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8 Environmental and biological controls on Mg and Li in
9 deep-sea scleractinian corals
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46 **Abstract**

47

48 Deep-sea scleractinian corals precipitate aragonite skeletons that provide valuable
49 archives of past ocean conditions. During calcification biological mediation causes variability in
50 trace metal incorporation and isotopic ratios of the aragonite such that signals caused by
51 environmental controls can be overwhelmed. This complicates the interpretation of geochemical
52 proxies used for paleo-reconstructions. In this study we examine the environmental controls on
53 the Mg/Li ratio of 34 individuals from seven genera of deep-sea scleractinian corals:

54 *Desmophyllum*, *Balanophyllia*, *Caryophyllia*, *Enallopsammia*, *Flabellum*, *Trochocyathus*, and
55 *Lophelia*. In addition we examine the distributions of Mg and Li in *Desmophyllum* and
56 *Balanophyllia* using laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS).

57 Both Mg/Ca and Li/Ca ratios increased by more than a factor of 2 in the center of
58 calcification regions compared to the outer, fibrous regions of the coral skeleton. As a result,
59 replicate ~10 mg subsamples of coral show less variability in the Mg/Li ratio than Mg/Ca.
60 Microscale Mg and Li results are consistent with Rayleigh-type incorporation of trace metals
61 with additional processes dominating composition within centers of calcification. Comparison of
62 Mg/Li to seawater properties near the site of collection shows that the ratio is not controlled by
63 either carbonate ion or salinity. It appears that temperature is the major control on the Mg/Li
64 ratio. For all 34 samples the temperature correlation ($R^2=0.62$) is significantly better than for
65 Mg/Ca ($R^2=0.06$). For corals of the family *Caryophyllidae* the R^2 value increases to 0.82 with
66 the exclusion of one sample that was observed to have an altered, chalky texture. Despite this
67 excellent correlation the scatter in the data suggests that the Mg/Li ratio of deep-sea corals

68 cannot be used to reconstruct temperature to better than approximately $\pm 1.6^{\circ}\text{C}$ without better
69 temperature control and additional calibration points on modern coral samples.

70

71 Keywords: biomineralization, paleoceanography, deep-sea coral, Mg/Ca, Mg/Li, thermometry

72

73 **1. Introduction**

74

75 Ice core records show that there have been large and abrupt changes in atmospheric
76 temperature in the past, as particularly well documented during the last deglaciation (Grootes et
77 al., 1993). Since approximately half of the cross-equatorial heat transport occurs in the ocean
78 (Ganachaud and Wunsch, 2000) there are likely to be important links between ocean circulation
79 and climate (Rahmstorf, 2002). High-resolution, well-dated records of deep-water temperatures
80 are difficult to obtain because of the challenges of bioturbation and constructing precise
81 chronologies in sediment cores. Using the skeletons of deep-sea corals to reconstruct seawater
82 temperature in the past would circumvent some of these challenges given a suitable geochemical
83 proxy for temperature.

84 Deep-sea corals inhabit a wide variety of depths and geographic distributions, including
85 high-latitude regions and all ocean basins (Cairns, 2007). Scleractinian corals are made of
86 aragonite and datable using U/Th techniques, allowing accurate and precise ages to be calculated
87 for each individual (Cheng et al., 2000). Solitary deep-sea scleractinians have been collected
88 from diverse settings, and fossil specimens in existing collections span hundreds of thousands of
89 years (e.g., Robinson et al., 2006). *Desmophyllum dianthus*, a species that has been used in a

90 number of paleo-studies, live on the order of 100 years providing potential centennial-scale
91 archives (Adkins et al., 1998).

92 Despite the promise of deep-sea corals for reconstructing past ocean circulation,
93 paleoceanographic reconstruction of temperature has been complicated by vital effects. This
94 broad term represents the influence of biomineralization on skeletal composition. The near
95 constant environmental conditions of the deep sea over the roughly 100 year lifetime of modern
96 corals and the lack of photosymbionts mean that compositional variability in the coral skeleton
97 can be attributed entirely to the processes of coral biomineralization.

98 Stable isotopes and trace element ratios commonly used in paleoceanography, including
99 $\delta^{13}\text{C}$, $\delta^{18}\text{O}$, Mg/Ca, Sr/Ca, and U/Ca all vary across deep-sea scleractinian skeletal features
100 (Adkins et al., 2003; Cohen et al., 2006; Gagnon et al., 2007; Robinson et al., 2006; Sinclair et
101 al., 2006; Smith et al., 2000; Smith et al., 2002). The magnitude of this variation changes from
102 proxy to proxy, but for temperature reconstructions can overwhelm environmental controls.
103 Much of the compositional variability in coral follows differences in skeletal architecture. On the
104 macroscale, corals contain septal and thecal features, the former being comprised of septa
105 radiating about the axis of the coral and the latter comprising the thick walls of the coral
106 structure. Scleractinian coral skeletons have two distinctive micro-scale architectures: the
107 Centers of Calcification (COCs) which contain small, granular, disorganized crystals, and the
108 surrounding region which are comprised of fibrous bundles of aragonite needles (Gladfelter,
109 1982; Stolarski, 2003; Gladfelter, 2007). These features have also been documented in
110 scleractinian deep-sea corals (Adkins et al., 2003; Robinson et al., 2006; Gagnon et al., 2007;
111 Rollion-Bard et al., 2010). In the deep-sea coral *D. dianthus*, Mg/Ca nearly doubles in the COCs
112 (Gagnon et al., 2007), while $\delta^{13}\text{C}$ and $\delta^{18}\text{O}$ are depleted (Adkins et al., 2003). These results may

113 be interpreted as evidence that separate mechanisms control composition in different regions of
114 the coral skeleton. However, Sr/Ca seems unaffected by skeletal architecture (Gagnon et al.,
115 2007) while [U] is lower in regions associated with, but extending beyond, the COCs (Robinson
116 et al., 2006), suggesting some continuity of mechanism across the whole coral skeleton.

117 Geochemical models of biomineralization attempt to explain the mechanisms controlling
118 the distribution of trace metals and isotopes in coral skeletons. Models can be distinguished by
119 the most important step or process proposed to control skeletal composition. Some of these
120 processes include ion transport, pH-driven solution chemistry (e.g.: McConnaughey, 1989;
121 Adkins et al., 2003), precipitation from a closed system (e.g.: Gaetani & Cohen, 2006; Cohen et
122 al., 2006; Gagnon et al., 2007), the role of an organic matrix during precipitation, and the
123 presence of transient amorphous phases (Rollion-Bard et al., 2010). In this paper we examine Mg
124 and Li in *D. dianthus* accompanied by a discussion of mechanisms controlling the incorporation
125 and variation of these elements.

126 Most existing models agree that coral skeletogenesis occurs in a biologically controlled
127 space with a unique composition called the “extracytoplasmic calcifying fluid,” or ECF
128 (Johnson, 1982; McConnaughey, 1989; Adkins et al., 2003). The source and transport of skeletal
129 ions into the ECF has the potential to affect skeletal composition and cause vital effects. While
130 the extent of seawater leakage and its influence on calcification is a matter of debate, transport of
131 bulky fluorescent dyes to the skeleton implicates direct seawater transport (Erez & Braun, 2007).

132 Consideration of the ECF as a closed system has prompted the application of Rayleigh
133 fractionation principles to trace metal incorporation (Gaetani & Cohen, 2006; Cohen et al., 2006;
134 Gagnon et al., 2007; Rollion-Bard et al., 2009). This class of model describes a system in which
135 trace metals are precipitated into the aragonite lattice from an ECF that initially matches the

136 seawater composition, but is subsequently closed with regard to ion transport. As precipitation
137 proceeds the ECF becomes depleted in calcium and depleted or enriched in trace metals
138 depending on their distribution coefficient. Cohen et al. (2006) and Gagnon et al. (2007) have
139 applied this theory to model Sr/Ca and Mg/Ca variability in deep-sea corals where Sr has a
140 distribution coefficient >1 while that of Mg is $\ll 1$. Gagnon et al. (2007) concluded that Rayleigh
141 fractionation is consistent with their correlated Me/Ca variability outside of obvious *D. dianthus*
142 COCs. However, an equally important conclusion of that study is that large Mg/Ca variability
143 associated with COCs is dominated by a mechanism other than a Rayleigh process

144 The pathway for incorporation of Mg in aragonite is not well constrained (Fallon et al.,
145 2003). While Mg^{2+} may be directly substituted for Ca^{2+} as has been documented in calcite (Politi
146 et al., 2010) the relatively small ionic radius of Mg^{2+} severely distorts the 6-fold coordinated
147 calcite lattice, and it is likely that the aragonite lattice of scleractinian corals would be even more
148 severely distorted. However since Mg is typically much less abundant in aragonite than in
149 calcite, the effect of such distortions may be less significant. Finch & Allison (2008) employed
150 X-ray Absorption Fine Structure (XAFS) to determine the structural state of Mg in scleractinian
151 coral samples and the authors interpret their results to indicate that Mg is not directly hosted in
152 aragonite. These data may mean Mg is located in disordered a site, consistent with either organic
153 binding or the presence of amorphous calcium carbonate (ACC). Alternatively Mg^{2+} in corals
154 may be loosely bound in the aragonite lattice or adsorbed to crystal surfaces (Amiel et al., 1973;
155 Walls et al., 1977). Regarding Li incorporation, much less is known. Li^+ incorporation has been
156 observed in inorganic aragonite experiments (Okumura & Kitano, 1986), but Li^+ substitution for
157 Ca^{2+} is more complicated than Mg^{2+} because of the difference in charge. We compare some of
158 these mechanisms to our data in the discussion.

