Dear Reviewers,

First of all, we would like to thank you for your detailed comments that helped to improve the manuscript. We think we have addressed all of them. You will find the new version of the manuscript, with changes highlighted attached to this letter, along with a clean version of the manuscript.

To comment on a few points:

One of the main comments we adressed is the sensitivities of $\delta^{11}B_{carbonate}$ to $\delta^{11}B_{borate}$, >1 for *G. ruber* and *T. sacculifer*. We acknowledge that the sensitivities are not statistically different from published literature. When doing the bootstrap analysis on all compiled data, the regression for *G. ruber* was similar to Raitzsch et al., (2018), with a sensitivity of 0.46 ± 0.34 (updated table 3) compared to 0.45 ± 0.16 for Raitzsch et al., (2018) and the sensitivity was of 0.83 ± 0.48 for *T. sacculifer* in line with Martinez-Boti et al., (2015b) and Raitzsch et al., (2018). We also take into account the pCO₂ based $\delta^{11}B$ reconstructions against the Vostok pCO₂ record as pointed out. We also have clearly written that possible sensitivities of >1 are speculative.

Another major change is the reconstruction of microenvironment pH for each species (Figure 8) which we believe are in line with results of a study of photosynthesis/symbiont density from Takagi et al., (2019) for different species. This also helps to explain a possible light-limiting threshold for calibrations. We have also added discussion around symbionts-host interactions of different species.

We do observe changes in calcification depth with thermocline depth for most of the deeper-dwelling species (*G. ruber*, *T. sacculifer*) which might be problematic for future reconstructions over periods when there is evidence for changes in stratification. However, our data are too limited to derive a robust trend.

We understand that our discussion on the Western Equatorial Pacific in the earlier draft was limited as it was based on a single site. In order to strengthen the discussion, we added two other sites, ODP Sites 806A and 807A. The data record similar, low values to what we reported for site WPO7, and add to our confidence in the results.

Also, a reviewer pointed out a potential issue with the circularity of some of the calculations presented in the last manuscript. In order to address this issue, we recalculate for each species a calibration that excludes results for a single site, and then use the modified calibration that contains a subset of calibration data to reconstruct vertical profiles of carbonate chemistry as shown in Figure 9. What Fig. S5 shows is the comparison of vertical profiles using these two approaches, which yield consistent results. On average, the difference in δ ¹¹B is 0.8% for ¹¹B_{borate}, 0.2% for pH and 5% for pCO₂. The fact that using a subset of the calibration data and all of the calibration data yields similar results validates the work.

Reviews are adressed in the commented version of the manuscript and in our response to reviewers. We thank you for your time.

Seawater pH reconstruction using boron isotopes in multiple planktonic foraminifera species with different depth habitats and their potential to constrain pH and pCO₂ gradients Maxence Guillermic^{1,2}, Sambuddha Misra^{3,4}, Robert Eagle^{1,2}, Alexandra Villa^{2,5}, Fengming Chang⁶, Aradhna Tripati 1,2 ¹ Department of Earth, Planetary, and Space Sciences, Department of Atmospheric and Oceanic Sciences, Institute of the Environment and Sustainability, UCLA, University of California - Los Angeles, Los Angeles, CA 90095 USA ² Laboratoire Géosciences Océan UMR6538, UBO, Institut Universitaire Européen de la Mer, Rue Dumont d'Urville, 29280, Plouzané, France ³ Indian Institute of Science, Centre for Earth Sciences, Bengaluru, Karnataka 560012, India ⁴ The Godwin Laboratory for Palaeoclimate Research, Department of Earth Sciences, University of Cambridge, UK ⁵ Department of Geology, University of Wisconsin-Madison, Madison, WI 53706 USA ⁶ Key Laboratory of Marine Geology and Environment, Institute of Oceanology, Chinese Academy of Sciences, Qingdao 266071, China Submitted to Biogeosciences *Corresponding author: E-mail address: maxence.guillermic@gmail.com

44 **ABSTRACT**

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Boron isotope systematics of planktonic foraminifera from core-top sediments and culture experiments have been studied to investigate the sensitivity of $\delta^{11}B$ of their calcite tests to seawater pH. However, our knowledge of the relationship between $\delta^{11}B$ and pH remains incomplete for many taxa. Thus, to expand the potential scope of application of this proxy, we report δ^{11} B data for 7 different species of planktonic foraminifera from sediment core-tops. We utilize a method for the measurement of small samples of foraminifera and calculate the $\delta^{11}B$ calcite sensitivity to pH for Globigerinoides ruber, Trilobus sacculifer (sacc or w/o sacc), Orbulina universa, Pulleniatina obliquiloculata, Neogloboquadrina dutertrei, Globorotalia menardii and Globorotalia tumida, including for unstudied core-tops and species. The sensitivity of $\delta^{11}B_{carbonate}$ to $\delta^{11}B_{borate}$ (eg. $\Delta \delta^{11} B_{carbonate} / \Delta \delta^{11} B_{borate}$ in core-tops is consistent with previous studies for T. sacculifer and G. ruber and close to unity for N. dutertrei, O. universa and combined deep-dwelling species. Deep-dwelling species closely follow the core-top calibration for O. universa, which is attributed to respiration-driven microenvironments, likely caused by light limitation and/or symbiont/host interactions. These taxa have diverse ecological preferences and are from sites that span a range of oceanographic regimes, including some that are in regions of air-sea equilibrium and others that are out of equilibrium with the atmosphere. Our data support the premise that utilizing boron isotope measurements of multiple species within a sediment core can be utilized to constrain vertical profiles of pH and pCO2 at sites spanning different oceanic regimes, thereby constraining changes in vertical pH gradients and yielding insights into the past behavior of the oceanic carbon pumps.

