An improved boron isotope pH proxy calibration for the deep-sea coral *Desmophyllum dianthus* through sub-sampling of fibrous aragonite

**Joseph A. Stewart a,b,\*, Eleni Anagnostou a, Gavin L. Foster a**

a Ocean and Earth Science, National Oceanography Centre, University of Southampton. SO14 3ZH, UK.

a National Institute of Standards and Technology, Hollings Marine Laboratory, 331 Fort Johnson Rd, Charleston, SC, 29412, USA

\*Corresponding author. Tel: +1 843 725 4833, Email: Joseph.Stewart@noaa.gov

Highlights

* δ11B of deep sea coral is potentially a powerful tool to reconstruct intermediate depth seawater pH
* Microstructural heterogeneities result in uncertainty in pH derived from bulk skeletal δ11B
* We present δ11B and trace element composition of micro-sampled fibrous aragonite from *D. dianthus*
* δ11B of fibres lowest in Mg for each coral reveal a stronger, and better-defined dependence on ambient seawater pH
* *D. dianthus* is an archive of precise palaeo-pH (<0.1 pH units), providing that suitable sampling strategies are applied

Abstract

The isotopic composition of boron (δ11B) in marine carbonates is well established as a proxy for past ocean pH, however, its robust application to palaeo-environments relies on the generation of species-specific calibrations. Existing calibrations utilising the deep-sea coral (DSC) *Desmophyllum dianthus* highlight the potential application of this pervasive species to pH reconstructions of intermediate depth waters. Nevertheless, considerable uncertainty remains regarding the estimation of seawater pH from these bulk skeletal δ11B measurements, likely resulting from microstructural heterogeneities in δ11B of *D. dianthus*. To circumvent this problem, thus improving the reliability of the *D. dianthus* δ11B-pH calibration, we present a new δ11B calibration of micro-sampled fibrous aragonite from this species.

Modern coral specimens recovered from the Atlantic, Pacific, and Southern Oceans, micro-sampled using microdrilling, micromilling, and laser cutting extraction, were analysed for trace element (B/Ca, Mg/Ca, Sr/Ca, and U/Ca) and boron isotopic composition. We find the best calibration against the δ11B of borate in local ambient seawater (a function of pH and taken from hydrographic data sets; pH range 7.57 to 8.05) utilises δ11B measurements of fibres with likely slow growth rates and minimal contamination from adjacent microstructures (identified by low Mg/Ca) for each coral specimen. This new calibration exhibits a stronger, and better-defined dependence on ambient seawater pH compared to bulk coral δ11B; δ11B*fibre* = (**0.93** ±0.17) × δ11Bborate + (**12.02** ±2.63). We suggest that the majority of the variability in measured δ11B between replicate bands of fibrous aragonite from a *D. dianthus* specimen can be explained by small incorporation of non-fibrous aragonite and surface impurities during microsampling and growth rate effects. This study confirms the utility of *D. dianthus* as an archive of precise palaeo-pH (±0.07 pH units), provided that suitable sampling strategies are applied.

Keywords: *Desmophyllum dianthus*; deep-sea coral; pH; boron isotopes; trace element; fibrous aragonite

## Introduction

Intervals of increasing atmospheric CO2 concentration in the recent and geological past are generally associated with decreases in surface ocean pH following CO2 incursion into surface waters ([Doney et al., 2009](#_ENREF_19); [Hönisch et al., 2012](#_ENREF_42)). Consequently, there is a pressing need to understand the pathways by which the deep ocean has previously sequestered or released atmospheric carbon ([Barker et al., 2010](#_ENREF_9); [Burke and Robinson, 2012](#_ENREF_11); [Martínez-Botí et al., 2015](#_ENREF_50); [Skinner et al., 2010](#_ENREF_70); [Thiagarajan et al., 2014](#_ENREF_74)) and also the physiological and ecological response of marine calcifying organisms to lower pH; particularly under the other added pressures associated with anthropogenic CO2 emissions (*e.g.* warming and pollution; Orr et al., 2005; Hoegh-Guldberg et al., 2007). To this end, much attention has been focused on ocean carbonate system proxy development to allow reconstruction of seawater pH in the past. Of particular promise is the boron isotope pH-proxy based on the measurement of the boron isotopic composition (expressed as 11B, representing the 11B/10B ratio of samples relative to the 11B/10B of standard NIST SRM 951 in parts per thousand) of marine calcifying organisms ([Hemming and Hönisch, 2007](#_ENREF_34)). Despite this obvious utility, reliable use of this proxy is predicated on robust calibration of δ11B measurements of modern marine calcifying organisms to the pH of ambient seawater to account for commonly observed vital effects.

Previous studies have described the systematics of the δ11B pH-proxy ([Hemming and Hanson, 1992](#_ENREF_33); [Klochko et al., 2006](#_ENREF_44); [Zeebe and Wolf-Gladrow, 2001](#_ENREF_80)). Briefly, boron in seawater exists almost exclusively in two phases, boric acid (B(OH)3) and the borate ion (B(OH)4−), with an isotopic equilibrium fractionation between the two species such that B(OH)3 is enriched by 27.2‰ in the heavier 11B isotope (fractionation factor αB = 1.0272; Klochko et al., 2006). The relative abundance of these two boron species is pH dependent ([Dickson, 1990](#_ENREF_17)) and because the isotopic composition of the total boron in seawater (δ11Bsw = 39.61‰; Foster et al., 2010) must be maintained the δ11B of each species of boron is also a function of pH. δ11B measurements of marine carbonates show a strong similarity to the boron isotopic composition of the borate ion (δ11Bborate) forming the basis for a model whereby it is the charged borate ion that is predominantly incorporated into CaCO3 ([Hemming and Hanson, 1992](#_ENREF_33)). This understanding has recently been validated in aragonite using inorganic precipitation experiments ([Noireaux et al., 2015](#_ENREF_59)).

Boron isotope pH-proxy calibration work to date has predominantly focused on organisms calcifying in the surface ocean including planktonic foraminifera ([Henehan et al., 2013](#_ENREF_35); [Sanyal et al., 1996](#_ENREF_67); [Sanyal et al., 2000](#_ENREF_68)) and shallow-water corals ([Hönisch et al., 2004](#_ENREF_41); [Krief et al., 2010](#_ENREF_45); [Reynaud et al., 2004](#_ENREF_63)). Yet only limited attention has been given to δ11B-pH calibrations for deep/intermediate dwelling organisms such as benthic foraminifera ([Rae et al., 2011](#_ENREF_62)) calcitic octocorals ([Farmer et al., 2015](#_ENREF_21)), and aragonitic deep-sea corals (DSCs; Anagnostou et al., 2012; McCulloch et al., 2012b). Aragonitic DSCs are rich in boron (up to 10 times that of calcitic foraminifera; Blamart et al., 2007; Rae et al., 2011) and of sufficient mass to permit radiometric dating ([Cheng et al., 2000](#_ENREF_13); [Douville et al., 2010](#_ENREF_20); [Lomitschka and Mangini, 1999](#_ENREF_48)), making them an attractive substrate for recent intermediate water pH reconstructions.

Corals do not utilise ions derived directly from ambient seawater to build skeletal aragonite; instead corals calcify from a restricted extracellular calcifying fluid (ECF) in the sub-calicoblastic space ([Cohen and McConnaughey, 2003](#_ENREF_14)), which, although influenced by the external seawater environment, it is primarily controlled by the organism ([Comeau et al., 2013](#_ENREF_15); [Gagnon et al., 2012](#_ENREF_27); [Tambutté et al., 2011](#_ENREF_73)). One of the pathways by which the pH and chemical composition of this internal seawater pool is biologically modified is through enzymatic activity (Ca-ATPase pump) transporting Ca2+ into the ECF in exchange for H+ ions so as to optimise conditions for aragonite precipitation ([Al-Horani et al., 2003b](#_ENREF_3); [Allemand et al., 2004](#_ENREF_4); [Allison et al., 2010](#_ENREF_5); [Cohen and McConnaughey, 2003](#_ENREF_14); [McConnaughey, 1989](#_ENREF_52); [McCulloch et al., 2012b](#_ENREF_54)). Thus, the pH of the ECF is often greater than that of ambient seawater ([Al-Horani et al., 2003b](#_ENREF_3); [Venn et al., 2013](#_ENREF_77)), prohibiting estimation of ambient seawater pH derived from coral δ11B measurements by simply applying the theoretical relationship between δ11Bborate and pH ([Hemming and Hanson, 1992](#_ENREF_33)). Rather, robust, species-specific, coral δ11B calibrations to well constrained seawater pH must be implemented for reliable use of this proxy in coral (*e.g.* [D'Olivo et al., 2015](#_ENREF_15)).