159 Bryan & Marchitto (2008) discovered a strong Mg/Li-to-temperature correlation in the
160 shells of calcitic and aragonitic foraminifera which was significantly better than that for Mg/Ca.
161 This study evaluates major environmental controls on Mg/Li by comparing the measured ratio of
162 seven genera of deep-sea corals with ambient seawater. We also build on our understanding of
163 coral vital effects and geochemistry by analyzing the microscale distribution of Mg and Li in *D.*
164 *dianthus* and *Balanophyllia*.

165

166 **2. Materials and Methods**

167

168 *2.1 Sample selection*

169

170 Thirty-four separate coral individuals representing seven scleractinian genera were
171 selected: *Desmophyllum* (n=22), *Balanophyllia* (n=4), *Caryophyllia* (n=3), *Flabellum* (n=1),
172 *Trochocyathus* (n=1), *Enallopsammia* (n=2), and *Lophelia* (n=1) (Table 1). The first five are
173 solitary corals while the latter two are colonial. The samples were selected to represent diverse
174 oceanographic settings in all major ocean basins (Figure 1), and cover a depth range of 120 to
175 1818 meters. Environmental parameters, including temperature, salinity, and carbonate ion
176 concentration were derived from the GLODAP database (Key et al., 2004, Table 1). Typically,
177 the match between coral and water column location was within 3.5° latitude and longitude and
178 closer than 30 meters in depth.

179 Corals in this study were either loaned from the Smithsonian Museum of Natural History
180 by Stephen Cairns or collected on recent cruises to the North Atlantic or Southern Ocean. Corals
181 from recent cruises were either collected live or were radiocarbon dated to be recently dead

182 (Burke et al., in press, available online). For the case of the Southern Ocean samples temperature
183 and salinity were also available from nearby CTD casts taken on the same cruise, *Nathaniel B.*
184 *Palmer* 0805.

185 The temperature range for all samples is from 1.8 to 17.0°C. Salinity ranged from 34.04
186 to 36.54 ‰ and carbonate ion concentration from 1.2 to 192.7 µmol/kg (Table 1). Samples from
187 the Smithsonian museum were assumed to be modern, although two (48738 and 77019)
188 exhibited a visibly chalky texture indicating that they may have experienced post-mortem
189 alteration.

190

191 *2.2 Sample preparation*

192

193 Coral samples were prepared for two different types of analyses: bulk solution ICP-MS
194 and laser ablation ICP-MS. For bulk solution ICP-MS, samples ranging from 10 to 136 mg were
195 cut from the coral using a dremel tool. Subsamples came from both thecal and septal regions, and
196 some coral individuals were subsampled more than once. At most four subsamples were taken
197 from a single coral individual. All corals were chemically cleaned at WHOI using the oxidative
198 and reductive steps of Cheng et al. (2000). After cleaning, samples were dissolved in 5% Optima
199 Grade nitric acid, diluted to 60 ppm Ca, and spiked to 0.5 ppb In.

200 To prepare for ICP-MS laser ablation, sections of coral were cut perpendicular to the
201 coral growth axis with a dremel saw, bound to a microscope slide using epoxy (polymerized
202 Araldite resin, Ernest F. Fullam, Inc.) and polished with 0.18 µm silica. Images of each section
203 were taken under reflected light.

204

205 *2.3 Standard mixing for ICP-MS*

206

207 Gravimetric elemental solutions were prepared for two standard curves to account for
208 matrix effects, background corrections and differences in elemental ionization. The first set had
209 constant Me/Ca ratios diluted into five solutions with concentrations of Ca varying from 8.8 to
210 79.6 ppm. The second standard series contained constant Sr, Ba, Mg, and Li concentrations but
211 varying Ca concentrations over a range of 0 to 175 ppm, encompassing ranges broadly observed
212 in coral skeletons (McCulloch et al., 2003; Gagnon et al., 2007; Allison et al., 2007; Mitsuguchi
213 et al., 2008). A separately mixed Fe (16 ppb), Mn (16 ppb), and Ca (57 ppm) standard was also
214 prepared. All standards and solution samples were spiked to 0.5 ppb In to monitor ICP-MS
215 intensity drift.

216

217 *2.4 Bulk solution ICP-MS*

218

219 Solution samples were run on a Thermo Element2 ICP-MS, tuned to yield at least
220 275,000 counts per second (cps) for a 0.5 ppb In solution. Full standard curves were run at the
221 beginning and end of each session and between every 10-15 samples. A consistency standard,
222 S13, and 5% Optima Grade nitric acid blanks were run between every four samples. Every coral
223 sample aliquot was measured twice. The isotopes ^7Li , ^{25}Mg , ^{48}Ca , ^{88}Sr , and ^{138}Ba were measured
224 in low resolution with a mass resolving power of 300 ($M/\Delta M$ at 5% peak height). Medium-
225 resolution mode was used for ^{55}Mn and ^{56}Fe . For each isotope the sampling time was 0.03
226 seconds, except for ^7Li which was increased to 0.30 seconds. Counting mode was used to
227 measure ^7Li , ^{115}In , ^{138}Ba , ^{55}Mn , and ^{56}Fe while ^{48}Ca and ^{88}Sr were measured in analog mode.

228 ^{25}Mg varied between counting and analog modes ($\sim 4 \times 10^6$ cps). The method collected 600 cycles of
229 data per sample giving a total analysis time of ~ 3 minutes. Background corrections were made
230 by subtracting the intensity of bracketing acid blanks from standard and sample intensities. Rinse
231 time was three minutes between each sample resulting in background ^7Li counts less than 100
232 cps, only $\sim 0.5\%$ of our typical signal. Relative standard deviations for raw ratios were better than
233 2% for all samples and standards and were typically better than 1.5% ($1-\sigma$) for $^{25}\text{Mg}/^7\text{Li}$,
234 $^{25}\text{Mg}/^{48}\text{Ca}$, and $^7\text{Li}/^{48}\text{Ca}$. Since this study focuses on Mg, Li, and Ca, henceforth we only discuss
235 these elements. Fe and Mn results are semi-quantitative and are used only for the purposes of
236 checking for contamination.

237

238 *2.5 Data correction and reproducibility*

239

240 After the background correction, a number of additional corrections were made to the raw
241 data to account for dilution, matrix effect, element-specific ionization efficiency, and
242 instrumental drift. To account for small variations in the In spikes, the measured intensities were
243 corrected to the gravimetrically determined In concentration to the intensity expected for a 0.5
244 ppb solution. This correction allows direct comparison with standards for calculating elemental
245 concentrations. However, we only report final data as ratios and not concentrations.

246 The intensity of each isotope was normalized to the average value of In during each ICP-
247 MS run to account for drift. For the standard curves, each “Observed” ratio was compared to the
248 gravimetrically determined “Expected” ratio to ascertain the relative ionization efficiency. This
249 efficiency drifted by a maximum of $\sim 5\%$ on any one day. Within the Ca concentration range of
250 40 to 100 ppm the Observed/Expected ratios showed that there are no biases from matrix effects

251 or from the background correction. Our samples were therefore diluted to 60 ppm. A consistency
252 standard (S13) was used to test the reproducibility and accuracy, which were always within 2%
253 of expected values, and $^{25}\text{Mg}/^7\text{Li}$ was generally within 1%.

254 The same solution standards were used for laser ablation standardization, although a
255 blank subtraction was not applied because the solution acid blanks are not representative of the
256 blank levels in the laser ablation chamber. We expect the laser ablation background to be smaller
257 than for solution analyses, for which the background corrections were ~0.5%.

258

259 *2.6 Laser ablation ICP-MS*

260

261 Laser ablation was carried out using a 193 nm laser coupled to the same Thermo
262 Element2 ICP-MS with the same isotopes and ratios measured as for solution analyses. The laser
263 spot size was 50 μm and was run at a repetition of 20 Hz. Laser power was run at 100% capacity
264 and scan speed was set at 3 $\mu\text{m}/\text{sec}$. Ablated material was mixed with a 5% Optima grade acid
265 stream to provide a wet plasma equivalent to standard introduction. Solution standards are not
266 ideal since they are not matrix matched to the samples, so measured laser ablation values are
267 therefore assumed to be relative but not absolute. Even with this potential bias, our laser ablation
268 values are well within the range of the solution data for the same coral individuals (Figure 2,
269 Figure 3).

270 Three coral cross-sections were chosen for laser ablation. These included two sections
271 from a single *D. dianthus* individual (DMC08a and DMC08b) and one from a *Balanophyllia*.
272 Both corals were collected from the same site in the Southern Ocean (*Nathaniel B. Palmer* 0805,
273 TB04). Long laser lines were run perpendicular to coral density strata (i.e. across skeletal

274 features) and short lines were run parallel to coral density strata (i.e. within single skeletal
275 features). Each short line was run once and the standard deviations of the ratios were calculated
276 from the discrete analysis spots along the line. These standard deviations were generally 5% or
277 lower for $^{25}\text{Mg}/^{48}\text{Ca}$, $^7\text{Li}/^{48}\text{Ca}$, and $^{25}\text{Mg}/^7\text{Li}$, though in two cases the *Balanophyllia* $^{25}\text{Mg}/^7\text{Li}$
278 ratio approached 10%. Standard deviations of discrete analysis points for the long lines are not
279 representative of the analytical precision since each ablation line crosses regions with large
280 Me/Me variability. Instead the long lines were repeated up to three times in adjacent series,
281 providing an approximation of replicate measurements (Figure 2). The long lines proved to be
282 reproducible, often with 5% or less disagreement between adjacent lines (Figure 2).