1. Introduction

The oceans are absorbing a substantial fraction of anthropogenic carbon emissions resulting in declining surface ocean pH (Fig. 1; IPCC, 2014). Yet there is a considerable uncertainty over the magnitude of future pH change in different parts of the ocean and the response of marine biogeochemical cycles to physio-chemical parameters (T, pH) caused by climate change (Bijma et al., 2002; Ries et al., 2009). Therefore, there is an increased interest in reconstructing past seawater pH (Hönisch and Hemming, 2005; Liu et al., 2009; Wei et al., 2009; Douville et al., 2010), in understanding spatial variability in aqueous pH and carbon dioxide (pCO_2) (Foster et al., 2008; Martinez-Boti et al., 2015b; Raitzsch et al., 2018), and in studying the response of the biological carbon pump utilizing geochemical proxies (Yu et al., 2007, 2010, 2016).

Although proxies for carbon cycle reconstruction are complex in nature (Pagani et al., 2005; Tripati et al., 2009, 2011; Allen and Hönisch, 2012), the boron isotope composition of foraminiferal tests is emerging as one of the more robust candidates (Hönisch et al., 2005, 2009; Ni et al., 2007; Foster et al., 2008, 2012; Bartoli et al., 2011; Henehan et al., 2013; Martinez-Boti et al., 2015b; Chalk et al., 2017). The study of laboratory cultured foraminifera has demonstrated a systematic dependence of the boron isotope composition of tests on ambient pH (Sanyal et al., 1996, 2001; Henehan et al., 2013, 2016). Core-top measurements on globally distributed samples also show a δ^{11} B sensitivity to pH with taxa-specific offsets from the theoretical fractionation line of borate ion (Rae et al., 2011; Henehan et al., 2016; Raitzsch et al., 2018).

Knowledge of seawater pH, in conjunction with constraints on one other carbonate system parameter (Total Alkalinity (TA), DIC (dissolved inorganic carbon), [HCO₃⁻], [CO₃²-]), can be utilized to constrain aqueous pCO₂. Application of empirical calibrations for boron isotopes, determined for select species of foraminifera from core-tops and laboratory cultures, has resulted in accurate reconstructions of pCO₂ utilizing downcore samples from sites that are in quasi-equilibrium with the atmosphere at present. δ^{11} B_{carbonate} based reconstructed values of pCO₂ are analytically indistinguishable from ice core CO₂ records (Hönisch et al., 2005, 2009; Foster et al., 2008; Henehan et al., 2013; Chalk et al., 2017).

Therefore, the last decade has produced several studies aiming at reconstructing past seawater pH using boron isotopes to constrain atmospheric pCO_2 in order understand the changes in the global carbon cycle (Hönisch et al., 2005, 2009; Foster et al., 2008, 2012, 2014; Seki et al., 2010; Bartoli et al., 2011; Henehan et al., 2013; Martinez-Boti et al., 2015a, 2015b; Chalk et al., 2017). In addition to reconstructing atmospheric pCO_2 , in a few studies, the $\delta^{11}B$ proxy has been applied to mixed-layer planktonic foraminifera at sites out of equilibrium with the atmosphere to constrain past air-sea fluxes (Foster et al., 2014; Martinez-Boti et al., 2015b). A small body of work has examined whether data for multiple species in core-top (Foster et al., 2008) and down-core samples could be used to constrain vertical profiles of pH through time (Palmer et al., 1998; Pearson and Palmer, 1999).

In this study, we make critical additions to the emerging pool of boron isotope data of core-top planktonic foraminifera from different oceanographic regimes, including data for species that have not previously been examined. We utilize a low-blank (15 pg B to 65 pg B), high precision (2sd on the international standard JCP-1 is $0.20 \, \%$, n=6) $\delta^{11}B_{carbonate}$ analysis method (down to ~250 µg CaCO₃), modified after Misra et al. (2014), to study multiple species of planktonic foraminifera from sediment core-tops that span a range of oceanographic regimes, including open-ocean oligotrophic settings and marginal seas. We constrain calibrations for different species, and compare results to published work (Foster et al., 2008; Henehan et al., 2013; Henehan

et al., 2016; Martinez-Boti et al., 2015b; Raitzsch et al., 2018). We also test whether these data support the application of boron isotope measurements of multiple species within a sediment core as a proxy for constraining vertical profiles of pH and pCO₂.

2. Background

2.1 Planktonic foraminifera as archives of seawater pH

Planktonic foraminifera are used as archives of past environmental conditions within the mixed layer and thermocline, as their chemical composition is correlated with the physio-chemical parameters of their calcification environment (Ravelo and Fairbanks, 1992; Elderfield and Ganssen, 2000; Dekens et al., 2002; Anand et al., 2003; Sanyal et al., 2001; Ni et al., 2007; Henehan et al., 2013, 2015, 2016; Howes et al., 2017; Raitzch et al., 2018). The utilization of geochemical data for multiple planktonic foraminifera species with different ecological preferences to constrain vertical gradients has been explored in several studies. The framework for such an approach was first developed using modern samples of planktonic foraminifera for oxygen isotopes, where it was proposed as a tool to constrain vertical temperature gradients and study physical oceanographic conditions during periods of calcification (Ravelo and Fairbanks, 1992).