The solitary, azooxanthellate, DSC, *Desmophyllum dianthus* is ideally suited for δ11B-pH reconstructions being both prevalent in the deep ocean and geographically and bathymetrically extensive (up to 70° of latitude and up to 2.5 km water depth; Figure 1; Stanley and Cairns, 1988; Cairns, 1994; Försterra et al., 2005; Thresher et al., 2011). Despite the palaeoceanographic potential of this taxon, recent geochemical studies suggest that DSCs contain considerable microstructural heterogeneity. Optically opaque centres of calcification (COC; Figure 2) generally yield low δ11B (*e.g.* COC in *Lophelia pertusa* and *Madrepora oculata* respectively up to 8‰ and 3‰ lower; Blamart et al., 2007) and U/Ca ratios, and high Mg/Ca ratios with respect to the adjacent fibrous aragonite that radiates from these internal early mineralisation zones ([Gagnon et al., 2007](#_ENREF_28)). It is postulated that variable mixing between these different microstructural components is the cause of the relatively large spread in the δ11B data used in previous attempts to characterise the δ11B-pH relationship for bulk *D. dianthus* ([Anagnostou et al., 2012](#_ENREF_6); [McCulloch et al., 2012b](#_ENREF_54)), limiting the utility of the bulk *D. dianthus* δ11B proxy to characterising palaeoceanographic-pH shifts (for example the ~0.15 pH unit shift in seawater from last glacial maximum to Holocene; Hönisch et al., 2008; Yu et al., 2010; Rae et al., 2011). It has been suggested that restricting δ11B analysis to a single microstructural component in DSC specimens, such as the fibrous aragonite, will help to minimise the scatter inherent within the bulk *D. dianthus* δ11B analyses ([Anagnostou et al., 2012](#_ENREF_6)), thus allowing the δ11B-pH calibration to be refined and enhance the palaeoceanographic utility of this species. The relatively slow growth-rates of DSCs (*D. dianthus* extension rate up to ~2 mm yr-1; Adkins et al., 2004) and sub-millimetre thickness of fibrous aragonite bands mean that attainment and processing of microstructurally uniform, sub-milligram, DSC samples presents clear challenges. Sampling must be fastidious enough to reject extraneous coralline phases from the fibrous aragonite whilst at the same time ensuring that sample sizes are sufficient for accurate δ11B analysis (ideally >0.1 mg CaCO3).

Here we present δ11B data from subsampled fibrous aragonite from the DSC *D. dianthus*. A combination of extraction methodologies are presented and critiqued as to their reliability in obtaining small (sub-milligram) fibre-only samples from DSC, whilst rejecting extraneous skeletal phases. We use these new δ11B data to explore the potential causes of microstructural heterogeneity in DSCs and to refine bulk *D. dianthus* δ11B-pH calibrations, thereby enhancing the precision and accuracy of intermediate water pH reconstructions.

## Methodology

### Sampling strategy

Coral samples for this study were obtained from the National Museum of Natural History (Smithsonian Institution, Washington, D.C.) and from the National Institute for Water and Atmosphere (NIWA), Greta Point, Wellington, NZ. The sample set covers a wide variety of depth habitat and latitudinal distribution (Figure 1; Table 1). Only samples collected alive were selected for analyses (*i.e.* with associated organic tissue; excluding specimen 94069) to ensure hydrographic data are as representative as possible. Many of the *D. dianthus* specimens used in this study were analysed for bulk septa δ11B by Anagnostou et al., ([2012](#_ENREF_6)) allowing results to be directly compared.

Where necessary calyx subsamples comprising one to two primary S1 septa (Figure 2 A; see Cairns, 1994 for structural nomenclature of solitary scleractinia) were detached using a <1 mm cut-width (diamond coated) rotary blade. The majority of surficial organic matter was mechanically removed from coral septa using a water-pick and a soft brush. Samples were then mounted in an epoxy resin and sectioned parallel to the primary growth axis, orthogonal to the plane of the S1 septa, using a low-speed saw (Buehler®) fitted with a 0.4 mm diamond wafering blade (Figure 2 A). Cut sample surfaces were polished using wetted silicon carbide fixed abrasive papers (Buehler®) to remove surface irregularities. A variety of microsampling approaches were applied here with the shared aim of exclusively sampling coral fibrous aragonite but with sufficient CaCO3 recovery to yield precise δ11B results. We now discuss these microdrilling, micromilling and laser cutting approaches in order of decreasing simplicity and speed of sampling (Figure 2 B, C and D).

#### Microdrilling (large fibre bands)

The surfaces of sectioned corals were wiped with methanol and thoroughly rinsed with MilliQ ultrapure water (18.2 MΩ) to remove dust and surface contaminants. Lines of fibrous aragonite (0.5 to 7 mm length along primary growth axis; 250 µm depth) were then directly microdrilled from cleaned, polished blocks of corals using the ESI New Wave MicroMill and a Brasseler H1621.31.008 Scriber drill bit (cut width 100 µm), leaving approximately 50 µm between the sample area and adjacent COCs or external coating. The resulting powered fibre sample was recovered in a droplet of MilliQ water using an acid cleaned pipette tip. Between samples, residual carbonate dust was removed using compressed air and the drill bit was cleaned in weak acid and rinsed thoroughly in MilliQ water to minimise sample cross-contamination.

Direct microdrilling of fibrous aragonite is desirable because, if performed successfully, this technique avoids powdering the unwanted COCs and external septum coating altogether. In practice however, the narrow diameter drill bit used is susceptible to slipping towards lines of weakness (such as the surface between coral and epoxy), potentially leading to unintentional entrainment of non-fibrous carbonate. Although this occasionally occurred, any samples where non-target COC/coatings were visibly incorporated were rejected. This led to microdrilling being reserved for samples with atypically thick zones of fibrous aragonite (*i.e.* >500 µm thickness).

#### Micromilling (intermediate fibre bands)

Polished and cleaned sample blocks possessing thinner zones of fibre (100 to 400 µm thick) required a more stable micromilling approach using a broader Brasseler 850 016 drill bit (cut width 800 µm) to provide drill bit stability. For this milling approach, a preliminary sampling trench was drilled into the epoxy at a depth of 300 µm overlapping the outermost 50 µm of the coral to completely remove the external coating and avoid epoxy contamination of the final milled powder. The sample block was then removed from the stage, thoroughly cleaned again with methanol and MilliQ to remove any drilling residue from the trench, before returning to the stage for sampling. The cleaned drill bit was then used to sample progressively towards the COCs (as finely as 20 µm slices) at a depth in the z-direction of 250 µm. This technique yielded far greater sampling precision for those *D. dianthus* samples with thin fibrous zones.

The progressive nature of sampling by the micromilling technique allowed multiple samples to be extracted across singles band of fibrous aragonite in two *D. dianthus* specimens (19168 and 83583). Radial extension rates in DSCs are complex ([Adkins et al., 2004](#_ENREF_1)) however, we assume that our sampling approximately 50 μm at a time from the fibres close the sample edge, to the fibres close to COCs represents a general transition from most recent to oldest fibre growth. For these two specimens, once multiple fibre samples had been removed, this progressive sampling was continued to extract a predominantly COC sample for comparison to adjacent fibres.

#### Laser cutting (small fibre bands)

Laser cutting of fibres was applied to samples with particularly narrow fibre zones (<100 µm thick). The principal advantage of this approach was it allowed even greater accuracy of sampling and enabled the recovery of whole pieces of aragonite (rather than milled/drilled powder) that result in minimal losses during subsequent chemical cleaning. Polished, surface-cleaned, coral thin sections, of uniform 200 µm thickness, were mounted in the New Wave UP193FX Laser Ablation device and individual pieces of fibrous aragonite were detached from the surrounding exterior coating and COC through laser drilling using a 50 µm spot size, 20 Hz, a fluence of ~6 J/cm2 and scanning speed of 5 µm/sec speed. This laser cutting technique is only recommended for very small samples, owing to its time consuming nature (24 hours to cut one sample) and associated expense.

### Analytical techniques

All analytical techniques used in this study were carried out at the University of Southampton and follow the protocols previously described in Foster ([2008](#_ENREF_23)). Extracted fibrous aragonite samples, weighing between 60 and 660 µg (typically 300 µg, ~12 ng of B; Table 2), were subject to oxidative cleaning in warm 1% H2O2 (buffered in ammonium hydroxide) to chemically remove remaining organic matter. The fibre samples were then given a weak acid leach (0.0005 M HNO3) to remove any re-adsorbed ions. Once cleaned, samples were dissolved in a minimal volume of 0.5 M HNO3 before centrifuging and transferring into clean PFA vials.