283

284 **3. Results**

285

286 *3.1 Laser ablation ICP-MS*

287

288 Solution standardization of laser ablation is not ideal as it is not matrix matched, so we
289 restrict our discussion of laser data to relative changes within each coral section and qualitative
290 comparisons to biomineralization models. However, there is reasonable agreement between the
291 data collected by laser ablation and bulk solution (Figure 2, Figure 3).

292 Consistent with previous studies we observed large Mg/Ca variation associated with
293 changes in the skeletal architecture of the coral in *D. dianthus*. An overlay of trace metal data
294 was created by matching time-stamped data to coral features based on notes taken during the
295 laser ablation run (Figure 4). Mg/Ca was higher by more than a factor of 2 in COC regions as
296 opposed to outer fibrous bands. The Li/Ca ratio also increased by a more than a factor of 2 in the

297 COCs. In both cases the highest values were in the central, widest band in the coral, with local
298 maxima in less well-developed COCs. The Mg/Li ratio also shows distinct variability across the
299 *D. dianthus* skeletal sections but with a change of only a factor of 1.4. The largest maxima of
300 Mg/Li are associated with the most well developed COCs, but local maxima also appear in
301 regions not dominated by well-developed COCs (Figure 4a, b).

302 The microstructure of *Balanophyllia* is composed of COCs and fibrous material that
303 radiate outward (Figure 4c, Brahmi et al., 2010). The reflected light image of the *Balanophyllia*
304 has light and dark regions that appear to be compositionally equivalent to those in *D. dianthus*
305 (Brahmi et al., 2010). The geometry of *Balanophyllia* is more complex than for *D. dianthus* so
306 the laser ablation line was curved to stay perpendicular to the skeletal features. For this reason,
307 plots of trace metal data are not overlain on reflected light images (Figure 4c). Similar to *D.*
308 *dianthus* Mg/Ca and Li/Ca ratios vary by more than a factor of 2. Mg/Li variations in
309 *Balanophyllia* were observed throughout the skeleton and are much more frequent than those of
310 Mg/Ca or Li/Ca; it does not seem that Mg/Li variability is aligned with any specific skeletal
311 features (Figure 4c).

312 Overall LA-ICP-MS Mg/Ca ratios exhibit a strong positive relationship to Li/Ca for both
313 coral species: $R^2=0.94$ for *D. dianthus* and $R^2=0.75$ for *Balanophyllia*. Similarly there is a
314 positive correlation in multiple solution analyses from the same corals. There are two notable
315 deviations from the overall trend, one in each coral species. In *D. dianthus* section DMC08b, the
316 deviation from the regular slope is observed in the highest Mg/Ca region where Li/Ca reaches a
317 plateau. By comparing this to the reflected light image it can be seen that the location of this part
318 of the data is the central, most well-developed COC. In this area there is a distinct peak in
319 Mg/Ca, but Li/Ca does not continue to increase (Figure 4b). By contrast the *Balanophyllia*

320 sample exhibits a deviation in the lowest Mg/Ca region where again Mg/Ca varies and Li/Ca
321 does not (Figure 4c).

322

323 *3.2 Solution ICP-MS*

324

325 In total we analyzed 34 separate corals, 7 of which were subsampled more than once. For
326 all corals the Mg/Ca ratio ranged from 1.60 to 3.51 mmol/mol, Li/Ca from 7.05 to 14.99
327 $\mu\text{mol/mol}$, and Mg/Li from 0.20 to 0.35 mol/mmol (Table 1). Replicate analyses of Mg/Ca,
328 Li/Ca and Mg/Li for the same aliquots were always within our estimated reproducibility from the
329 internal standard S13. Analyses of separate subsamples varied by at most 11% for Mg/Li, 27%
330 Mg/Ca, and 29% for Li/Ca, demonstrating variability within coral individuals. For example coral
331 48738, in which 4 subsamples were analyzed, ranged from 0.19 to 0.22 mol/mmol Mg/Li, 1.77
332 to 2.83 mmol/mol Mg/Ca, and 9.15 to 13.7 $\mu\text{mol/mol}$ Li/Ca.

333 The average Mn/Ca and Fe/Ca values were 0.6 $\mu\text{mol/mol}$ and 2.5 $\mu\text{mol/mol}$, respectively
334 and the maximum values were 3.9 $\mu\text{mol/mol}$ and 53.9 $\mu\text{mol/mol}$, respectively. These ratios were
335 analyzed to determine potential contamination from ferromanganese overgrowths or detrital
336 grains. In all cases our observed Mn/Ca values fell well below the commonly applied upper limit
337 of 100 $\mu\text{mol/mol}$ (Ferguson et al., 2008; Boyle, 1983). Likewise, the maximum Fe/Ca value of
338 53.9 $\mu\text{mol/mol}$ is below the established upper limit of 175 $\mu\text{mol/mol}$ (Ferguson et al., 2008; Lea
339 et al., 2005; Bice et al., 2005). Both limits were developed for foraminifera, but in the absence of
340 equivalent limits for corals we found their values to be useful benchmarks. In addition we
341 followed the convention of Steinke et al. (2010) in testing for a correlation between Mg/Ca and
342 either Mn/Ca or Fe/Ca. In both cases no significant correlation was observed (Mg/Ca vs. Mn/Ca,

343 $R^2=0.003$; Mg/Ca vs. Fe/Ca, $R^2=0.01$). Our samples likely have low ferromanganese
344 contamination because the specimens are modern and therefore have had little time to develop
345 ferromanganese crusts.

346

347 **4. Discussion**

348

349 *4.1 Coral calcification models*

350

351 The information from the laser ablation microscale distribution of Mg coupled to Li
352 allows us to explore modes of calcification, including continual replenishment of the ECF,
353 Rayleigh fractionation, and ACC precursors or incorporation in organic materials.

354 If the ECF is entirely open, then Mg/Ca and Li/Ca should be uniform across skeletal
355 features in the absence of other driving processes; this prediction is inconsistent with our
356 observations. If Rayleigh fractionation describes the trace metal variability, then log-log plots of
357 the Me/Ca data should be linear (Gagnon et al., 2007). Both Mg and Li are preferentially
358 excluded from aragonite (i.e.: have a distribution coefficient $\ll 1$) so a Rayleigh model would
359 predict a positive correlation between the two elements, as observed in our laser data (Figure 3).

360 The positive slopes of the Mg/Ca versus Li/Ca data from DMC08a, DMC08b, and *Balanophyllia*
361 are consistent with a Rayleigh fractionation model following that of Gagnon et al. (2007) (Figure
362 3). The uncertainty within the DMC08 laser data cloud and the relative insensitivity of the model
363 to D_{Li} allows for greater than 10-fold variability of D_{Li} within the context of the model slope.

364 However most of the data are consistent with Rayleigh fractionation starting from seawater with
365 Li, Mg, and Ca concentrations of 25.9 $\mu\text{mol/kg}$ (Stoffynegli & MacKenzie, 1984), 52.7 mmol/kg

366 (Broecker & Peng, 1982), and 10.27 mmol/kg (Broecker & Peng, 1982), respectively. These
367 concentrations give rise to D_{Li} and D_{Mg} equal to 2.7×10^{-3} and 2.9×10^{-4} , respectively. If regions of
368 the coral are not controlled by Rayleigh fractionation then they should plot away from this
369 relationship. In *D. dianthus* there appears to be a break in slope at the highest Mg/Ca for each
370 coral (within COCs) that indicates that Rayleigh fractionation is not the only process controlling
371 coral Mg and Li distributions. This is consistent with previous studies of Sr/Ca and Mg/Ca in *D.*
372 *dianthus* that suggest a process other than Rayleigh fractionation dominates Mg/Ca behavior in
373 the COCs (Gagnon et al., 2007). Thus, our data are consistent with Rayleigh fractionation
374 occurring across the entire skeleton, with additional mechanisms for trace metal incorporation
375 occurring within COCs.

376 Rollion-Bard et al. (2009) conclude that Rayleigh-type fractionation is insufficient to
377 explain observed Li/Ca variability coupled with near constant δ^7Li values across the skeletons of
378 deep-sea corals. Since aragonite precipitation strongly discriminates against Li incorporation, the
379 Li concentration of the ECF is largely unaffected during Rayleigh fractionation. Therefore, δ^7Li
380 should remain constant (within analytical uncertainty) during Rayleigh fractionation while Li/Ca
381 varies according to the calcium concentration of the calcifying fluid. The results measured by
382 Rollion-Bard et al. (2009) are therefore not inconsistent with Rayleigh fractionation.