Because planktonic foraminifera species complete their lifecycle in a particular depth habitat due to their ecological preference (Ravelo and Fairbanks, 1992; Farmer et al., 2007), it is theoretically possible to reconstruct water column profiles of pH using data from multiple taxa (Palmer and Pearson, 1998; Anagnostou et al., 2016). The potential use of an analogous approach to reconstruct past profiles of seawater pH was first highlighted by Palmer and Pearson (1998) on Eocene samples to constrain pH-depth gradients. However, in these boron isotope-based studies, it was assumed that boron isotope offset from seawater and foraminiferal carbonate were constant, which is an assumption not supported by subsequent studies (e.g., Hönisch et al., 2003; Foster et al., 2008; Henehan et al., 2013, 2016; Raitszch et al., 2018; Rae, 2018). Furthermore, δ^{11} B differences between foraminifera species that inhabit waters that are the same pH makes the acquisition of more core-top and culture data essential for applications of the proxy.

2.2 Boron systematics in seawater

Boron is a conservative element in seawater with a long residence time ($\tau_B \sim 14$ Myr) (Lemarchand et al., 2002a). In seawater, boron exists as trigonal boric acid B(OH)₃ and tetrahedral borate ion B(OH)₄ (borate). The relative abundance of boric acid and borate ion is a function of the ambient seawater pH. At standard open ocean conditions (T = 25 °C and S = 35), the dissociation constant of boric acid is 8.60 (Dickson, 1990), implying that boron mainly exists in the form of boric acid in seawater. Since the pK_B and seawater pH (e.g., ~8.1, NBS) values are similar, it implies that small changes in seawater pH will induce strong variations in the abundance of the two boron species (Fig. 2).

Boron has two stable isotopes, ^{10}B and ^{11}B , with average relative abundances of 19.9 and 80.1 %, respectively. Variations in B isotope ratio are expressed in conventional delta (δ) notation:

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$$\delta^{11} B (\%) = 1000 x \left(\frac{{}^{11} B / {}^{10} B_{Sample}}{{}^{11} B / {}^{10} B_{NIST 951-a}} - 1 \right)$$
 (1)

where positive values represent enrichment in the heavy isotope ¹¹B, and negative values enrichment in the light isotope ¹⁰B, relative to the standard reference material. Boron isotope values are reported versus the NIST SRM 951 (Cantazaro et al., 1970).

B(OH)₃ is enriched in ¹¹B compared to B(OH)₄ with a constant offset between the two chemical species, within the range of physio-chemical variation observed in seawater, given by the fraction factor (α). The fractionation (ϵ) between B(OH)₃ and B(OH)₄ of 27.2 ± 0.6 % has been empirically determined by Klochko et al., (2006) in seawater. Note, Nir et al., (2015) calculate this fractionation, using an independent method, to be 26 ± 1 %, which is within the analytical uncertainty of the Klochko et al., (2006) value.

2.3 Boron isotopes in planktonic foraminifera calcite

Many biogenic carbonate-based geochemical proxies are affected by "vital effects" or biological fractionations (Urey et al., 1951). The $\delta^{11}B_{carbonate}$ in foraminifera exhibits species-specific offsets (see Rae et al., 2018 for review) compared to theoretical predictions for the boron isotopic composition of B(OH)₄- (α =1.0272, Klochko et al., 2006). As the analytical and technical aspects of boron isotope measurements have improved (Foster et al., 2008; Rae et al., 2011; Misra et al., 2014; Lloyd et al., 2018), evidence for taxonomic differences have not been eliminated, but have become increasingly apparent (Foster et al., 2008, 2018; Henehan et al 2013, 2016; Noireaux et al., 2015; Foster et al., 2016; Rae et al., 2018; Raitzsch et al., 2018).

At present, culture and core-top calibrations have been published for several planktonic species including *Trilobatus sacculifer*, *Globigerinoides ruber*, *Globigerina bulloides*, *Neogloboquadrina pachyderma*, *Orbulina universa* (Foster et al., 2008; Henehan et al., 2013; Henehan et al., 2015; Sanyal et al., 1996; Sanyal et al., 2001). Although the boron isotopic composition of several species of foraminifera are now commonly used tools for reconstructing surface seawater pH, for other species, there is a lack of data constraining boron isotope sensitivity between foraminiferal carbonate and borate ion in seawater.

2.4 Origin of biological fractionations in foraminifera

Perforate foraminifera are calcifying organisms that maintain a large degree of biological control over their calcification space, and thus, mechanisms of biomineralization may be of significant importance in controlling the $\delta^{11}B$ of the biogenic calcite. The biomineralization of foraminifera is based on seawater vacuolization (Erez, 2003; de Nooijer et al., 2014) with parcels of seawater being isolated by an organic matrix thereby creating a vacuole filled with seawater. Recent work has also demonstrated that even if the chemical composition of the reservoirs is modified by the organism, seawater is directly involved in the calcification process with vacuoles formed at the periphery of the shell (de Nooijer et al., 2014). Culture experiments by Rollion-Bard and Erez., (2010) have proposed that the pH at the site of biomineralization is elevated to an upper pH limit of ~9 for the shallow-water, symbiont-bearing benthic foraminifera *Amphistegina lobifera*, which would support a pH modulation of a calcifying fluid in foraminifera. We acknowledge this is speculative as it is based upon benthic foraminifer experiments.

For taxa with symbionts, the microenvironment surrounding the foraminifera is chemically different from seawater due to photosynthetic activity (Jorgensen et al., 1985; Rink et al., 1998; Köhler-Rink and Kühl, 2000). Photosynthesis by the symbionts elevates the pH of the microenvironment (Jorgensen et al., 1985; Rink et

al., 1998; Wolf-Gladrow et al., 1999; Köhler-Rink and Kühl, 2000), while calcification and respiration decrease it (Equation 2 and 3).

$$Ca^{2+} + 2HCO_3^- \leftrightarrow CaCO_3 + H_2O + CO_2 \text{ or } Ca^{2+} + CO_3^{2-} \leftrightarrow CaCO_3$$
 [calcification] (2)
 $CH_2O + O_2 \leftrightarrow CO_2 + H_2O$ [respiration/photosynthesis] (3)

 $\delta^{11}B$ in foraminifera is primary controlled by seawater pH, but is also dependent of the pH alteration of microenvironments due to calcification, respiration and symbiont photosynthesis. $\delta^{11}B$ should therefore reflect the relative dominance of these processes and may account for species-specific $\delta^{11}B$ offsets. Theoretical predictions from Zeebe et al. (2003) and foraminiferal data from Hönisch et al., (2003) highlighted the dominance of microenvironment pH in $\delta^{11}B$ signature of foraminifera. Their work also suggested that for a given species, there should be a constant offset observed between the boron isotope composition of foraminifera and borate ion over a large range of pH, imparting confidence in utilizing species-specific boron isotope data as a proxy for seawater pH.