#### Elemental analysis

Prior to isotopic analysis, a small aliquot of each sample solution (~7%) was taken for elemental analysis using the Thermo Scientific Element 2 ICP-MS. Aliquots were diluted to an equal concentration of Ca and bracketed by well-characterised, matrix-matched synthetic standard solutions to yield B/Ca, Mg/Ca, Sr/Ca and U/Ca ratios for samples and to assess external reproducibility (2σ uncertainties of B/Ca = ± 5%, Mg/Ca = ± 2%, Sr/Ca = ± 2%, U/Ca = ± 4%). The JCp-1 reference material is a finely powdered coral (<250 µm; crushed by ball mill over 4 days; Okai et al., 2002) that is a similar size fraction to micromilled/drilled samples in this study. Eight identical JCp-1 powders (~5 mg in weight) and nine coarsely powdered samples from bulk *D. dianthus* specimen 19168 (superficially crushed by pestle and mortar; fragments up to ~500 µm) were analysed for trace element chemistry to assess the influence of cleaning; some received oxidative treatment as outlined above and some did not. These trace element values were used to assess the impact of organic matter removal on trace element values and measurement accuracy. Only Sr/Ca ratios measured in JCp-1 were found to be consistently different to those documented in the interlaboratory comparison study by Hathorne et al., ([2013](#_ENREF_32)), therefore a correction factor of +4.7% is applied to all Sr/Ca results in this study (Table 3). Ancillary Al/Ca, Fe/Ca, and Mn/Ca measurements (2σ respectively ± 8%, ± 7%, and ± 7%) were also made to ensure samples were free from contamination derived from detrital clay (*i.e.* Al/Ca < 200 µmol/mol) and oxide rich coatings (*i.e.* Fe/Ca < 100 µmol/mol and Mn/Ca < 50 µmol/mol).

#### Boron isotope analysis

The boron in the remaining sample solutions was separated from the carbonate matrix using 20 μl micro-columns containing Amberlite IRA 743 boron-specific anionic exchange resin ([Kiss, 1988](#_ENREF_43)). All boron must be recovered from columns to avoid isotopic fractionation, therefore following elution of the boron fraction, additional elutions were checked to ensure >99% of sample boron was recovered in the sample. Resultant solutions for boron isotope analyses used for the δ11B-pH calibration ranged from 3 to 33 ng of boron.

The δ11B of purified boron samples were measured in duplicate on a Thermo Scientific Neptune multi-collector (MC)-ICPMS against NIST SRM 951. The uncertainty on the average of the δ11B of the duplicates is dependent on boron intensity (long-term JCp-1 standard reproducibility for the laboratory while data in this study were collected, estimated using from voltage measured on 11B cup; n>40; typically 500 mV for 35 ppb B) and is calculated as,

(1)

A total procedural blank was also measured alongside batches of 8 samples throughout this study. A total procedural blank adjustment was applied to the coral fibre samples, however in all cases the impact on δ11B results was small (*i.e.* less than analytical uncertainty).

### Hydrographic data

For the δ11B-pH calibration we included coral samples only from localities which were in close proximity to well-constrained water sample measurements of alkalinity and dissolved inorganic carbon (DIC) in the CDIAC ocean carbon system database; (<http://cdiac.ornl.gov>). The *D. dianthus* specimens lacking reliable hydrographic data were reserved for the assessment of microstructural δ11B variability. The hydrographic alkalinity and DIC data for each coral shown in Table 1 were used in conjunction with temperature, salinity, and nutrient data to calculate pH (total scale), carbonate ion concentration, and aragonite saturation state at each coral location in the Seacarb package in R ([Gattuso et al., 2015](#_ENREF_29)) using the dissociation constants from Lueker et al. ([2000](#_ENREF_49)) and the boron/salinity relationship of Lee et al. ([2010](#_ENREF_46)). Calculated seawater pH values were then used to calculate ambient δ11Bborate rearranging equation (2) ([Zeebe and Wolf-Gladrow, 2001](#_ENREF_80)) to give equation (3), where αB is the fractionation factor between boric acid to borate (1.0272; Klochko et al., 2006) and p*K*B\* is the dissociation constant of the two boron species calculated using the Seacarb package in R with site-specific temperature, salinity, and pressure data.

(2)

(3)

We estimate the impact of anthropogenic CO2 invasion on pH estimates (*i.e.* Sabine et al., 2004), through subtraction of the accumulated anthropogenic contribution to DIC averaged across a 1° box (±50 m of coral depth) in the GLODAP gridded database (http://cdiac.ornl.gov/oceans/glodap), to be small (typically <0.01 pH units) and can therefore be omitted (Note: 1° box for site 47409 is extended to ±100 m to account for discrepancy between coral sample depth and hydrographic bottle depth). The same 1° boxes from the GLODAP gridded database were used to estimate the local pH variability at each coral locality; calculating multiple pH estimates from each paired value of alkalinity and DIC within the defined box. This method yielded uncertainty in ambient pH of < ±0.05 pH units (2σ) that manifest in uncertainty on δ11Bborate of < ±0.24‰ (Table 1).

## Results

### Cleaning experiments

Trace element results of peroxide treated and untreated coarsely powdered bulk 19168 *D. dianthus* samples and finely powdered *Porites* coral JCp-1 are shown in (Table 3). Trace element results replicate well within treatments (up to only 3% RSD), suggesting that both of these coarse and finely powdered carbonates were successfully homogenised. No significant difference was observed between trace element values of cleaned and uncleaned *D. dianthus* when coarsely powdered (pestle and mortar; low surface area; low cleaning efficacy), whereas B/Ca and Mg/Ca ratios in finely powdered JCp-1 (ball-milled for 4 days) were seen to decrease by approximately 25% (−113 and −990 µmol/mol respectively) upon removal of organics. Oxidative cleaning was found to have minimal impact on Sr/Ca and U/Ca values in either powdered carbonate.

### δ11B*fibre* *vs.* δ11Bborate

In Figure 3 A we show previous carbonate δ11B measurements regressed against the δ11Bborate ambient seawater (a function of seawater pH; Equation 3) including bulk *D. dianthus* δ11B (δ11B*bulk*; Anagnostou et al., 2012; McCulloch et al., 2012b), surface corals ([Hönisch et al., 2004](#_ENREF_41); [Krief et al., 2010](#_ENREF_45); [Reynaud et al., 2004](#_ENREF_63); [Trotter et al., 2011](#_ENREF_76)), and calcitic bamboo octocorals ([Farmer et al., 2015](#_ENREF_21)). The δ11B of bamboo corals falls close to the 1:1 line between carbonate δ11B and δ11Bborate suggesting that these biocalcifiers simply incorporated borate ion from seawater with little modification ([Farmer et al., 2015](#_ENREF_21)). This is in contrast to bulk *D. dianthus* values that are relatively enriched in 11B with respect to the 1:1 line (by approximately +11‰); more than 2‰ heavier than shallow-water coral taxa growing at equivalent pH (Figure 3; Anagnostou et al., 2012; McCulloch et al., 2012b).

Our new δ11B*fibre* and trace element data are shown in Table 2 and δ11B*fibre* data are compared in detail to the δ11B*bulk* data of [Anagnostou et al. (2012)](#_ENREF_6) in Figure 3 B. Comparative regression analysis for each of these calibrations are summarised in Table 4. The majority of δ11B*fibre* values fall within 1‰ of δ11B*bulk* values where identical specimens were sampled and we find no consistent δ11B offset between these two data sets. Rather, we document average δ11B*fibre* values that are both more than 0.5‰ higher in some cases (samples 19168, 47409, 94069) and lower in others (samples 82065, 48740, 47407) than their respective bulk measurement. Like δ11B*bulk* (R2 = 0.42) our new *D. dianthus* δ11B*fibre* data are well correlated with δ11Bborate if considered as solitary measurements (R2 = 0.48), with the sensitivity to pH (inferred from the slope of the regression; 0.72) nearly identical to that of δ11B*bulk*. We note however that if the mean of replicate δ11B*fibre* data for each coral sample is taken, the correlation with δ11Bborate improves greatly, with δ11Bborate explaining 69% of the variance in the average δ11B*fibre* data. Moreover, filtering these δ11B*fibre* data so as to only include the δ11B*fibre* sample replicate with lowest measured Mg/Ca (denoted by † symbol in Table 2) further strengthens the correlation with δ11Bborate (R2 = 0.83); yielding a greater sensitivity to pH (steeper slope that is closer to 1) and narrower confidence intervals at the 95% level (Figure 3 B).