383 Rollion-Bard et al. (2010) suggested that ACC formation could account for observed
384 variation in trace metal ratios. ACC has been suggested as a precursor during biogenic
385 calcification (Addadi et al., 2003; Meibom et al., 2004). While evidence for ACC in coral
386 biomineralization remains inconclusive, this precursor phase does provide a possible mechanism
387 for high Mg since Mg is enriched in ACC (Addadi et al., 2003; Wang et al., 2009). If ACC is
388 indeed present during coral skeletogenesis, then ACC formation in COC regions could account for

389 the high Mg within COCs observed in this study. Alternatively, scleractinian coral skeletons are
390 known to contain a small amount of organic matrix (Cuif et al., 2003; Puverel et al., 2005).
391 Given the suggestion that organic matter may host Mg in coral skeletons (Finch & Allison,
392 2008), a second possibility is that COCs are rich in Mg-hosting organics. Consistent with this
393 explanation, Brahmi et al. (2010) find organic matter to be elevated in *Balanophyllia* COCs. Our
394 observed overall similarity in Mg/Ca and Li/Ca distributions suggest that whatever mechanism is
395 controlling Mg also controls Li, perhaps unsurprising given the similar ionic radius of the two
396 elements. These incorporation and variation pathways remain speculative, but nevertheless
397 provide possible mechanisms to elevate Mg and Li within COCs.

398 Data on Mg and Li incorporation in *D. dianthus* are complemented by several previous
399 studies, which document $\delta^{13}\text{C}$, $\delta^{18}\text{O}$, [U], and Sr/Ca variability across skeletal features (Adkins et
400 al., 2003; Robinson et al., 2006; Gagnon et al., 2007). Collectively, these patterns should
401 constrain vital effect mechanisms. Unfortunately the compositional effect of ACC, organic
402 matrix components, and several other competing mechanisms on multiple proxies remains
403 unquantified. It is our hope that future laboratory-based experiments will yield testable
404 predictions to compare different mechanisms to skeletal variability across COCs.

405

406 *4.2 Environmental controls on trace metal incorporation into deep-sea corals*

407

408 *4.2.1 Strategies for reducing the biases from vital effects*

409

410 Our new data corroborate previous studies which have reported significant trace metal
411 variation across skeletal features (Adkins et al., 2003; Sinclair et al., 2006; Cohen & Gaetani,

412 2006; Gagnon et al., 2007; Rollion-Bard et al., 2009; Brahmi et al., 2010). This distinct internal
413 variability of trace metals and isotopes means that separate subsamples of the same *D. dianthus*
414 have markedly different values (e.g. Smith et al., 2000; Figure 5). For example, if we use Mg/Ca-
415 to-temperature relationships for surface scleractinian corals (Mitsuguchi et al., 1996) and the
416 range of Mg/Ca values obtained by LA-ICP-MS on DMC08, we calculate corresponding
417 temperatures of 13°C and 2°C. Such a large range in apparent temperature obviously precludes
418 Mg/Ca as a paleothermometer and is in line with previous literature expressing concerns
419 regarding the Mg/Ca temperature proxy (Quinn & Sampson, 2002; Fallon et al., 2003;
420 Mitsuguchi et al., 2008; Finch & Allison, 2008, Gagnon et al., 2007). We find a negative Li/Ca-
421 to-temperature relationship, consistent with a previous study (Marriott et al., 2004a). However, it
422 exhibits extensive scatter (Figure 6i).

423 Analyzing specific features of the coral skeleton could avoid the complications presented
424 by vital effects. For example, sampling purely COC or alternatively fibrous material may allow
425 for repeatable measurements and development of a useful environmental proxy. Unfortunately,
426 we have found that such microsampling is difficult, if not impossible, on typical samples by the
427 methods described in this study. Laser ablation techniques can come close to analyzing single
428 architectural units, but with the 50 µm spot size used in our study we rarely sampled only COC
429 material even within the widest band (Figure 4). Previous work using NanoSIMS with spot size
430 0.4 µm has revealed Mg concentrations up to 10 times higher in COCs than fibrous regions
431 (Meibom et al., 2004). Targeting the fibrous bands may be somewhat easier because they tend to
432 be wider, however secondary COCs are common and sometimes difficult to identify. In
433 particular, our *Balanophyllia* samples showed broad gradients between the optical bands.

434 Additionally, the necessity of achieving a small spot diameter to analyze discrete parts of the
435 coral decreases the precision of the analytical measurement.

436 An alternative sampling strategy for overcoming the vital effect is to take many small
437 subsamples. Smith et al. (2000) used this approach to determine the relationship between $\delta^{13}\text{C}$
438 and $\delta^{18}\text{O}$ in *D. dianthus*, allowing them to determine the equilibrium values related to
439 precipitation from seawater. Adkins et al. (2003) used a similar strategy but subsampled a thin
440 section using a micromill, and were able to subsample finely enough to detect a break in the
441 slope between $\delta^{13}\text{C}$ and $\delta^{18}\text{O}$ within COCs. Gagnon et al. (2007) followed this strategy for trace
442 metals and were able to discern a similarly placed break in slope of Mg/Ca to Sr/Ca. However,
443 deciding whether any one subsample is purely COC or not requires the assignment of arbitrary
444 cut off values. In addition, this method is labor intensive and likely not to yield the most extreme
445 values within the skeleton. In our study, four randomly sampled 15-20 mg fragments of coral
446 DMC08 measured by solution ICP-MS yielded Mg/Ca values from 2.05 to 2.34 mmol/mol that
447 do not come close to capturing the full range of Mg/Ca variability detected by 50 μm spot sizes
448 using LA-ICP-MS.

449 Another approach is to find geochemical proxies less affected by biomineralization. Our
450 in situ LA-ICP-MS data show that Mg and Li behave similarly within the coral skeleton, with
451 high values in the COC regions and low values in fibrous areas. Since deep-sea scleractinian
452 corals appear to incorporate Mg and Li similarly, for reasons discussed above, the ratio of these
453 two elements has the potential to be less variable within the skeleton of a single coral. Laser
454 ablation results show less dramatic Mg/Li variation across the coral skeleton than for either
455 Mg/Ca or Li/Ca. For example DMC08a shows three distinct Mg/Ca peaks with the largest in the
456 central band which is 2.1 times greater than the local Mg/Ca minimum in the adjoining fibrous

457 layer. The Mg/Li ratio also shows three peaks, with the central one being the largest excursion,
458 however the magnitude is reduced to a factor of 1.4, and the two other peaks are reduced to an
459 even greater extent. In the second section, DMC08b, the broad Mg/Ca maximum from 500 to
460 1100 μm is not as well developed in Mg/Li and the entire section is characterized by $\sim 10\%$
461 Mg/Li variability on a $< 100 \mu\text{m}$ scale. This switch from few, discrete Mg/Ca peaks to higher
462 frequency Mg/Li variability is particularly pronounced in the *Balanophyllia* section (Figure 4c).

463 Mg/Li is less variable across each coral section, and that variability is more evenly
464 distributed throughout the skeleton. Therefore a randomly cut subsample is more likely to
465 contain a representative proportion of high and low Mg/Li and be closer the average coral value
466 than for Mg/Ca. For example, four separate subsamples comprising septal and thecal material of
467 the coral 48738 (*D. dianthus*) gave standard deviations for Mg/Ca, Li/Ca and Mg/Li of 22%,
468 18% and 5% respectively. As expected in cases where multiple subsamples were taken, the
469 Mg/Li ratios were more reproducible than were Mg/Ca (Figure 5). Since the Mg/Li variability is
470 distributed within the skeleton, $\sim 10 \text{ mg}$ samples are able to give a reasonable approximation of
471 the average value for that coral. To achieve the best possible estimate of the true ratio in each
472 whole coral we average all data points for separate subsamples. For corals with large skeletal
473 features, larger samples or multiple replicates should be made to provide the most representative
474 Mg/Li estimate.

475

476 *4.3 Environmental controls on Mg/Li ratios*

477

478 Having established that measured Mg/Li ratios are reasonably representative of the coral
479 skeleton, we now examine the external controls on the average coral Mg/Li. Our results from 34

480 coral individuals show a Mg/Li range of 0.20 to 0.35 mol/mmol. First we consider contamination
481 by ferromanganese crust overgrowths and detrital grains using Mn/Ca and Fe/Ca. Both ratios
482 were low, indicating that all corals were cleaned thoroughly and allowing us to rule out
483 contamination. Second, we explore the environmental controls on the trace metal incorporation
484 at the time of growth. To do this analysis we compare the Mg/Li ratio to parameters that may
485 influence Mg incorporation during precipitation, including salinity, carbonate ion concentration,
486 and temperature.

487 Recent studies have highlighted the potential control of salinity on Mg incorporation in
488 calcitic foraminifera (Kisakurek et al., 2008; Ferguson et al., 2008). Despite a wide salinity range
489 of 34.04 to 36.54 ‰, we do not see any correlation between salinity and Mg/Ca or Mg/Li in our
490 samples (Figure 6). The carbonate ion concentration has also been shown to play a controlling
491 role on Mg/Ca (Martin et al., 2002; Elderfield et al., 2006; Rosenthal et al., 2006) and on Mg/Li
492 in foraminifera (Bryan & Marchitto, 2008). We compare our data to $[\Delta\text{CO}_3]$, following the
493 convention of previous discussions (Elderfield et al., 2006; Rosenthal et al., 2006; Bryan &
494 Marchitto, 2008). This value is calculated using derived $[\text{CO}_3]$ values from in situ data (Key et
495 al., 2008) and equations from Broecker & Peng (1982), where $[\Delta\text{CO}_3] = [\text{CO}_3]_{\text{in situ}} -$
496 $[\text{CO}_3]_{\text{saturation}}$ and for aragonite, $[\text{CO}_3]_{\text{saturation}} = 120 * e^{(0.16(Z-4))}$ where Z is the depth in km. We
497 find no significant correlation between $[\Delta\text{CO}_3]$ and the Mg/Ca, Li/Ca, or Mg/Li ratios (Figure 6).
498 In all cases there is no improvement in correlation whether we differentiate by species or by
499 ocean basin.

