragonitic Stylasteridae corals. We show that a similar exponential temperature dependency also exists for Li/Mg ratios across all high-Mg calcitic taxa. Future implementation of these proxies for paleo-temperature estimates in fossilised biogenic carbonates, will therefore deepen our knowledge of the thermal history of the ocean.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary material

Supplementary material related to this article can be found online at https://doi.org/10.1016/j.epsl.2020.116412.

References

tion in Porites on coral Sr/Ca and δ18O paleotemperometry. Paleoceanography 27.
E11023.
effects correlated with skeletal architecture in a scleractinian deep-sea coral and
Guo, W., 2020. Kinetic clumped isotope fractionation in the DIC-H2O-CO2 system:
Hall, J.M., Chan, L.H., 2004. Li/Ca in multiple species of benthic and plank-
tonic foraminifera: thermocline, latitudinal, and glacial-interglacial variation.
aragonite skeletons of massive Porites corals: a new tool to reconstruct tropical
sea surface temperatures. Paleoceanography 28, 143–152.
Hathorne, E.C., Gagnon, A., Felis, T., Atkins, J., Asami, Y., Boer, W., Caillon, N., Case,
D., Cobb, K.M., Douville, E., deMenocal, P., Eisnerhauer, A., Garbe-Schönberg,
D., Geibert, W., Goldstein, S., Huguen, K., Inoue, M., Kawahata, H., Kölling, M.,
Corne, F.L., Linsley, B.K., Mersch, H.V., Montagna, P., Nishita, I.S., Quinn, T.M.,
Raddatz, J., Rehbach, H., Robinson, L., Sadekova, A., Sherrrell, R., Sinclair, D.,
study for coral Sr/Ca and other element/Ca ratio measurements. Geochim. Geo-
Jurikova, H., Liebetrau, V., Raddatz, J., Fietzke, J., Trotter, J., Rocholl, A., Krause, S.,
of the cold-water coral Lophelia pertusa along the Norwegian margin: zooming
into a potential pH-proxy by combining bulk and high-resolution approaches.
Chem. Geol. 513, 143–152.
Key, R.M., Kozyr, A., Sabine, C.L., Lee, K., Wanninkhof, R., Bullister, J.L., Feely, R.A.,
Locarnini, R.A., Mishonov, A.V., Antonov, J.I., Boyer, T.P., Garcia, H.E., Baranova, O.K.,
Zheng, M.M., Faver, C.R., Reager, J.T., Johnson, D.R., Hamilton, M., Seidov, D.,
NOAA Atlas NESDIS.
biominerolization model to explain Li, Mg, and Sr incorporation into aragonitic
Marriott, C.S., Henderson, C.M., Belshaw, N.S., Tudhope, A.W., 2004. Temperature de-
pendence of δ3Li, 48Ca and Li/Ca during growth of calcitic corals. Earth
McConnaughey, T., 1980, 18O and 16O isotopic disequilibrium in biological carbon-
McCulloch, M.T., Tudhope, A.W., Esat, T.M., Mortimer, C.G., Chappell, J., Pillans, B.,
during the Pemulwuy Deglaciation at Huon Peninsula. Science 283, 202.
Montagna, P., McCulloch, M., Douville, E., López Correa, M., Trotter, J., Rodolfo-
Metalpa, R., Dissard, D., Ferrier-Pagès, C., Frank, N., Freiwald, A., Goldstein, S.,
Mazzoli, C., Reynaud, S., Rüggeberg, A., Russo, S., Taviani, M., 2014. Li/Mg sys-
tematics in scleractinian corals: calibration of the thermometer. Geochim. Cos-
Poggelein, D.W., Hathorne, E.C., Nürnberg, D., Frank, M., Bruhn, L., Reebig, S., Bahr,
A., 2017. Rapid deglacial injection of nutrients into the tropical Atlantic via
Raddatz, J., Liebetrau, V., Rüggeberg, A., Hathorne, E., Krabbenhöft, A., Eisnerhauer,
A., Böhm, F., Völstaedt, H., Fietzke, J., López Correa, M., Freiwald, A., Dullo, W.C.,
2013. Stable Sr-isotope, Sr/Ca, Mg/Ca, Li/Ca and Mg/Li ratios in the scleractinian
cold-water coral Lophelia pertusa. Chem. Geol. 352, 143–152.
Rae, J.W.B., Foster, G.L., Schmidt, D.N., Elliott, T., 2011. Boron isotopes and B/Ca in
benthic foraminifera: proxies for the deep ocean carbonate system. Earth Plan.
Sci. Lett. 302, 401–413.
Roberts, J., Gottschalk, J., Skinner, L.C., Peck, V.L., Kender, S., Elderfield, H., Wael-
density and chemical stratification across the last deglaciation. Proc. Natl. Acad.
Sci. USA 113, 514.
geology of cold-water coral ecosystems. Science 312, 543–547.
Ross, C.L., DeCarlo, T.M., McCulloch, M.T., 2019. Calibration of Sr/Ca, Li/Mg and Sr-U
Paleothermometry in branching and foliose corals. Paleoenviron. Paleoclima-
tol. 34, 1271–1291.
v1.22, NASA.
in a physicochemical calcification system: application to coral bimineralization
Sponer, P.T., Guo, W., Robinson, L.F., Thiagarajan, N., Hendry, K.R., Rosenheim, B.E.,
Leng, M.J., 2016. Clumped isotope composition of cold-water corals: a role for
 calibration for the deep-sea coral Desmosophyllum diadexus through sub-sampling
for boron isotope geochemistry (abstract). In: AGU Fall Meeting, San Francisco,
USA.
Bolling–Allerød warming and circulation changes in the deep ocean. Nature 511,
75.
aragonite and seawater: developing a novel kinetic oxygen isotope fractionation
Watson, E.B., 2004. A conceptual model for near-surface kinetic controls on the
trace-element and stable isotope composition of aragonitic calcite crystals.