### Spatial chemical variability in individual fibres

Systematic changes in δ11B*fibre* and trace element variability (B/Ca, Mg/Ca, Sr/Ca and U/Ca) during coral growth were further assessed, using the multiple samples taken across single bands of fibrous aragonite, in *D. dianthus* specimen 19168, possessing the thickest bands of fibrous aragonite and specimen 83583 from the lowest pH site (Figure 4). Sample 1 from 19168 shows decreases in δ11B and Mg/Ca (of 0.9 ‰ and 0.15 mmol/mol respectively) sampling fibres progressively away from the COCs while Sr/Ca and U/Ca both increase (by 0.63 mmol/mol and 0.15 μmol/mol respectively). The other two specimens measured in this manner (19168 Sample 2 and 83583) show more consistent intra-fibre chemistry, however similar trends are also present in these specimens, particularly δ11B and U/Ca in sample 83583, with more muted change found in 19168 Sample 2 (Figure 4).

The sampling methods described in this study are not of sufficient spatial resolution to allow the acquisition of a “pure” COC sample containing no fibrous aragonite component. However, the purest COC sample obtained was from the centre of 19168 Sample 1 that possessed the most prominent COC. This COC sample yielded higher B/Ca (+20%; +42 μmol/mol) and Mg/Ca (+80%; +1.10 mmol/mol), similar Sr/Ca (only 10% or 0.77 mmol/mol lower), and lower U/Ca (−30%; −0.84 μmol/mol) relative to adjacent fibrous aragonite; a finding consistent with previous *D. dianthus* microstructural trace-element studies ([Anagnostou et al., 2011](#_ENREF_7); [Blamart et al., 2007](#_ENREF_10); [Gagnon et al., 2012](#_ENREF_27)). The less prominent COC sampled from 83583 does not exhibit any clear chemical differentiation between COC and adjacent fibrous aragonite (with the exception of lower U/Ca in the COC), however the narrower COC sampled here will represent a significant mixing with the fibrous aragonite end member. Despite the disparate trace element chemistry between COC and fibres we observe relatively small differences (<0.9‰; *c.f.* *Lophelia pertusa* data by Blamart et al., 2007) between δ11B in fibres and COCs using our sampling strategies (Figure 4; Table 2).

### Trace elements in *D. dianthus* fibres

We compare our new *D. dianthus* δ11B*fibre* data to measured trace element ratios for each dissolved fibre sample (Figure 5). *D. dianthus* δ11B*fibre* data are positively correlated with B/Ca and Mg/Ca and negatively correlated with Sr/Ca (albeit with a relatively weak correlation) and U/Ca. We note that the strength and direction of trace element correlation to δ11B*fibre* appears to be dependent on the partition coefficient of each trace element into aragonite (*D*X = (X/Ca)Aragonite/(X/Ca)Seawater): Elements that more preferentially remain in solution during the calcification process (*i.e.* B and Mg; *D* values <<1) yield positive correlations. Conversely, elements favourably incorporated into the aragonite lattice (*i.e.* U; *D* value > 1) are negatively correlated with δ11B*fibre*, while elements incorporated close to unity with seawater (*i.e.* Sr; *D* value only slightly above 1) show the poorest degree of correlation.

There is no systematic trace element or isotopic offset between *D. dianthus* fibrous aragonite sampling techniques with the exception of samples collected via the laser cutting technique that yield anomalously high B/Ca values (+300 µmol/mol) compared to milled/drill samples of similar δ11B (Figure 5; Table 2; compare samples from 19249).

## Discussion

### An improved δ11B-pH calibration for *D. dianthus*

In line with previous DSC studies ([Anagnostou et al., 2012](#_ENREF_6); [McCulloch et al., 2012b](#_ENREF_54)), our total dataset yields a skeletal coral δ11B to δ11Bborate slope of <1 (Table 4). Together, our new *D. dianthus* data support suggestions that the pH of the calcifying fluid in this DSC is biologically upregulated ([Anagnostou et al., 2012](#_ENREF_6); [Blamart et al., 2007](#_ENREF_10); [McCulloch et al., 2012a](#_ENREF_53)), promoting calcification and partially mitigating the effects of the low aragonite saturation states (ΩArag) found in the deep waters these taxa inhabit ([Guinotte et al., 2006](#_ENREF_30)).

If our new fibre data are considered as solitary measurements, these data appear similar to absolute δ11B values of bulk septa and exhibit a similar sensitivity to pH (Figure 3; Table 4). The calibration by linear regression of δ11B*fibre* to δ11Bborate has a comparable spread in its residuals (R2 = 0.48) to that of δ11B*bulk* (R2 = 0.42) suggesting that δ11B*fibre* data provide seemingly little improvement to the original δ11B*bulk*-pH calibration. Yet, if replicate δ11B*fibre* data for each coral sample are averaged the δ11B*fibre* calibration (R2 = 0.69) reveals a marked improvement upon the δ11B*bulk* calibration. Once averaged, each point on the average δ11B*fibre* calibration represents a different mass of averaged fibre CaCO3 (between 0.25 and 1.44 mg). An alternative regression, weighted by fibre sample mass, further improves the calibration (R2 = 0.76) with minimal impact on the slope or intercept of the regression.

The benefits to the δ11B-pH calibration of microsampling fibrous aragonite in *D. dianthus* are clear, yet the approach of averaging δ11B*fibre* data described above still encompasses all of the chemical variability within fibre replicates; the source of which we explore in detail in Section 4.2. Low Mg/Ca ratios in *D. dianthus* substructures are indicative of the purest fibrous aragonite ([Gagnon et al., 2007](#_ENREF_28)) and/or slow growth rates (*e.g.* inorganic aragonite; Gabitov et al., 2008; and corals; Montagna et al., 2014). Therefore, when we take full advantage of our microsampling approach and filter our δ11B*fibre* data to include only those replicates with the lowest Mg/Ca value (denoted by † in Table 2), we improve the calibration further with δ11Bborate explaining 83% of the variance and suggesting a stronger sensitivity to pH than δ11B*bulk* measurements (*i.e.* greater slope). It is presumed the lowest Mg/Ca fibre samples are least contaminated by Mg-rich COCs and were formed closest to thermodynamic equilibrium. We note that the majority of these low-Mg fibre replicates also correspond to high U/Ca further suggesting that these samples are largely free from COC contamination.

This low-Mg δ11B*fibre* calibration for pH is a considerable improvement upon the δ11B*bulk* calibration (residual standard error of 0.44 and 0.93 respectively); therefore this clearly enhances the utility of the *D. dianthus* calibration permitting yet smaller perturbations in intermediate water pH in the past to be resolved (*e.g.* ~0.15 pH unit shift from last glacial maximum to Holocene changes in seawater pH; Hönisch et al., 2008; Yu et al., 2010; Rae et al., 2011). Application of our new low-Mg fibre calibration to typical coral δ11B measurements of ~26‰, the 95% confidence intervals suggest that seawater pH can be reconstructed to within ±0.07 pH units (*c.f.* δ11B*bulk* ±0.34 pH units). Based on these results, for the most accurate δ11B-pH estimates using this taxon, we suggest one of the following approaches are followed: (i) Multiple fibre replicates are taken (n>3) for paired δ11B and trace element analyses so that samples with slow growth rates and/or minimal COC contamination (low Mg/Ca, high U/Ca) can be selected for pH estimation based on equation 4;

δ11B*low-Mg fibre* = (**0.93** ±0.17) × δ11Bborate + (**12.02** ±2.63) (4)

(ii) Alternatively, if elemental data are unavailable, powders from these multiple fibre replicates should be combined to make an average fibre sample (ideal total sample mass >0.7 mg; average of this study) and pH should be estimated using the (sample mass weighted) average fibre regression (equation 5).

δ11B*fibre* = (**0.75** ±0.17) × δ11Bborate + (**14.69** ±2.57) (5)

Although low-Mg δ11B*fibre* data have clear benefits when it comes to reconstructing accurate seawater pH in the following section we seek further explanation for δ11B*fibre* variability within single coral calyxes.

### Causes of δ11B variability within fibrous aragonite

Measurements of δ11B*fibre* and trace element composition across single bands of fibrous aragonite in *D. dianthus* specimen 19168 Sample 1 (and to a lesser extent 19168 Sample 2 and 83583) show that fibre chemistry is not homogenous across a single band of fibrous aragonite (Figure 4). Rather, trace element and isotopic data show trends of decreasing δ11B and Mg/Ca and increasing Sr/Ca and U/Ca towards the exterior of the septum. Possible sources of δ11B*fibre* and trace element variation include (i) site specific differences in hydrography and food supply during growth, (ii) contamination of fibrous aragonite samples with organic matter, COC, or coating during sampling, (iii) Rayleigh fractionation, and/or (iv) growth rate induced changes in skeletal chemistry during calcification. Below we discuss the extent to which these possible sources of variability explain our trace element and isotope data from *D. dianthus* fibres.