Comparison of boron isotope data for multiple planktonic foraminiferal species indicate that taxa with high levels of symbiont activity such as T. sacculifer and G. ruber show higher $\delta^{11}B$ values than the $\delta^{11}B$ of ambient borate (Foster et al., 2008, Henehan et al., 2013, Raitzsch et al., 2018). The sensitivities $(\Delta\delta^{11}B_{carbonate}/\Delta\delta^{11}B_{borate})$ referred to as the slope) of existing calibrations suggest a different species-specific sensitivity for these species compared to other taxa (Sanyal et al., 2001; Henehan et al., 2013; Henehan et al., 2015; Raitzsch et al., 2018). For example, *Orbulina universa* exhibits a lower $\delta^{11}B$ than *in situ* $\delta^{11}B$ values of borate ion (Henehan et al., 2016), consistent with the species living deeper in the water column characterized by reduced photosynthetic activity.

It is possible that photosynthetic activity by symbionts might not be able to compensate for changes in calcification and/or respiration, leading to an acidification of the microenvironment. It is interesting to note that for *O. universa* the slope determined for the field-collected samples is not statistically different from unity (0.95 \pm 0.17) (Henehan et al. 2016), while culture experiments report slopes of \leq 1 for multiple species including *G. ruber* (Henehan et al., 2013), *T. sacculifer* (Sanyal et al., 2001), and *O. universa* (Sanyal et al., 1999). More core-top and culture calibrations are needed to fully understand why different slopes are observed, which is part of the motivation for this study.

2.5 Planktic foraminifera depth and habitat preferences

The preferred depth habitat of different species of planktonic foraminifera depends on their ecology, which in turn relies on the hydrographic conditions. For example, *G. ruber is* commonly found in the mixed layer (Fairbanks and Wiebe, 1980; Dekens et al., 2002; Farmer et al., 2007) during the summer (Deuser et al., 1981) whereas *T. sacculifer* is present in the mixed layer until mid-thermocline depths (Farmer et al., 2007) during spring and summer (Deuser et al., 1981, 1989). Specimens of *P. obliquiloculata* and *N. dutertrei* are abundant during winter months (Deuser et al., 1989), with an acme in the mixed layer (~60m) for *P. obliquiloculata*, and at mid-thermocline depths for *N. dutertrei* (Farmer et al., 2007). In contrast, *O. universa* tends to record annual average conditions within the mixed layer. Specimens of *G. menardii* calcify within the seasonal thermocline (Fairbanks et al., 1982, Farmer et al., 2007, Regenberg et al., 2009), and in some regions in

the upper thermocline (Farmer et al., 2007), and records annual temperatures. *G. tumida* is found at the lower thermocline or below the thermocline and records annual average conditions (Fairbanks and Wiebe, 1980; Farmer et al., 2007, Birch et al., 2013).

3. Materials and Methods

3.1 Localities studied

Core-top locations were selected to span a broad range of seawater pH, carbonate system parameters, and oceanic regimes. Samples from Atlantic Ocean (CD107-A), Indian Ocean (FC-01a and FC-02a), Arabian Sea (FC-13a and FC-12b) and Pacific Ocean (WP07-01, A14, and Ocean Drilling Program 806A and CO7A) were analyzed; characteristics of the sites are summarized in Table 1 and S7, Fig. 3, and Fig. 4.

Atlantic site CD107-a (CD107 site A) was drilled in 1997 by the Benthic Boundary Layer program (BENBO) (K.S. Black et al., 1997 - cruise report RRS Charles Darwin Cruise 107). Arabian Sea sites FC-12b (CD145 A150) and FC-13a (CD145 A3200) were retrieved by the *Charles Darwin* in the Pakistan Margin in 2004 (B.J. Bett et al., 2003 - cruise report n°50 RRS Charles Darwin Cruise 145). A14 was recovered by box corer in the southern area of the South China Sea in 2012. Core WP07-01 was obtained from the Ontong Java Plateau using a giant piston corer during the Warm Pool Subject Cruise in 1993. Holes 806A and 807A were retrieved on Leg 130 by the Ocean Drilling Program (ODP). The top 10 cm of sediment from CD107-A have been radiocarbon dated to be Holocene <3 ky (Thomson et al., 2000). Samples from multiple box cores from Indian Ocean sites were radiocarbon dated as Holocene <7.3 ky (Wilson et al., 2012). Samples from western equatorial Pacific Site 806B, close to site WP07-01, are dated to between 7.3-8.6 ky (Lea et al., 2000). Arabian Sea and Pacific core-top samples were not radiocarbon dated but are assumed to be Holocene.

3.2 Species

Around 50-100 foraminifera shells were picked from the 400-500 µm fraction size for *Globorotalia menardii* and *Globorotalia tumida*, >500 µm for *Orbulina universa*, from the 250-400 µm fraction size for *Trilobatus sacculifer* (w/o sacc, without sacc-like final chamber), *Trilobatus sacculifer* (sacc, sacc-like final chamber), *Globigerinoides ruber* (white, sensu stricto), *Neogloboquadrina dutertrei*, *Pulleniatina obliquiloculata*. The samples picked for analyses were visually well preserved.