500 The correlation between Mg/Li and temperature gives an R^2 value of 0.62 with all 34 data
501 points, and 0.82 when we exclude two visibly chalky coral samples (Figure 7a, b). The only
502 comparable Mg/Li data set comes from Bryan & Marchitto (2008), who measured the Mg/Li

503 ratio of the aragonitic foraminifera, *Hoeglundina elegans*, and found a similar temperature
504 correlation (Bryan & Marchitto, 2008; Figure 7b).

505 There does not appear to be any clear difference between data from solitary
506 (*Desmophyllum*, *Balanophyllia*, *Caryophyllia*, *Flabellum*, and *Trochocyanthus*) versus colonial
507 (*Enallopsammia* and *Lophelia*) coral groups (Figure 7a). Our in situ laser ablation data set
508 showed some subtle differences between the geochemistry of *D. dianthus* and *Balanophyllia*, so
509 secondly we consider only *Desmophyllum dianthus* and *Caryophyllia spp.* which are both within
510 the *Caryophyllidae* family and have similar skeletal architecture. Within this subset of corals
511 (n=25, 22 of which are *D. dianthus*) we show an almost identical correlation as when all species
512 are combined (Figure 7a, c). This result is not surprising since 25 of our 34 samples are either *D.*
513 *dianthus* or *Caryophyllia spp.* Again correlations of $R^2=0.68$ including all 25 and $R^2=0.82$
514 excluding one chalky sample demonstrate a strong control of temperature on the Mg/Li ratio of
515 these corals. As a third approach, we examine *Caryophyllidae* family corals by ocean basin to
516 see whether sample provenance plays a role in the Mg/Li-to-temperature correlation. There are
517 no obvious differences in the Mg/Li ratio from basin to basin demonstrating that the temperature
518 control is global in nature (Figure 7d). In addition, if we only consider live-collected or
519 radiocarbon dated samples in this analysis the resulting relationship to temperature does not
520 change.

521 We now consider factors that might account for the positive relationship between Mg/Li
522 and temperature. Our laser data show that COC regions have higher Mg/Li than fibrous regions
523 (Figure 4). Increasing the amount of COCs relative to fibrous regions could, therefore, raise the
524 overall coral Mg/Li ratio, essentially linear mixing between two endmembers (Sinclair & Risk,
525 2006). The most extreme measured Mg/Li values in DMC08 are 0.17 mol/mmol and 0.27

526 mol/mmol, but we observe values greater than 0.3 mol/mmol in some corals allowing us to rule
527 out this simple explanation.

528 As an alternative explanation, we examine the combined effect of Rayleigh fractionation
529 and temperature-driven changes to D_{Mg} and D_{Li} . For most of the coral skeleton, Mg/Ca and Li/Ca
530 ratios are consistent with Rayleigh fractionation. Since both Mg and Li are strongly
531 discriminated against during skeletal precipitation, Mg and Li concentrations are predicted to
532 remain nearly constant in the ECF regardless of the extent of precipitation. Thus Mg/Li ratios
533 minimize the effect of Rayleigh fractionation, allowing the expression of other biomineralizing
534 processes and/or environmentally driven changes to Mg/Li

535 If our Rayleigh interpretation is accurate, then the apparent relationship between Mg/Li
536 and temperature could be caused by a temperature dependence of D_{Mg} or D_{Li} . For this
537 mechanism to be consistent with our data, increasing temperature would need to cause an
538 increase in the ratio of D_{Mg} to D_{Li} by changing one or both parameters. There are few studies that
539 attempt to quantify D_{Li} or D_{Mg} in inorganic aragonite, and even fewer that explore their
540 temperature dependence. Regarding D_{Mg} , Gaetani & Cohen (2006) report a decrease in D_{Mg} with
541 increasing temperature over the appropriate range for inorganically precipitated aragonite. By
542 contrast, Reynaud et al. (2007) document a 1.6-fold increase of Mg/Ca over 10°C using cultured
543 shallow water corals. Similarly coral aragonite exhibits a D_{Li} decrease of a factor of ~2 over
544 10°C (Marriott et al., 2004a; Marriott et al., 2004b). We are not aware of any data that measured
545 D_{Li} in inorganically precipitated aragonite. For both D_{Mg} and D_{Li} the coral experiments were
546 performed in the range 20-30°C, and there are no empirical data at lower temperatures. Although
547 the data are conflicting there do appear to be temperature dependencies of D_{Mg} and D_{Li} in
548 inorganic and coral aragonite. Thus we can conclude that a temperature dependence of the ratio

549 D_{Mg}/D_{Li} may cause our observed Mg/Li to temperature correlation but we cannot ascertain the
550 controlling mechanism on these distribution coefficients.

551

552 *4.4 Mg/Li as a potential temperature proxy*

553

554 Empirical Mg/Li-to-temperature relationships for deep-sea corals suggest a control by
555 temperature. This ratio is readily measureable by laboratories with trace metal analytical
556 capabilities and can be analyzed on ~10 mg subsamples with a reasonable reproducibility
557 although larger-featured samples may be preferable on large corals. As such it is an appealing
558 candidate for a new paleothermometer in aragonitic deep-sea corals. The equation of the best fit
559 line of the *Caryophyllidae* family corals (*D. dianthus* and *Caryophyllia spp*) (Figure 8c) is:

560

$$561 \quad \text{Mg/Li (mol/mmol)} = 0.009 \pm 0.0008T (\text{°C}) + 0.19 \pm 0.006 \quad (4)$$

562

563 On average (excluding the one *Caryophyllidae* chalky sample), the corals exhibit 0.012
564 mol/mmol scatter from the Mg/Li trendline, equivalent to $\pm 1.3^\circ\text{C}$. When the one *Caryophyllidae*
565 chalky sample is included this scatter increases to 0.014 mol/mmol or $\pm 1.6^\circ\text{C}$. Some of this
566 range may be due to uncertainty in the ambient seawater temperature that was taken from global
567 data sets and not from the exact location of the coral.

568

569 **5. Conclusion**

570

571 Our results demonstrate that the microscale distributions of Mg/Ca and Li/Ca vary
572 similarly in two genera of deep-sea scleractinian corals. The Mg/Li ratio therefore exhibits lower

573 amplitude variation. Our data are consistent with trace metal incorporation that varies according
574 to Rayleigh fractionation with a potential role for trace metal association with ACC or organic
575 matter depending upon skeletal architecture.

576 Solution ICP-MS analyses from ~10 mg samples of 34 corals show less variability in the
577 Mg/Li ratio than Mg/Ca. Neither salinity nor carbonate ion concentration seem to control the
578 Mg/Li ratio. However, temperature correlates to Mg/Li with $R^2=0.62$ for all corals and with
579 $R^2=0.82$ for the *Caryophyllidae* family alone. We therefore suggest a temperature control on
580 Mg/Li, but note that paleotemperatures cannot be reconstructed to better than $\pm 1.6^\circ\text{C}$ if using the
581 *Caryophyllidae* trendline.

582

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593

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745 *of the National Academy of Sciences* 106, 21511-21516.

Table 1. All samples analyzed including their identifying information, biology, environmental parameters, and measured trace metal content. The completely averaged bulk solution Me/Me ratios are reported for every coral. Where more than one subsample was analyzed, the average values for each subsample are also presented. Collection temperature is assumed to be growth temperature since corals were either collected live or dated to be modern. Uncertainty in temperature is given in the table. Uncertainty in Mg/Ca, Li/Ca, and Mg/Li are ± 0.5 mmol/mol, ± 0.01 mol/mol, and ± 1.2 μ mol/mol, respectively, based on the maximum likely error (Figure 5).