#### Changes in hydrography or food supply during growth

Cold-water corals are known to change their distribution and calcification rates depending on temperature and/or nutrient availability ([Dodds et al., 2007](#_ENREF_18); [Mienis et al., 2014](#_ENREF_55); [Mortensen et al., 2001](#_ENREF_58); [Roberts et al., 2006](#_ENREF_64)), hence it is possible that *D. dianthus* δ11B is influenced by hydrographic variables other than pH alone. We therefore explore our δ11B*fibre* data for further correlation with ambient seawater conditions. To this end, we regress residuals of the ordinary least squares regression of δ11B*fibre* *vs.* δ11Bborate against hydrographic variables such as temperature, salinity, and [PO4-], yet we find no significant relationships (residuals *vs.* Temperature, R2 < 0.01; Salinity, R2 =0.01; [PO4-], R2 =0.01; Supplementary Figure S 1). Further, the amount of variance observed for replicate δ11B*fibre* results in each *D. dianthus* sample bears no clear relation to the variance in the available hydrographic data (Supplementary Figure S 2).

The ability of a DSC to upregulate internal pH could be limited by metabolic energy availability ([Al-Horani et al., 2003a](#_ENREF_2)), hence food supply may in part drive pH upregulation and therefore some of the δ11B*fibre* variability. Yet, we find no clear evidence for food supply (either amount or intra-annual variability estimated from site specific particulate organic carbon flux) driving discrepancies between δ11B*fibre* replicates (See Supplementary information).

#### Contamination of fibres with organics, COC, or surface coating

It was important that organic phases were effectively eradicated from biogenic carbonate samples in this study to ensure these trace-element-rich phases were excluded (*i.e.* high Mg; Barker et al., 2003), and to preserve the micro-columns used for boron purification. The importance of effective oxidative cleaning on coral samples is clear when we compare trace element data from the cleaned and uncleaned JCp-1 standard (Table 3). Our warm buffered peroxide treatment greatly reduced resultant B/Ca and Mg/Ca ratios of this finely powdered standard (*e.g.* Holcomb et al., 2015), suggesting that coral organic phases are potentially rich in these two trace elements and that oxidative cleaning must be applied to extract trace element ratios representative of the aragonite phase. We note that trace element ratios of JCp-1 replicates cleaned in this manner are remarkably consistent (*i.e.* within instrumental analytical uncertainty) suggesting that the cleaning method detailed above yields near-complete and reproducible removal of organics matter in samples.

All micromilled and microdrilled samples in this study were finely powdered (similar surface area to the JCp-1 standard) and therefore we assume successfully eradicated of organics. The lower B/Ca ratios measured in cleaned micromilled/drilled *D. dianthus* samples in this study (300 µmol/mol) compared to *in situ* analyses (e.g. laser-ablation where pre-cleaning was not applied; Montagna et al., 2005) in this species are consistent with a boron rich organic phase being removed. The controls on the boron isotopic composition of coral organic matter are likely complex. Although light 10B is preferentially adsorbed onto organic surfaces (~25‰ lower than the ambient fluid at pH 8; Lemarchand et al., 2005), the study by [Stoll et al. (2012)](#_ENREF_72) on coccolithophorids points to membrane permeable boric acid (11B enriched with respect to seawater) as the major source of boron in organic cells. Despite large reductions in B/Ca, results of a recent interlaboratory comparison study reveal only a small reduction in δ11B of JCp-1 powder (−0.2‰) from oxidative cleaning ([Gutjahr et al., 2014](#_ENREF_31)), also suggesting that, at least in this case, boron in coral organic matter is isotopically heavier than skeletal carbonate. Although well homogenised, the bulk 19168 sample, superficially crushed by pestle and mortar (large fragments), yielded no such reduction in B/Ca or Mg/Ca after cleaning (Table 3), suggesting that the low surface area of this test sample hindered the efficacy of organic matter removal. Persistence of organic matter in coarse samples likely explains higher B/Ca values measured in *D. dianthus* fibres sampled via the laser cutting methodology in this study (Figure 5) that were similarly cleaned as larger intact fragments. Presence of residual organic phases that are both rich in boron and isotopically heavy may explain why in some cases bulk δ11Bvalues for *D. dianthus* samples (also homogenised by pestle and mortar; Anagnostou et al., 2012) are higher than δ11B*fibre* measurements in this study. The laser cut δ11B*fibre* value of sample 19249 falls within the range of microdrilled aliquots of the same sample (only 0.3‰ apart), suggesting that, in this case, δ11B*fibre* values are impacted little by choice of microdrilling or laser cutting protocols. While these two laser-cut data points cannot account for the δ11B variability in the majority of samples that were micromilled/drilled and effectively oxidised, this further suggests that samples collected by laser cutting be used with caution.

If *D. dianthus* corals are chemically similar to the DSC *Lophelia pertusa*, then varying proportions of COCs ([Blamart et al., 2007](#_ENREF_10)) will cause significant shifts in measured δ11B*fibre* values. Special care was made to avoid COCs with the fibrous aragonite sampling strategies described above; certainly in comparison to previous bulk sampling approaches ([Anagnostou et al., 2012](#_ENREF_6); [McCulloch et al., 2012b](#_ENREF_54)). Additionally, our stringent quality control of sampling protocols ensured that any sample where it was visible that COC had been mistakenly incorporated was rejected prior to analysis. Nevertheless, because filtering of δ11B*fibre* data based on Mg content has significant impacts on the calibration with pH, it is conceivable that small amounts of COC are unavoidably incorporated into the micro-samples simply due to averaging of unseen skeletal material in the z-direction and the possibility that minor COCs were not visible on the polished surface. In the *D. dianthus* specimen 19168 these opaque COC bands were found to be similar to adjacent fibre samples differing by less than 0.9‰ in δ11B composition, despite the large trace element shifts mentioned above (Figure 4). Although analytical techniques are different and our spatial sampling resolution is greatly averaged in comparison to ion microprobe analysis, we suggest that at least for this coral specimen, the difference in δ11B between COCs and fibres in *D. dianthus* is far smaller than that measured in *Lophelia pertusa* ([Blamart et al., 2007](#_ENREF_10)). In this regard, we suggest that *D. dianthus* is perhaps more akin to the DSC *Madrepora oculata* for which more minor (~3‰) microstructural differences are reported ([Rollion-Bard and Blamart, 2014](#_ENREF_65)). Despite only modest differences measured between δ11B of COC and fibrous aragonite in *D. dianthus*, we document considerable improvement in the δ11B-pH calibration by removing the high-Mg analyses presumably contaminated by COC microstructures. This suggests that the current poorly constrained relationship of bulk skeletal δ11B to seawater pH in *Lophelia pertusa* ([McCulloch et al., 2012b](#_ENREF_54)) could be equally improved by applying similar sampling techniques to this microstructurally heterogeneous taxon.

Fibrous aragonite samples where the microdrill track was seen to visibly deviate into (or close to) the line of weakness between the sample edge and surrounding epoxy were considered potentially contaminated by oxide-rich sample coatings and therefore not included in the δ11B-pH calibration. However, analysis of coating contaminated samples such as these reveal measured δ11B to be up to 3‰ lower than “pure” adjacent fibres samples while deviating little in B/Ca and other trace element chemistry (see 47413 and 94069 “Coating”; Table 2). In light of the detrimental effect of coating contamination on δ11B*fibre* measurements, our sampling techniques focused on the exclusion of these coating phases. High Fe/Ca and Mn/Ca (>200 and >50 μmol/mol respectively) measurements in the 94069 “Coating” sample show that these ancillary data are useful diagnostic tools for identifying substructures strongly contaminated by oxide coatings, however this is not the case for all coating contaminated samples (*e.g.* 47413 “Coating”). Therefore, in a similar argument to that of potential COC contamination, while trace contamination of coating phases in our “pure” fibre samples cannot be discounted, diligent sampling protocols make it unlikely that the spread in replicate δ11B*fibre* measurements is down to incorporation of contaminants alone.

#### Rayleigh fractionation during growth

Spatial chemical profiling across bands of fibrous aragonite reveals that the presumed oldest fibre growth close to COCs has high δ11B and Mg/Ca and low Sr/Ca and U/Ca with respect to presumed newer growth close to the edge of the septum (Figure 4). Such co-varying chemical trends are echoed more generally across the entire fibrous aragonite data set with the concentration of elements more preferentially incorporated into the aragonite lattice with respect to seawater (*i.e.* U) being inversely correlated with δ11B while elements less readily incorporated into the coral (*i.e.* B and Mg) are positively correlated with δ11B (Figure 5). The implied dependency of these elemental-isotopic relationships on partition coefficients suggests a common systematic effect on these parameters during calcification; one candidate for this is Rayleigh fractionation.