3.3 Sample cleaning

Briefly, picked foraminifera were gently cracked open, clay removed and checked for coarse-grained silicates. The next stages of sample processing and chemical separation were performed in a class 1000 clean lab equipped with boron-free HEPA filters. Samples were then cleaned using full reductive and oxidative cleaning (Boyle and Keigwin, 1955) Barker et al., 2003). A final leaching step with 0.001N HCl was done before dissolution in 1N HCl. Each sample was divided into two aliquots: an aliquot for boron purification and one aliquot for trace element analysis.

3.4 Reagents

Double-distilled HNO₃ and HCl acids (from Merck® grade) and a commercial bottle of HF Ultrapure grade were used at Brest. Double-distilled acids were used at Cambridge. All acids and further dilutions were

prepared using double-distilled 18.2 M Ω .cm-1 MQ water. Working standards for isotope ratio and trace element measurements were freshly diluted on a daily basis with the same acids used for sample preparation to avoid any matrix effect.

32.5 Boron isotopes

Boron purification for isotopic measurement was done utilizing microdistillation method developed by Gaillardet et al., (2001), for Ca-rich matrices by Wang et al., (2010) and adapted at Cambridge by Misra et al., (2014a). 70 μ L of dissolved carbonate sample was loaded on a cap of a clean fin legged 5 mL conical beaker upside down. The tightly closed beaker was put on a hotplate at 95°C for 15 hours. The beakers were taken off the hotplate and were allowed to cool for 15 min. The cap where the residue formed was replaced by a clean one. Then, 100 μ L of 0.5% HF were added to the distillate.

Boron isotopic measurements were carried out on a Thermo Scientific ®Neptune+ MC-ICP-MS at the University of Cambridge. Neptune+ was equipped with Jet interface and two 10^{13} Ω resistors. The instrumental setup included Savillex® 50μ l/min C-flow self-aspirating nebulizer, single pass Teflon® Scott-type spray chamber constructed utilizing Savillex® column components, 2.0 mm Pt injector from ESI®, Thermo® Ni 'H' type sample cone and 'X' type skimmer cones. Both isotopes of boron were determined utilizing 10^{13} Ω resistors (Misra et al., 2014a; Lloyd et al., 2018).

The sample size for boron isotope analyses typically ranged from 10 ppb B (~5 ng B) to 20 ppb B samples (~10 ng B). Instrumental sensitivity for 11 B was 17 mV/ppb B (eg. 170 mV for 10ppb B) in wet plasma at 50μ l/min sample aspiration rate. Intensity of 11 B for a sample at 10ppb B was typically $165\text{mV} \pm 5\text{mV}$ closely matched the $170\text{mV} \pm 5\text{mV}$ of the standard. Due to the low boron content of the samples extreme care was taken to avoid boron contamination during sample preparation and reduce memory effect during analysis. Procedural boron blanks ranged from 15pg B to 65 pg B (contributed to less than <1% of the sample signal). The acid blank during analyses was measured at $\leq 1\text{mV}$ on the 11 B, meaning a contribution < 1% of the sample intensity, no memory effect was observed within and across sessions.

Analyses of external standards were done to ensure data quality. For $\delta^{11}B$ measurements two carbonate standards were utilized: the JCP-1 (Geological Survey of Japan, Tsukuba, Japan) international standard (Guishr et al., 2014) and the NEP internal coral (Porites sp., $\delta^{11}B = 26.12 \pm 0.92$ ‰, 2SD, n=33 Holcomb et al., 2015 and Sutton et al., 2018, Table S2) from University of Western Australia/Australian National University. Certified boron isotopes liquid standard, the ERM® AE121 ($\delta^{11}B = 19.9 \pm 0.6$ ‰, SD, certified) was used to monitor reproducibility and drift during each session (Vogl and Rosner, 2011; Foster et al., 2013; Misra et al., 2014). Results for the isotopic composition of the NEP standard are shown in Table S2, average values are $\delta^{11}B_{NEP} = 25.70 \pm 0.93$ ‰ (2SD, n=22) over different 7 analytical sessions with each number representing an ab-initio processed sample - this study). Our results are within error of published values of 26.20 ± 0.88 ‰ (2SD, n = 27) and 25.80 ± 0.89 ‰ (2SD, n = 6) by Holcomb et al. (2015) and Sutton et al. (2018) respectively. Chemically cleaned JCP₁ samples were measured at 24.06 ± 0.20 (2SD, n=6) and is within error of published values of 24.37 ± 0.32 ‰ and 24.42 ± 0.28 ‰ by Holcomb et al. (2015) and Sutton et al. (2018) respectively.

3.6 Trace elements

The calcium concentration of each sample was measured on an ICP-AES ® Ultima 2 HORIBA at the Pôle spectrometrie Océan (PSO), UMR6538 (Plouzané, France). Samples were then diluted to fixed calcium concentrations (typically 10 ppm or 30 ppm Ca) using 0.1 M HNO3 & 0.3 M HF matching multi-element standards Ca concentration to avoid any matrix effect (Misra et al., 2014b). Trace elements (e.g. X/Ca ratios) were analyzed on a Thermo Scientific ® Element XR HR-ICP-MS at the PSO, Ifremer (Plouzané, France).