Lab Label	Cruise Label	Genera	Species	Depth (m)	Latitude (°)	Longitude (°)	Salinity (‰)	[$\delta^{18}O$] (μ mol/kg)	Temperature (°C)	ΔT (°C)	Mg/Ca (mmol/mol)	Li/Ca (μ mol/mol)	Mg/Li (mol/mol)
DMC01	NBP0805-DR34-Dc-A-002	<i>Desmophyllum</i>	<i>dianthus</i>	869	-59.4	-60.4	34.36	-28.11	1.75	0.05	2.19	10.50	0.209
a											1.78	8.33	0.213
b											2.60	12.67	0.205
DMC05	NBP0805-DR35-Dc-A-001	<i>Desmophyllum</i>	<i>dianthus</i>	695	-59.4	-68.5	34.27	12.60	2.30	0.16	2.90	14.01	0.207
a											2.76	13.40	0.206
b											3.04	14.61	0.208
DMC08	IBP0805-TB08-BigBeauty-1	<i>Desmophyllum</i>	<i>dianthus</i>	816	-54.4	-62.1	34.29	-27.33	3.20	0.16	2.17	9.94	0.218
a											2.05	9.92	0.207
b											2.34	10.64	0.220
c											2.17	9.70	0.223
d											2.13	9.51	0.224
DMC09	NBP0805-DR40-Dc-A-1	<i>Desmophyllum</i>	<i>dianthus</i>	1323	-59.4	-68.6	34.65	-4.11	3.15	0.08	2.58	12.61	0.204
DMC10	NBP0805-TB04-Dp-A-3	<i>Desmophyllum</i>	<i>dianthus</i>	816	-54.4	-62.1	34.29	-27.33	2.05	0.16	2.67	12.51	0.214
DMC11	NBP0805-TB04-Dp-A-2	<i>Desmophyllum</i>	<i>dianthus</i>	816	-54.4	-62.1	34.29	-27.33	3.15	0.16	3.05	12.32	0.247
48738		<i>Desmophyllum</i>	<i>dianthus</i>	800	37.5	-25.5	35.50	51.80	10.25	0.85	2.31	11.11	0.206
a											1.77	9.20	0.192
b											2.81	13.13	0.214
c											1.98	9.65	0.205
d											2.66	12.45	0.213
47409		<i>Desmophyllum</i>	<i>dianthus</i>	659	-54.3	39.2	34.60		1.90	0.01	1.97	9.36	0.211
47413A		<i>Desmophyllum</i>	<i>dianthus</i>	421	-50.4	167.4	34.93	-36.20	7.30	0.30	3.02	11.36	0.265
80404		<i>Desmophyllum</i>	<i>dianthus</i>	390	35.3	-4.1	36.15		13.25	0.25	2.31	7.40	0.312
47407D		<i>Desmophyllum</i>	<i>dianthus</i>	549	-54.5	-129.5	34.29		2.75	0.15	2.63	12.09	0.218
REH_1	H13 DASS03#1	<i>Desmophyllum</i>	<i>dianthus</i>	1818	37.0	-60.0	34.98	24.75	3.50	0.50	3.06	13.87	0.220
a											2.99	13.64	0.219
b											3.13	14.09	0.222
REH_2	H13 DASS03#2	<i>Desmophyllum</i>	<i>dianthus</i>	1818	37.0	-60.0	34.98	24.75	3.50	0.50	1.81	8.37	0.216
a											1.83	8.61	0.213
b											1.79	8.13	0.219
REH_3	H13 DASS03#3	<i>Desmophyllum</i>	<i>dianthus</i>	1818	37.0	-60.0	34.98	24.75	3.50	0.50	2.58	11.59	0.223
a											2.45	10.80	0.227
b											2.70	12.38	0.218
AB132	CE0806 D19A-1a	<i>Desmophyllum</i>	<i>dianthus</i>	1720	58.0	-21.0	34.89		3.80	0.50	3.51	14.68	0.239
AB133	CE0806 D19A-2	<i>Desmophyllum</i>	<i>dianthus</i>	1720	58.0	-21.0	34.89		3.80	0.50	3.41	14.99	0.227
80358		<i>Desmophyllum</i>	<i>dianthus</i>	358	48.0	-8.0	35.53	-56.82	11.10	0.20	3.24	11.59	0.279
48739		<i>Desmophyllum</i>	<i>dianthus</i>	825	48.0	-7.0	36.54	-31.56	9.60	0.50	2.40	9.51	0.253
19249		<i>Desmophyllum</i>	<i>dianthus</i>	274	34.0	-120.0	34.04	-35.21	8.10	0.50	2.42	8.31	0.291
83383		<i>Desmophyllum</i>	<i>dianthus</i>	464	33.0	-128.0	34.11	-24.06	6.00	0.75	1.60	7.05	0.227
84818		<i>Desmophyllum</i>	<i>dianthus</i>	400	0.0	-90.0	34.72	-13.38	9.76	0.75	2.60	9.09	0.286
94069		<i>Desmophyllum</i>	<i>dianthus</i>	710	-31.0	-179.0	34.41	-69.47	7.40	0.50	2.61	8.59	0.304
DMC07	NBP0805-TB04-Dry Samples Modern Balan #1	<i>Balanophyllia</i>	<i>malouensis</i>	816	-54.4	-62.1	34.29	-27.33	3.15	0.16	1.91	9.13	0.209
a											1.80	8.64	0.208
b											2.03	9.61	0.211
DMC02	AB011-LMG0605 #20	<i>Balanophyllia</i>	<i>malouensis</i>	854	-53.5	-59.4	34.30		3.05	0.10	1.96	9.20	0.213
DMC03	QD024-LMG0605-Sin3#1	<i>Balanophyllia</i>	<i>malouensis</i>	120	-53.5	-61.5	34.21	-23.30	4.89	0.60	2.35	11.19	0.210
DMC04	JD030-LMG0605-Sin9-Box	<i>Balanophyllia</i>	<i>malouensis</i>	318	-54.3	-62.1	34.19	-25.34	4.51	0.25	2.15	9.62	0.223
1010252		<i>Carophyllia</i>	<i>ambrosia caribbeana</i>	785	28.2	-86.3	34.91	15.18	6.60	0.80	2.48	10.66	0.223
49020A		<i>Carophyllia</i>	<i>bertieriana</i>	201	12.5	-70.4	35.66	58.43	17.00	0.01	2.68	7.62	0.352
45923		<i>Carophyllia</i>	<i>ambrosia</i>	1318	7.4	-56.3	35.00	25.47	4.70	0.30	1.85	9.29	0.199
77019		<i>Enallapsammia</i>	<i>profunda</i>	494	30.2	-79.6	36.32	124.10	15.50	0.50	2.74	10.84	0.253
47531		<i>Enallapsammia</i>	<i>rostrata</i>	333	-51.0	162.1	35.00	-35.74	7.30	0.70	3.08	11.95	0.258
DMC06	NBP0805-TB04-Dry Samples Flabellum	<i>Flabellum</i>	<i>curvatum</i>	816	-54.4	-62.1	34.29	-27.33	3.15	0.16	1.73	8.78	0.197
a											1.86	9.33	0.199
b											1.60	8.22	0.194
61768B		<i>Trochocyanthus</i>	<i>rawsonii</i>	229	2.0	-47.0	35.37	87.37	10.00	2.00	3.29	10.39	0.317
46238I		<i>Lophelia</i>	<i>prolifera</i>	440	-25.0	-44.0	35.17	102.62	10.00	2.00	3.21	10.56	0.304

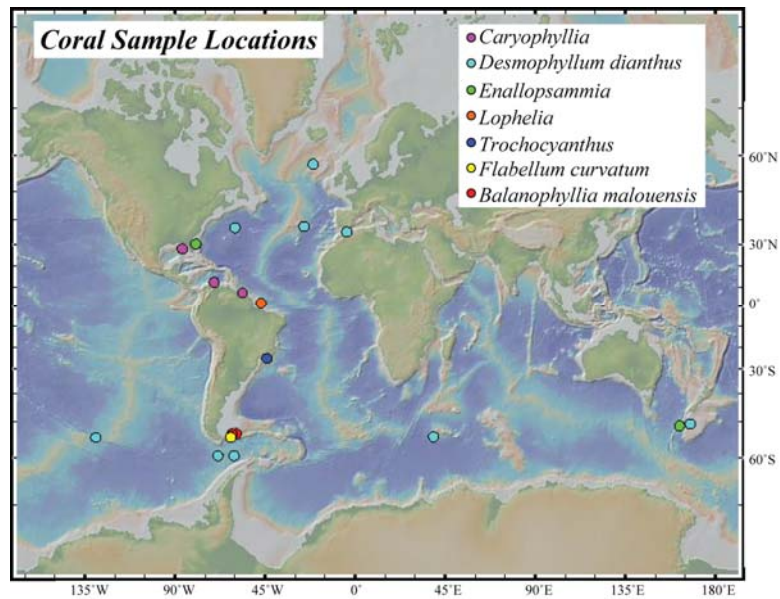


Figure 1. Location of coral sites colored by species. In cases where more than one species came from one site the location was colored according to the majority species found there (Table 1).

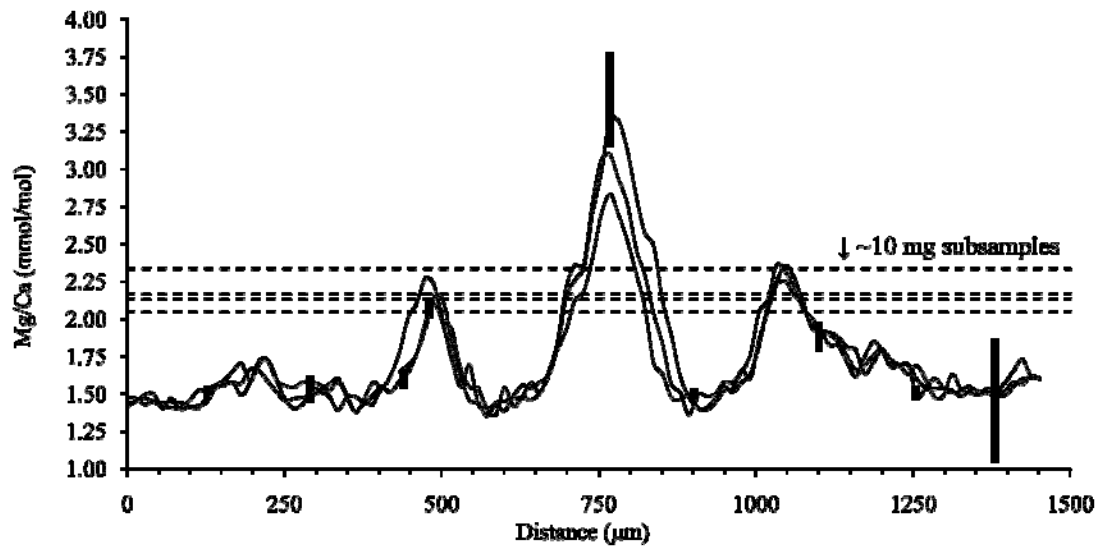


Figure 2. The Mg/Ca ratios analyzed in *D. dianthus* sample DMC08. Thin black lines are the closely spaced "long" laser lines run perpendicular to coral density strata. Thick black vertical bars are the Mg/Ca value from "short" laser lines. The length of the bar represents the uncertainty in those values. The four horizontal, dashed black lines are the Mg/Ca values obtained from four separate ~10 mg subsamples of the same coral analyzed by solution ICP-MS (Table 1).