Restricted flow of ions from ambient seawater to the ECF leads to Rayleigh-type fractionation as progressive precipitation of skeletal aragonite leaves the residual ECF proportionally enriched/depleted in its elemental ratios to calcium depending on the partition coefficient of the trace element in question. For example, the *D*B into aragonite from seawater is <<1, therefore the B/Ca ratio of the residual calcification fluid increases as progressively more aragonite is precipitated with a B/Ca ratio lower than the parent fluid. In this example, later forming aragonite precipitating from this fluid would have a higher B/Ca ratio than the initial precipitate. Previous Mg/Ca and Sr/Ca measurements by isotope dilution ICP-MS in *D. dianthus* fibrous aragonite are shown to track a closed system type Rayleigh fractionation model (Figure 6 A; Gagnon et al., 2007) hence we apply this closed system model to assess the extent to which simple Rayleigh-type processes can explain Mg/Ca, Sr/Ca, B/Ca and δ11B variability within the fibrous aragonite samples in this study (see Supplementary information for model description).

The exact fraction of Ca remaining in the ECF relative to ambient seawater concentration for each stage of fibrous aragonite calcification is an unknown therefore, following the approach of [Gagnon et al. (2007)](#_ENREF_28), we cross-plot the replicate Mg/Ca and Sr/Ca data for specimens with many replicates from sites 19168, 45669, 47413, and 47407 (Figure 6) to assess the fit to the Rayleigh models. Unlike the data from *D. dianthus* sample 47407 measured by [Gagnon et al. (2007)](#_ENREF_28), fibrous aragonite Mg/Ca and Sr/Ca data in this study do not fit well to Mg:Sr Rayleigh fractionation curves and in many cases show the opposite trend (i.e. positive correlation) to that predicted by Rayleigh fractionation. This further suggests that contaminant phases such as Mg-rich COCs are not being fully rejected by our microsampling techniques. Fibrous aragonite Sr/Ca ratios in this study are generally lower (by ~1 mmol/mol) than Sr/Ca measurements in sample 47407 by [Gagnon et al. (2007)](#_ENREF_28). The offset in Sr/Ca may, in part, represent discrepancies between analytical techniques, however these samples are difficult to compare directly. It remains unclear if our sample from site 47407 was taken from a different coral calyx and therefore calcifying from entirely different ECFs (perhaps with a subtly lower initial [Sr] in the ECF; −5%; Figure 6 A). Where fibrous aragonite was sampled similar to [Gagnon et al. (2007)](#_ENREF_28), sequentially across the same fibre band of sample 19168 (Figure 4; black squares Figure 6 A), Mg/Ca and Sr/Ca data cluster tightly and do fit a Rayleigh distribution albeit with very much lower initial [Sr] in the ECF (−15%). This result suggests that initial [Sr] in the ECF can be modified from ambient seawater and is variable between individuals.

Rayleigh models using low estimates for *D*B similar to inorganic aragonite (0.0037; Mavromatis et al., 2015; Holcomb et al., 2016) and an initial boron concentration in the ECF equal to ambient seawater are unable to describe the measured B/Ca and δ11B*fibre* results, incorporating far too little boron into the skeleton at each iterative step to cause and appreciable shift the residual δ11B of the ECF and δ11B*fibre* (example data from sample 19186 plotted in Figure 6 B). We note that the coral boron chemistry measured in this study can only be defined by tuning the Rayleigh model to unrealistically high *D*B (~0.23; two orders of magnitude greater than inorganic aragonite) and with an initial boron concentration in the ECF (7 μM) approximately 60 times lower than [B]sw (432.6 μM; Lee et al., 2010). Alternatively, it is possible that a biological process that strongly rejects seawater boron from the ECF would cause large isotopic fractionation of internal δ11Bsw, but varying the starting δ11Bsw in the modelled ECF between 35 and 45‰ does little to alter the gradient of the Rayleigh curve; merely shifting the modelled output up/down the *x* axis. Furthermore, addition of active Ca2+ pumping into the ECF (up to 10% replacement of precipitated Ca2+) can only go part way towards increasing the rate at which δ11B*fibre* values increase with respect to B/Ca (shaded region Figure 6 C), hence major reduction of the initial [B] in the ECF and extremely high *D*B values are still required.

Rare earth element spiked seawater culture experiments suggest that seawater transport to coral calcification sites is direct and rapid ([Gagnon et al., 2012](#_ENREF_27)). Therefore, without a mechanism to account for such strong biological modification of *D*B and [B] in the ECF, we must assume that the impact of Rayleigh fractionation on fibre boron chemistry is small.

#### Growth rate effects

Given that Rayleigh fractionation is an unlikely driver of the patterns we observe, an alternative explanation is suggested whereby elemental and isotopic variability are the result of two distinct but related mechanisms; seawater pH and crystal growth rate. While seawater pH is a clear driver of the δ11B in fibrous aragonite, pH also strongly covaries with ΩArag (Table 1). High saturation states in seawater boost carbonate precipitation rates ([Hoegh-Guldberg et al., 2007](#_ENREF_36)) which in turn promotes kinetic disequilibrium at the crystal surface, leading to “growth entrapment” of trace elements that would typically be rejected from the lattice (i.e. B and Mg; *D*<<1; Watson, 2004). A strong dependency on growth-rate has been observed for boron incorporation into inorganically precipitated aragonite ([Mavromatis et al., 2015](#_ENREF_51); [Noireaux et al., 2015](#_ENREF_59)). The decrease in Mg/Ca (and to a lesser extent B/Ca) paired with increases in Sr/Ca and U/Ca from the presumed oldest to the newest fibres, are potentially a result of a progressive decrease in calcification rate from the COC towards the sample edge as the septum is thickened. This change is also supported by the potential decrease in U/Ca-reconstructed carbonate ion concentrations (i.e. calibration by Anagnostou et al., 2011). It is likely therefore that the proposed change in growth rate is associated with a decrease in carbonate ion saturation state within the ECF over time ([Holcomb et al., 2009](#_ENREF_37)), with the rate of change expected to be related to ambient seawater ΩArag. This mechanism would also explain the more muted elemental offsets between fibres and COC in specimen 83583 as growth rates presumably remained slow throughout its life at this low pH site.

## Conclusions and implications

Sample size and shape dictates the choice of fibre sampling technique required to extract exclusively fibrous aragonite sub-samples from a *D. dianthus* calyx. Where possible we recommend the (broad drill bit) micromilling technique be used as it offers high sampling precision over (fine drill bit) microdrilling, without the cost and time burden of laser cutting.

Analyses of bulk carbonate *D. dianthus* samples yield a wide spread in measured δ11B ([Anagnostou et al., 2012](#_ENREF_6); [McCulloch et al., 2012b](#_ENREF_54)) that are likely the result of both microstructural heterogeneities (*i.e.* Blamart et al., 2007) and inherent variability in skeletal aragonite possibly linked to the calcification mechanism. Although microsampling proves a more labour intensive alternative, if regions within the coral with slow growth rate and/or minimal COC contamination are selected (*i.e.* low-Mg fibres), our δ11Bdata yield a more refined calibration against seawater pH in comparison to similar bulk sampling. Our new calibration therefore boosts the utility palaeoceanographic of *D. dianthus* as a substrate for intermediate water pH reconstruction, allowing shifts of < 0.1 pH to be fully resolved (*e.g.* LGM to Holocene).

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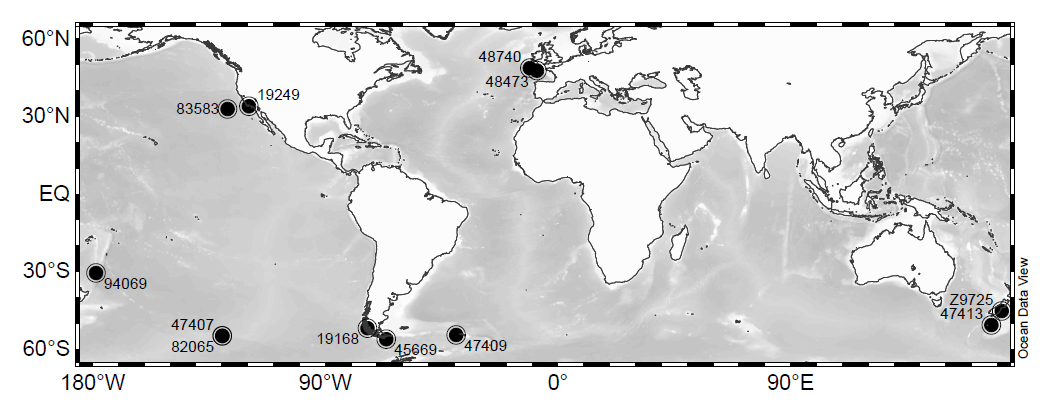