Trace element analyses were done at a Ca concentration of 10 or 30 ppm. The typical blanks for a 30 ppm Ca session were: ⁷Li < 2%, ¹¹B < 7%, ²⁵Mg < 0.2% and ⁴³Ca < 0.02%. Additionally, blanks for a 10 ppm Ca session were: ⁷Li < 2.5%, ¹¹B < 10%, ²⁵Mg < 0.4% and ⁴³Ca < 0.05%. Due to strong memory effect for boron and instrumental drift on the Element XR, long sessions of conditioning were done prior analyses. Boron blanks were driven below 5% of signal intensity usually after 4 to 5 days of centinuous analyses of carbonate samples. External reproducibility was determined on the consistency standard Cam-Wuellestorfi (courtesy of the University of Cambridge) (Misra et al., 2014b), Table S3. Our X/Ca ratio measurements on the external standard Cam-Wuellestorfi were all the time within error of the published value (Table S3) validating the robustness of our trace elements data. Analytical uncertainty of a single measurement was calculated from the reproducibility of the Cam-Wuellestorfi, measured during a particular mass spectrometry session. The analytical uncertainties on the X/Ca ratios are: 0.4 μmol/mol for Li/Ca, 7 μmol/mol for B/Ca and 0.01 mmol/mol for Mg/Ca (2SD, n=31) respectively.

3.7 Oxygen isotopes

Carbonate $\delta^{13}C$ and $\delta^{18}O$ were measured on a Gas Bench II coupled to a Delta V mass spectrometer at the stable isotope facility of Pôle spectrometrie Océan (PSO), Plouzané. Around 20 shells were weighed, crushed and clay removed. The recovered foraminifera were weighed in tubes and flushed with He gas. Samples were then digested in phosphoric acid and analyzed. Results were calibrated to the VPDB scale by international standard NBS19 and analytical precision on the in-house standard Ca21 was better than 0.11% for $\delta^{18}O$ (1SD, n=5) and 0.03% for $\delta^{13}C$ (1SD, n=5).

3.8 Calcification depth determination

We utilized two different chemo-stratigraphic methods to estimate the calcification depth in this study (Table S6 and S7). The first method, commonly used in paleoceanography, utilizes $\delta^{18}O$ measurements of the carbonate ($\delta^{18}O_c$) to estimate calcification depths (referred to as $\delta^{18}O$ -based calcification depths) (Schmidt et al., 2002; Mortyn et al., 2003; Sime et al., 2005; Farmer et al., 2007; Birsh et al., 2013). The second method utilizes Mg/Ca-based temperature estimates ($T_{Mg/Ca}$) to constrain calcification depths (Quintana Krupinski et al., 2017). In both cases, the postulate was that vertical profiles of seawater temperature are available for different seasons in ocean atlases and cruise reports, and that hydrographic data and geochemical proxy signatures can be compared to assess the depth in the water column that represents the species maximum abundance.

The two different methods to estimate calcification depth were then compared to published depth estimates for the basin, and where available, for the same site (Table S6). We chose literature values for calcification depths when available, or depths that were closest to what is known for the region or basin. As foraminifera can migrate in the water column along their ontogeny, we applied (based on uncertainties of our

measurements) an uncertainty of ± 10 m for calcification depths > 70 m and an uncertainty of ± 20 m when calcification depths < 70 m. The depth habitats utilized to derive *in situ* parameters are summarized in Table S7.

$3.9 \, \delta^{11} B_{borate}$

Two carbonate system parameters are needed to fully constrain the carbonate system. Following the approach of Foster et al., (2008) we used the GLODAP database (Key et al., 2004) corrected for anthropogenic inputs in order to estimate pre-industrial carbonate system parameters at each site. Temperature, salinity and pressure for each site are from the World ocean database 2013 (Boyer et al., 2013). We utilized the R^{\odot} code in Henehan et al, (2016) (courtesy of Michael Henehan) to calculate the $\delta^{11}B_{borate}$ and derive our calibrations. Uncertainty for $\delta^{11}B_{borate}$ utilizing the code was similar to the one calculated by applying 2 standard deviations of the calculated $\delta^{11}B_{borate}$ within the limits imposed by the calcification depth.

The Matlab[©] template provided by Zeebe and Wolf-Gladow, (2001) was used to calculate pCO₂ from TA; temperature, salinity and pressure were included into the calculations. Total boron was calculated from Lee et al., (2010), K_1 and K_2 were calculated from Mehrbach et al. (1973) refitted by Dickson and Millero (1987).

Statistical tests were made utilizing GraphPad[©] software, linear regressions for calibration where derived utilizing R[©] code in Henehan et al, (2016) (courtesy of Michael Henehan) with a k=500.

4. Results

4.1 Depth habitat

The calcification depths utilized in this paper are summarized in Tables S6 and S7, including a comparison of calcification depth determination methods. The calculated calcification depths are consistent with the ecology of each species and the hydrography of the sites. Specimens of *G. ruber* and *T. sacculifer* appear to be living in the shallow mixed layer (0-100 m), with *T. sacculifer* living or migrating deeper than *G. ruber* (down to 125 m). Specimens of *O. universa* and *P. obliquiloculata* are living in the upper thermocline; *G. menardii* is found in the upper thermocline until the thermocline depth specific to the location; *N. dutertrei* is living around the thermocline depth and specimens of *G. tumida* are found in the lower thermocline.

Data from both approaches implies that some species inhabit deeper environments in the Western Equatorial Pacific (WEP) relative to the Arabian Sea, which in turn are deeper dwelling than in the Indian Ocean. In some cases, we find evidence for differences in habitat depth of up to ~100m between the WEP and the Arabian Sea. This trend is observed for *G. ruber* and *T. sacculifer*, but not for *O. universa*.

Some differences in calcification depth are observed between the two calcification depth determination methods. These differences might be due to the choice of calibrations. Alternatively, our uncertainties for $\delta^{18}O$ implies larger uncertainties on the calcification depth determination using this approach, compared to Mg/Ca measurements.