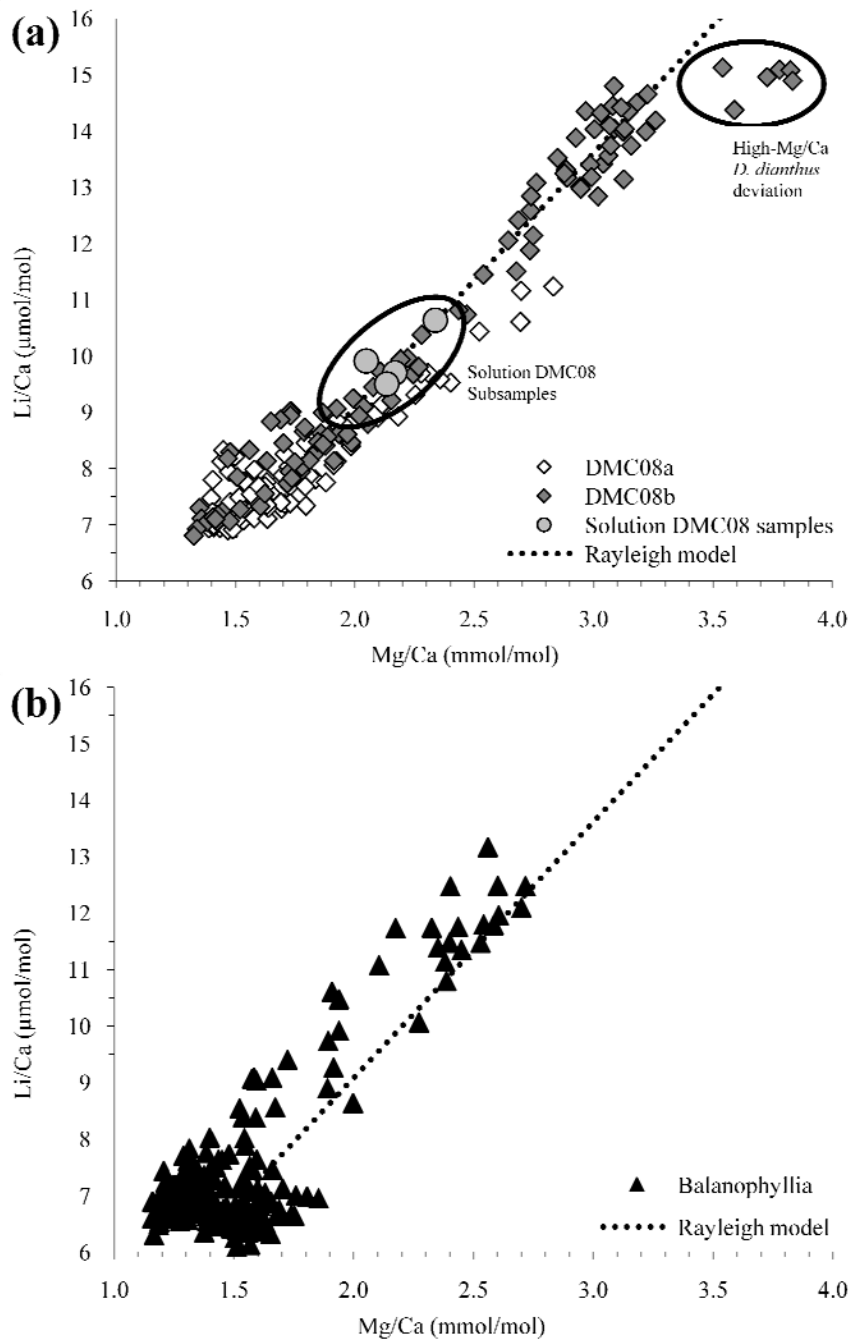


Figure 3. Laser ablation data for (a) DMC08a, DMC08b, and (b) *Balanophyllia*. Rayleigh models were calculated using D_{Li} of 0.00269 and D_{Mg} of 0.000292. The uncertainty on the solution DMC08 subsamples are smaller than the symbol size, and replicate laser lines have a reproducibility of $\sim 5\%$.

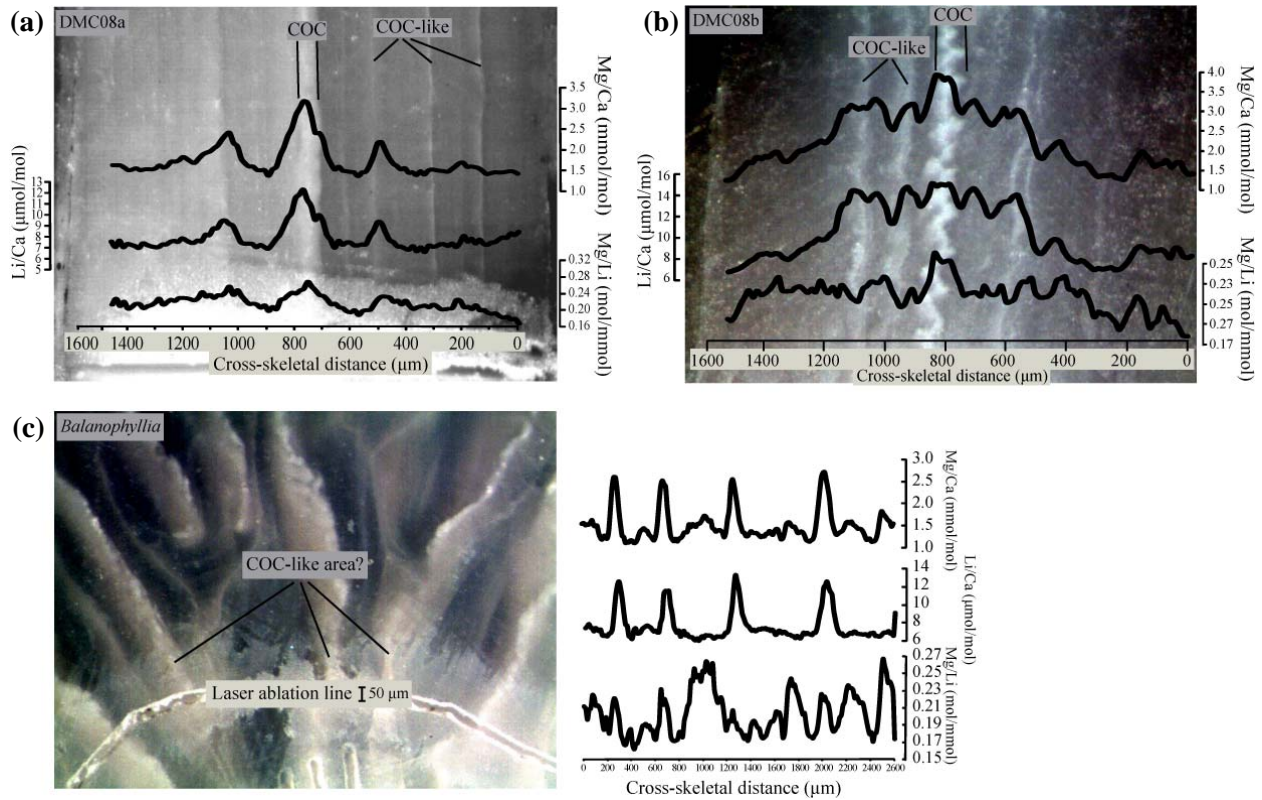


Figure 4. Laser ablation ICP-MS data overlaid on reflected light images of three coral sections. The first two sections (a) and (b) are from *D. dianthus* DMC08. The COCs are well developed and appear white in these sections. The x-axes indicate the location of the laser ablation line. (c) *Balanophyllia*. This laser ablation line was curved to stay perpendicular to the complex growth lines. In all cases, adjacent analyses indicate 5% error for Mg/Ca and Li/Ca ratios and 3% for Mg/Li. In addition regions which appear to exhibit similar geochemical behavior to COCs but which are not central to the skeleton are labeled “COC-like.”