Figure : Locations of the *D. dianthus* specimens used in this study. Map created using Ocean Data View software (R. Schlitzer, 2015, <http://odv.awi.de>)

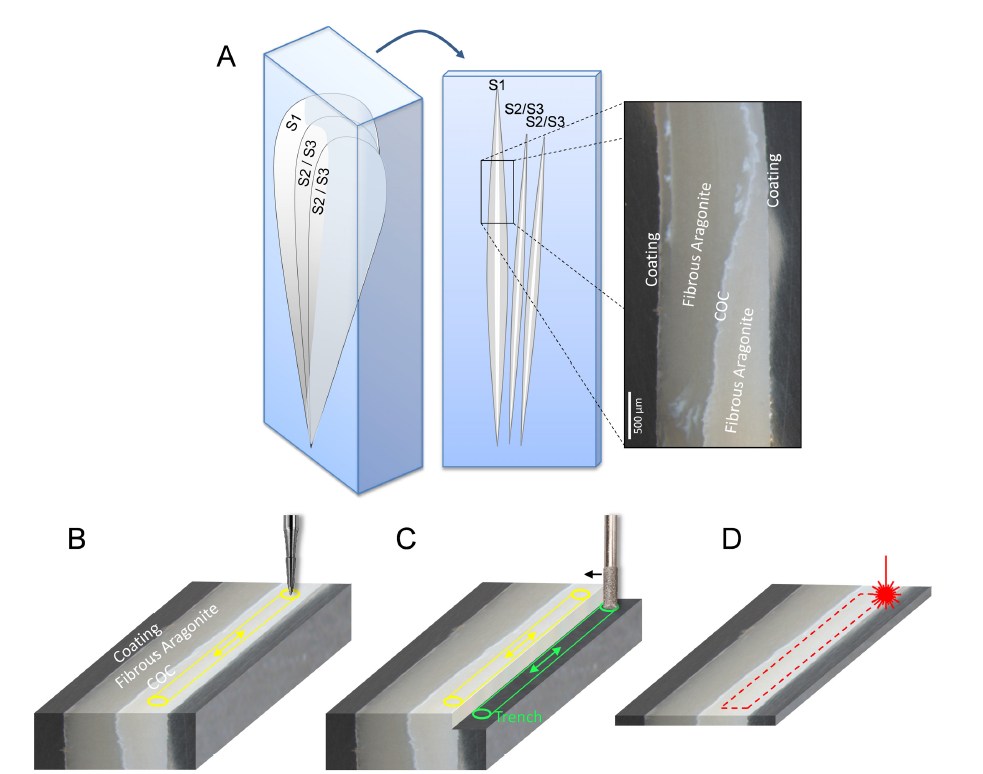


Figure : Sampling of DSC fibrous aragonite. Panel A: Sectioning of DSC septa to expose target fibrous aragonite along with centres of calcification (COC) and edge coatings. Light microscope image of coral sample 19186 used in this study. Panels B, C and D show different sampling techniques of polished sections described in the text. Microdrilling (B): direct drilling of fibres using fine drill bit to collect powdered sample. Micromilling (C): drilling and initial sampling trench (green) before milling towards fibre samples (yellow) using broad drill bit. Laser cutting (D): repeated laser shots are used to cut whole fibre section from a thin section of 200µm thickness.

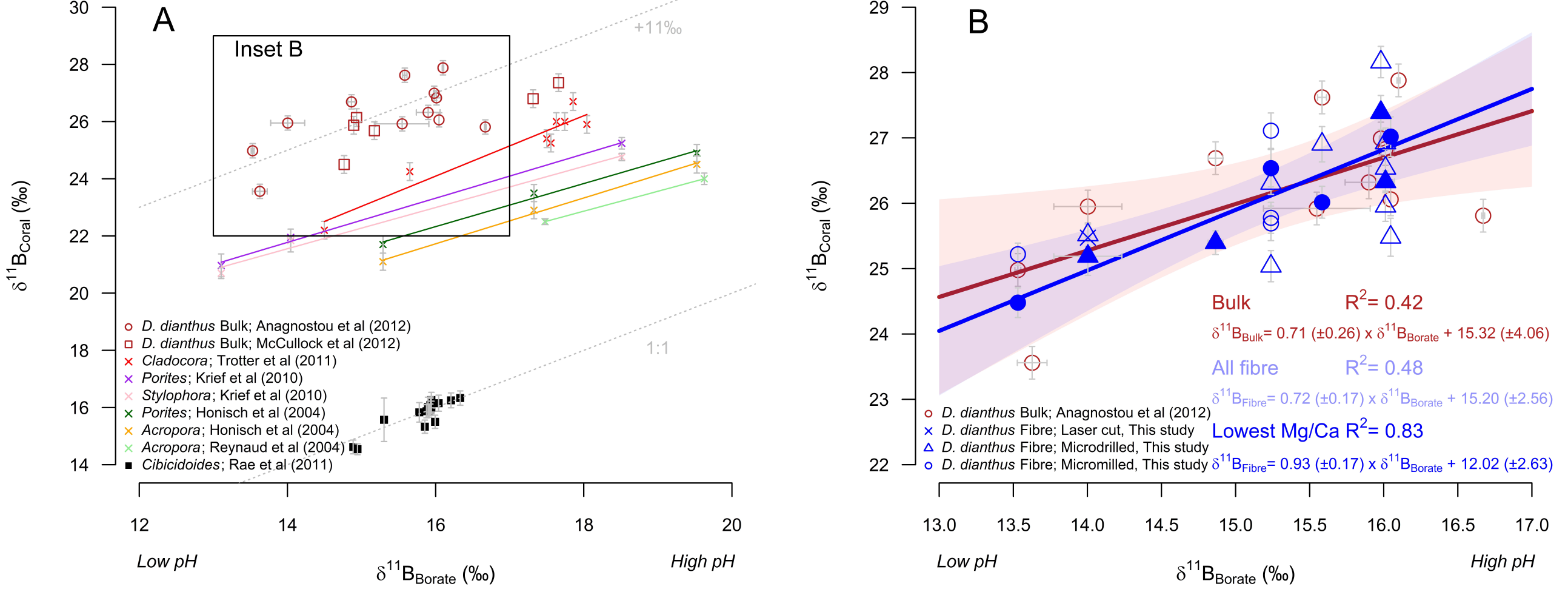


Figure : Measured δ11B of coral regressed against the δ11Bborate of ambient seawater. Panel A. δ11B of bulk *D. dianthus* ([Anagnostou et al., 2012](#_ENREF_6)), tropical corals ([Hönisch et al., 2004](#_ENREF_41); [Krief et al., 2010](#_ENREF_45); [Reynaud et al., 2004](#_ENREF_63); [Trotter et al., 2011](#_ENREF_76)), and bamboo coral ([Farmer et al., 2015](#_ENREF_21)). Grey dashed parallel lines in show 1:1 δ11B relationship and the same relationship offset by +11‰. Panel B. (inset of A) shows new fibrous aragonite *D. dianthus* data in comparison to bulk *D. dianthus* ([Anagnostou et al., 2012](#_ENREF_6)). Open blue symbols represent drilled (triangles), milled (circles) and laser cut fibres (crosses) sampling techniques (error bars show 2σ of each replicate). Filled blue symbols highlight the *D. dianthus* fibre replicate with the lowest Mg/Ca. Regression lines are least squares regression models and their 95% confidence intervals (shaded envelopes) for bulk and low Mg/Ca fibre data.