4.2 Empirical calibrations of foraminiferal $\delta^{11}B_{carbonate}$ to $\delta^{11}B_{borate}$

Results for the different species analyzed in this study are presented in Fig. 5, Fig. 6 and summarized in Table 2; additionally, published calibrations for comparison are summarized in Table 3.

4.2.1 G. vai er

Our results for *G. ruber* (Fig. 5) are in good agreement with published data from other core-tops, sediment traps, tows, and culture experiments for $\delta^{11}B_{borate}>19$ % (Foster et al., 2008, Henehan et al., 2013, Raitzsch et al., 2018). However, for $\delta^{11}B_{borate}<19$ % our results show lighter $\delta^{11}B_{carbonate}$ compared to published values. Whilst this species has been widely studied previously, the sites selected in this study allow us to extend the calibration. The positive offset from the 1:1 curve has been explained by the high photosynthetic activity (Hönisch et al., 2003; Zeebe et al., 2003). Two calibrations have been derived. Utilizing only our data, the sensitivity of $\delta^{11}B_{carbonate}$ to $\delta^{11}B_{borate}$ of our linear regression is not statistically different from 1, the uncertainty on this regression is important due to our small dataset and not inconsistent with the low sensitivity trend of the culture experiments from Sanyal et al., (2001) or Henehan et al., (2013). The second calibration made compiling all data from literature shows a sensitivity similar (e.g. 0.46 (\pm 0.34) to the one recently published by Raitzsch et al., (2018) (e.g. 0.45 (\pm 0.16), Table 3).

4.2.2 T. sacculifer

 $\delta^{11}B_{carbonate}$ results for *T. sacculifer* (sacc and w/o sacc) (Fig. 5) are compared to published data (Foster et al., 2008; Martinez-Boti et al., 2015b, Raitzsch et al., 2018). Results for *T. sacculifer* are in good agreement with the literature and fall above the 1:1 line. Linear regression on our data yields a slope of 1.3 ± 0.2 but is not statistically different to the results from Martinez-Boti et al., (2015b) (Table 3), (p>0.05). However, when compiled with published data using the bootstrap method a slope of 0.83 ± 0.48 is calculated, with a large uncertainty given the variability in the data. It is also noticeable that *T. sacculifer* (w/o sacc) samples from the WEP have a $\delta^{11}B_{carbonate}$ close or below the 1:1 line and are significantly lower compared to the combined *T. sacculifer* of other sites (p=0.01, unptiled d t-test).

4.2.3 O. universa and deeper dwelling species: N. dutertrei, P. obliquiloculata, G. menardii and G. tumida

Our results for *O. universa* (Fig. 5), *N. dutertrei*, *P. obliquiloculata*, *G. menardii* and *G. tumida* (Fig. 6) fall below the 1:1 line. These data for *O. universa* are not statistically different from the Henehan et al. (2016) calibration (p>0.05). Our results for *N. dutertrei* expand upon the initial measurements presented in Foster et al., (2008). The different environments experienced by *N. dutertrei* in our study permit us to extend the range and derive a calibration for this species; the slope is close to unity (0.93 \pm 0.55), and is similar to the (0.95 \pm 0.17) previously reported by Henehan et al., (2016) for *O. universa* and not statistically different (p>0.05). The data for *P. obliquiloculata* exhibits the largest offset from the theoretical line. The range of δ^{11} B_{borate} from the samples we have of *G. menardii* and *G. tumida* is not sufficient to derive calibrations, but the points are in good agreement with the *N. dutertrei* calibration and Henehan et al. (2016) calibration for *O. universa*.

For all species, the slopes are not statistically different from Henehan et al. (2016) (p>0.05) and are close to unity. If data for deep-dwelling foraminiferal species are pooled together with each other and with data from Henehan et al., (2016) and Raitzch et al., (2018), we calculate a slope of 0.95 (\pm 0.13) (R²=0.7987, p<0.0001); if only our data are used, we calculate a slope that is not significantly different (0.82 \pm 0.27; p<0.05). However, it may remain premature to assume that a unique calibration with a slope of ~0.9 can be used for all deeper-dwelling species; more data is needed for *P. obliquiloculata*, *G. menardii* and *G. tumida* to robustly test this assertion.

4.2.4 Comparison of core-top and culture data

The data for *G. ruber* and *T. sacculifer* from the core-tops we measured are broadly consistent with previous published results. The calibrations between these core-top derived estimates and culture experiments are not statistically different due to small datasets and uncertainties on the linear regressions (Henehan et al., 2013; Marinez-Boti et al., 2015; Raitzsch et al., 2018; Table 3). The sensitivities of the species analyzed are not statistically different and are close to unity.

4.3 B/Ca ratios

B/Ca ratios are presented in Table 2 and Fig. 7. Values are species-specific consistent with previous work (e.g., compiled in Henehan et al., 2016) with ratios higher for G. ruber > T. sacculifer > T. sacculifer (w/o sacc) > P. obliquicloculata > O. universa > > G. menardii > N. dutertrei > G. tumida > G. inflata > N. pachyderma > G. bulloides (Fig. 7). This study supports interspecific B/Ca ratios (Yu et al., 2007; Tripati et al., 2009, 2011; Allen and Hönisch, 2012; Henehan et al., 2016). Differences between surface- and deep-dwelling foraminifera are observed, with lower values and a smaller range for the deeper dwelling taxa (58-126 µmol/mol vs 83-190 µmol/mol for shallow dwellers), however, the trend for the surface-dwellers can also be driven by interspecies B/Ca variability. The B/Ca data for deep-dwelling taxa exhibits a significant correlation with $[B(OH)_4]/[HCO_3]$ (p<0.05), but no correlation with $\delta^{11}B_{carbonate}$ and temperature (Fig. S3). Surface-dwelling species have B/Ca ratios that exhibit significant correlations with [B(OH)₄-]/[HCO₃-], δ^{11} B_{carbonate} and temperature. The sensitivity of B/Ca to [B(OH)₄-]/[HCO₃-] is lower for deep-dwelling species compared to surface dwelling species. When all the B/Ca data are compiled, significant trends are observed with [B(OH)4-]/[HCO₃-], δ^{11} B_{carbonate} and temperature (Fig. S3). We also observe that if we compare data from all sites together, correlations exist between B/Ca and the water depths of the cores (not significant, Fig. S4) but these correlations may also be related to the different the depth habitats of different taxa in each region, a significant trend is observed when all the data are plotted (R²=0.11, p<0.05, Fig. S4).