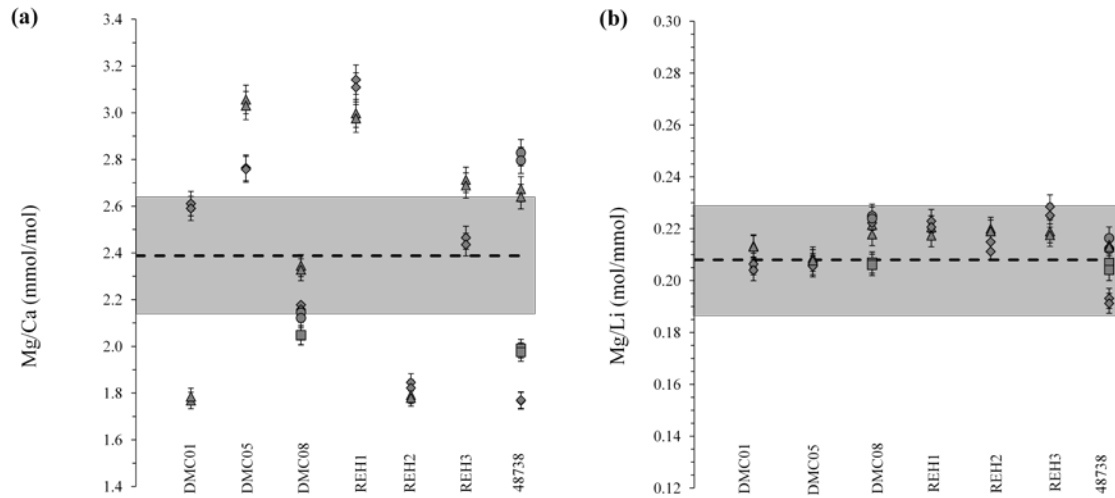


Figure 5. Intra-coral variability. **(a)** Mg/Ca and **(b)** Mg/Li. Only *D. dianthus* replicates are shown here. Within each coral individual symbols differentiate subsamples, while duplicate analyses of the same aliquot have the same symbol. Error bars for Mg/Li and Mg/Ca are 2%, each reflective of the standard deviation of reproducibility standards (S13). The average of plotted values is shown as a dashed line, and the gray box defines 10% deviations above and below the average.

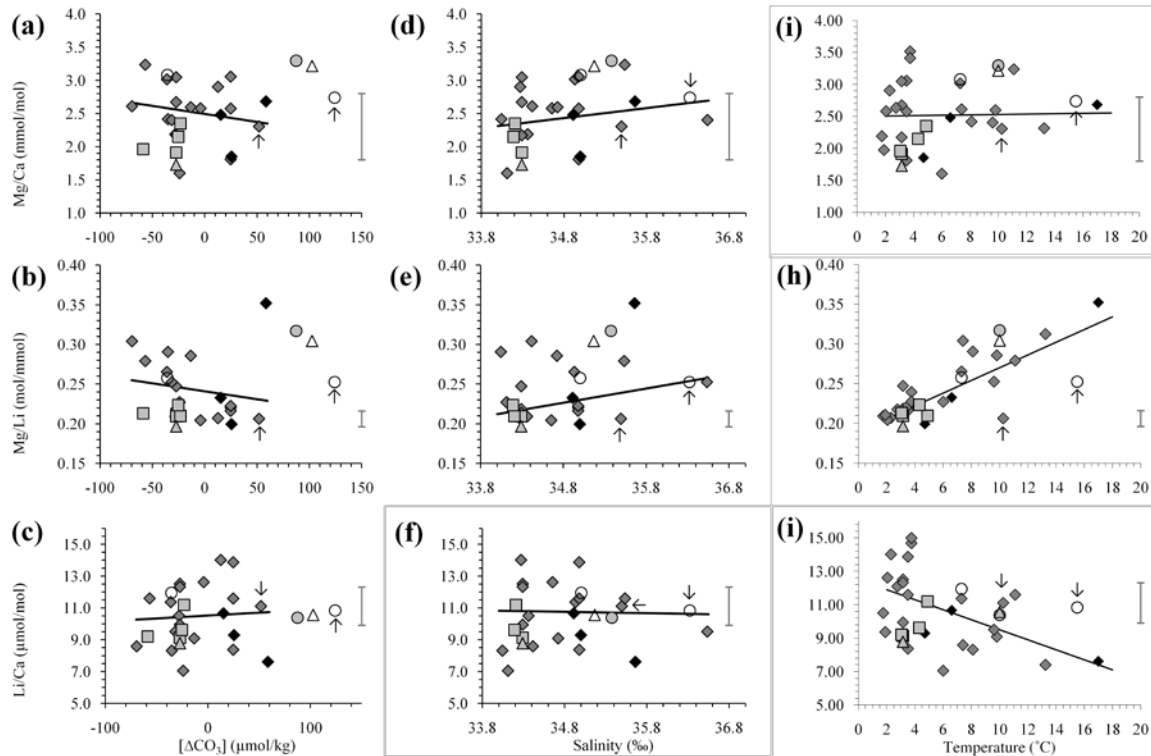


Figure 6. Subplots (a, b, c): effect of carbonate ion concentration on Mg/Ca, Mg/Li, and Li/Ca. Subplots (d, e, f): effect of salinity on the Mg/Ca, Mg/Li, and Li/Ca ratios. Subplots (g, h, i): effect of temperature on Mg/Ca, Mg/Li, and Li/Ca. Two chaly samples are shown with arrows. Symbols represent: *D. dianthus* (◆), *Caryophyllia* spp. (◆), *Balanophyllia* (■), *Flabellum* (▲), *Trochocyanthus* (●), *Enallopsammia* (○), *Lophelia* (△). Trendlines are given for the *Caryophyllidae* (◆ + ◆) data, for which the highest $[\Delta\text{CO}_3]$ value is 58.4. All $[\Delta\text{CO}_3]$ values are calculated using derived $[\text{CO}_3]$ values from in situ GLODAP data and equations from Broecker & Peng (1982). Gray bars at right indicate the maximum likely error in measurement, derived from data shown in Figure 5.

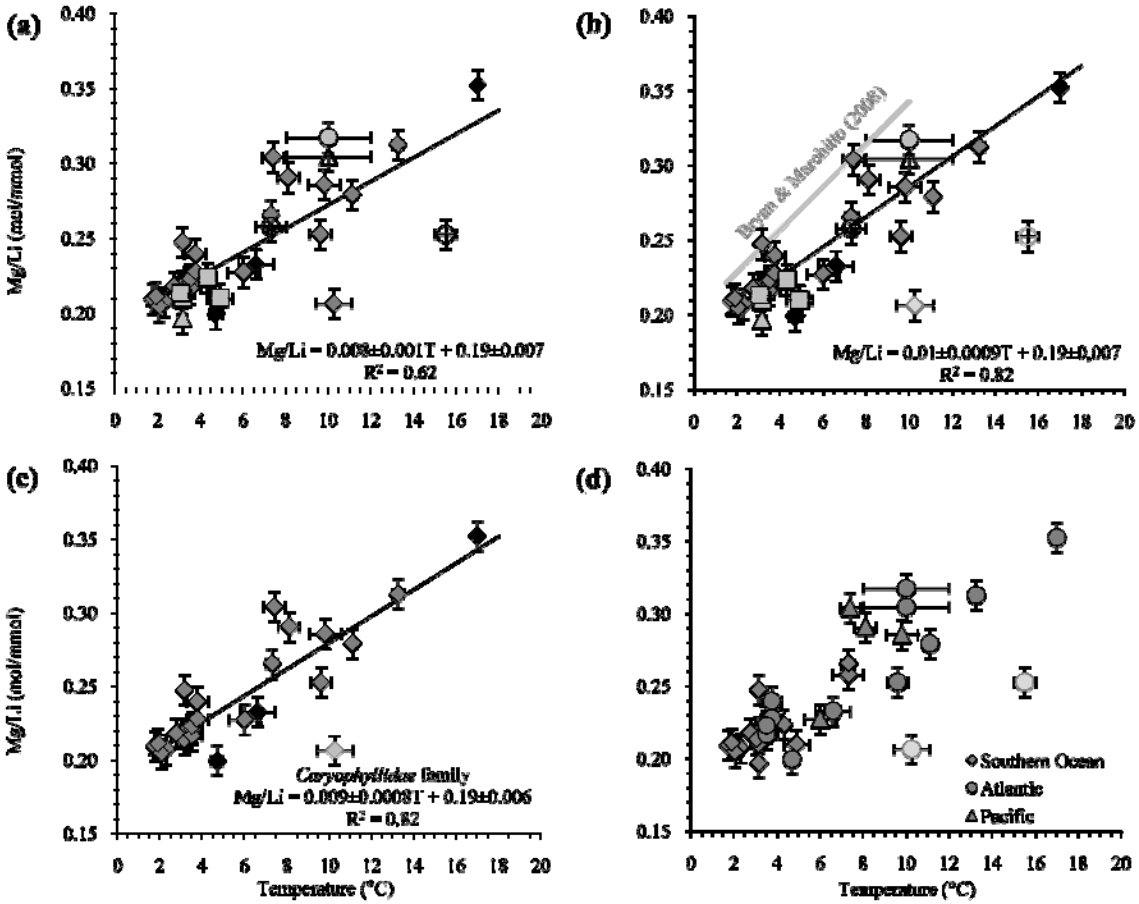


Figure 7. Mg/Li-to-temperature comparisons. Unless indicated in the subplot, symbols represent: *D. dianthus* (◆), *Caryophyllia spp.* (◆), *Balanophyllia* (■), *Flabellum* (▲), *Trochocyanthus* (●), *Enallopsammia* (○), *Lophelia* (△). Subplots display (a) Regression given for all data, separated by species; (b) Regression with the two visibly chalky samples removed (but shown faintly). The gray line represents the Mg/Li-to-temperature relationship given in Bryan & Marchitto (2008) for the aragonitic foraminifera *H. elegans*; (c) Regression for members of the *Caryophyllidae* family with one chalky sample shown faintly and not included in the regression (black line); (d) All samples, divided by major ocean basin, two chalky samples shown faintly. Temperature errors are derived from the GLODAP database (Key et al., 2004).