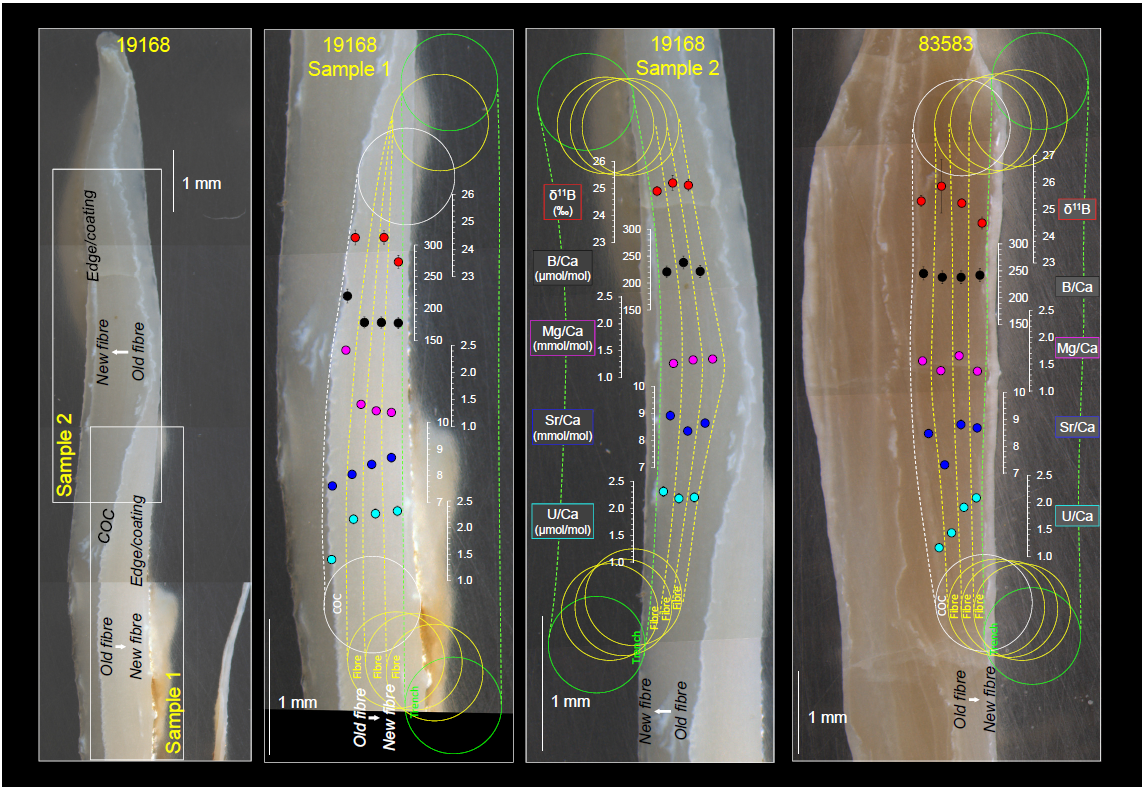


Figure : δ11B and trace metal composition across bands of fibrous aragonite in *D. dianthus* corals 19168 and 83583 (low pH site) sampled by micromilling from presumed newest to oldest fibre growth. The positions of sample trenches are highlighted in green; subsequent samples are then shown in yellow. The positions where COC samples were also taken are delineated by the connected white circles.

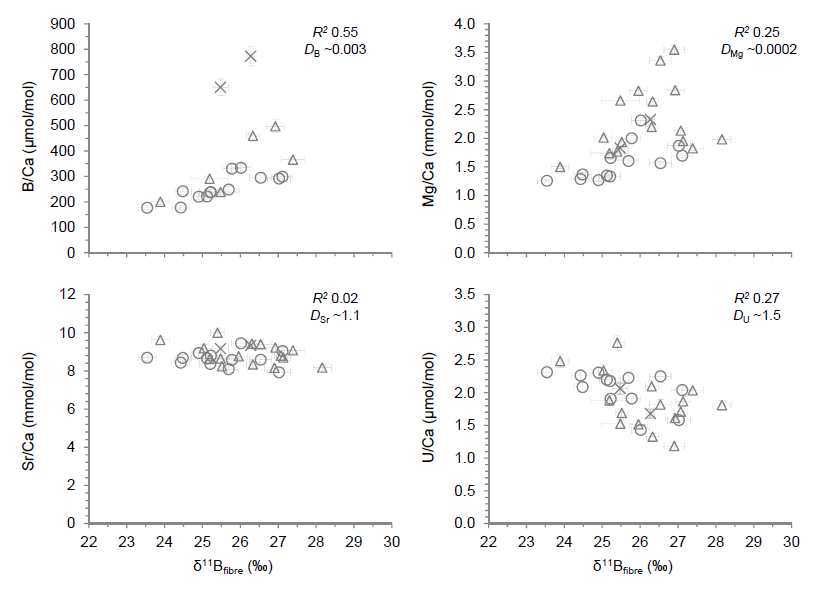


Figure : *D. dianthus* δ11B*fibre* data plotted against B/Ca, Mg/Ca, Sr/Ca and U/Ca trace element composition. Symbols represent fibre sampling technique as in Figure 3. Error bars show the 2σ external reproducibility. *D* values (*e.g.* Gaetani and Cohen, 2006) refer to the approximate partition coefficient of each trace element into aragonite (*D*X = (X/Ca)Aragonite/(X/Ca)Seawater). *R*2 values are representative of the linear regression of just the milled/drilled samples (*i.e.* excluding laser cut samples with anomalously high B/Ca).

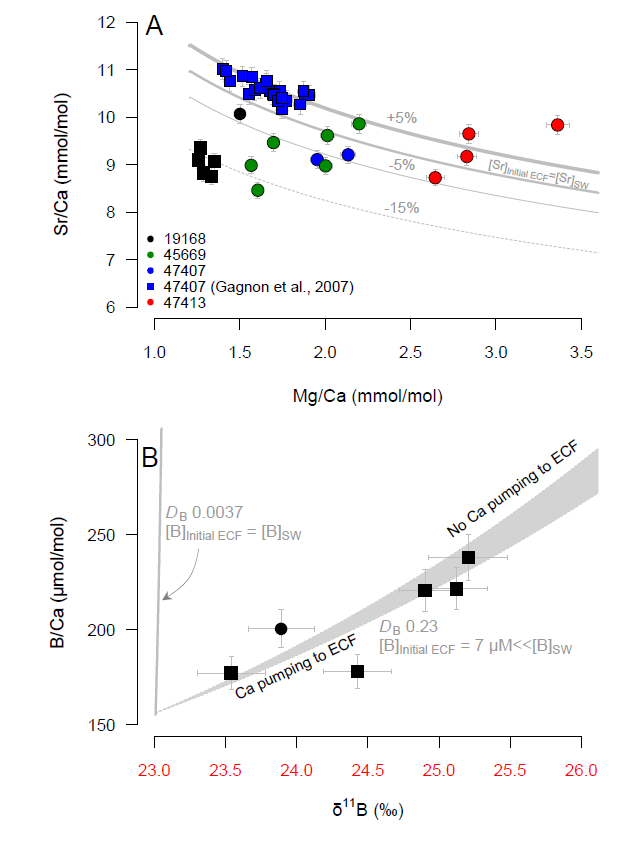


Figure : Rayleigh fractionation models for Mg, Sr, and B incorporation into *D. dianthus* (*e.g.* Gagnon et al., 2007). Panels A and B show theoretical coral Rayleigh fractionation curves for respectively, Mg/Ca *vs* Sr/Ca and B/Ca *vs* δ11B as [Ca] in the ECF reduces. We show the impact of changing initial elemental concentration of ECF (e.g. [Sr]Initial ECF +5%, −5% and −15%) and partition coefficients (*D*). The calcifying fluid assumptions are: [Ca]sw = 10.3 mM; [Sr]sw = 91 µM; [Mg]sw = 52.8 mM; [B]sw = 432.6 µM; δ11Bsw = 39.61 µmol/kg; αB = 1.0272; p*K*B\* = 8.79; internal pH = 8.62. Data for *D. dianthus* specimens in this study with many replicates (19168, 45669, and 47413) are plotted to show their fit to the Rayleigh fractionation models. Squares represent fibre replicates taken sequentially from the same fibre band (19168 this study; Figure 4; and 47407; Gagnon et al., 2007). Fibre replicates taken from different septa within the samples are shown as circles. The shaded region of panel B represents the impact of active Ca2+ pumping into the ECF replacing up to 10% of the Ca2+ lost to CaCO3 precipitation.

Table : *D. dianthus* specimens used in this study and previously measured boron isotope data for bulk *D. dianthus* ([Anagnostou et al., 2012](#_ENREF_6)). Proximal hydrographic bottle data are from the CDIAC ocean carbon system database. Additional carbonate system parameters based on specimen location are calculated using SeaCarb R package.



Table : Fibrous aragonite *D. dianthus* δ11B and trace element data. Measured δ11B data are corrected for the effect of the measured total procedural blank (TPB). Sampling method symbols correspond to those in Figure 3.



Table : Trace element results of cleaned/uncleaned bulk sample 19168 crushed by pestle and mortal and JCp-1 powdered coral standard. Results of uncleaned JCp-1 are compared to the interlaboratory comparison study by Hathorne et al., ([2013](#_ENREF_32)). All uncleaned JCp-1 results are within the robust standard deviation of Hathorne et al., ([2013](#_ENREF_32)) with the exception of Sr/Ca for which a correction factor of 1.047 must be applied to Sr/Ca results in this study.



Table : Regression summary of *D. dianthus* fibrous aragonite (this study) and bulk septa ([Anagnostou et al., 2012](#_ENREF_6)) calibrations. Regression in the form δ11Bcoral = *m* × δ11Bborate + *c* presented with one standard error uncertainties.



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