5. Discussion

5.1 Sources of uncertainty relating to depth habitat and seasonality at studied sites

5.1.1 Depth habitats and δ¹¹B_{borate}

Because foraminifera will record ambient environmental conditions during calcification, the accurate characterization of *in-situ* data is needed not only for calibrations, but also to understand the reconstructed record of pH or pCO₂. The species we examined are ordered here from shallower to deeper depth habitats: G. ruber > T. sacculifer (sacc) > T. sacculifer (w/o sac) > O. universa > P. obliquiloculata > G. menardii > N. dutertrei > G. tumida (this study; Birch et al., 2013; Farmer et al., 2007), although the specific water depth will vary depending on the hydrology of the site (Kemle-von and Oberhänhsli, 1999). We note that calculation of absolute calcification depths can be challenging in some cases as many species migrate during their ontogeny (Steinhardt et al., 2015).

We find that assumptions about the specific depth habitat a species of foraminifera is calcifying over, in a given region, can lead to differences of a few per mil in calculated isotopic compositions of borate (Fig. 4). Hence this can cause a bias in calibrations if calcification depths are assumed instead of being calculated (i.e., with δ^{18} O and/or Mg/Ca). Factors including variations in thermocline depth can impact depth habitats for some taxa. At the sites we examined, most of the sampled species live in deeper depth habitats in the WEP relative to the Indian Ocean, which in turn is characterized by deeper depth habitats than in the Arabian Sea. In the tropical Pacific, *T. sacculifer* is usually found deeper than *G. ruber* except at sites characterized by a shallow thermocline, in which case they tend to overlap their habitat (e.g., ODP Site 806 in the WEP which has a deeper thermocline than at ODP Site 847 in the Eastern Equatorial Pacific; EEP) (Rickaby et al., 2005). The difference in depth habitats for *T. sacculifer* and *N. dutertrei* between the WEP and EEP can be as much as almost 100 m (Rickaby et al., 2005).

5.1.2 Seasonality and *in-situ* δ¹¹B_{borate}

As discussed by Raitzsch et al., (2018), depending of the study area, foraminiferal fluxes can change throughout the year, so seasonality can have a major impact on hydrographic carbonate parameters calculations for any given water depth. We therefore recalculated the theoretical $\delta^{11}B_{\text{borate}}$ using seasonal data for temperature and salinity and annual values for TA and DIC for each depth at each site. The GLODAP (2013) database does not provide seasonal TA or DIC values.

The low sensitivity of $\delta^{11}B_{borate}$ to temperature and salinity means that calculated $\delta^{11}B_{borate}$ for each water depth at our sites were not strongly impacted (Fig. S1). Thus, these findings support Raitzsch et al. (2018), who concluded that calculated $\delta^{11}B_{borate}$ values corrected for seasonality was within error of non-corrected values for each water depth. As Raitzsch et al, (2018) highlight, seasonality might be more important at high latitude sites where seasonality is more marked, however, the seasonality of primary production will also be more tightly constrained due to the seasonal progression of winter light limitation and intense vertical mixing and summer nutrient limitation.

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Data for our sites suggests that most $\delta^{11}B_{borate}$ variability we observe does not come from seasonality but from the assumed water depths for calcification. With the exception of a few specific areas such as the Red Sea (Henehan et al., 2016, Raitzsch et al., 2018), at most sites examined, seasonal $\delta^{11}B_{borate}$ at a fixed depth does not vary by more than ~0.2‰. We conclude that seasonality is not an important factor impacting carbonate system parameters at the sites we examined.

5.2 δ¹¹B, microenvironment pH and depth hadiates

In planktonic foraminifera, algal symbiosis is the more common symbiotic relationship. For most of planktonic foraminifera, the host presents only one species of symbionts (Gast and Caron, 2001). The family Globigerinidae, including *G. ruber*, *T. sacculifer* and *O. universa*, commonly have dinoflagellates or chrysophyte algal symbionts (Anderson and Be, 1976; Spero, 1987). The families Pulleniatinidae, Globorotaliidae, including *N. dutertrei*, *P. obliquiloculata*, *G. menardii* and *G. tumida*, have chrysophyte algal symbionts (Gastrich, 1988).

The relationship between the symbionts and the host is complex by nature. Nevertheless, this symbiotic relationship provides energy (Hallock, 1981b) and promotes calcification of the foraminifera (Duguay, 1983;

Erez et al., 1983) by providing the inorganic carbon to the host (Jorgensen et al., 1985). Also, for *T. sacculifer* and *O. universa* photosynthesis increases with higher insolation (Jorgensen et al., 1985; Rink et al., 1998).

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Dinoflagellate-bearing foraminifera (*G. ruber*, *T. sacculifer* and *O. universa*) tend to have a higher symbiont density and photosynthesis activity while *P. obliquiloculata*, *G. menardii* and *N. dutertrei* have lowered symbiont density and *P. obliquiloculata*, *N. dutertrei* lower photosynthetic activity (Takagi et al., 2019). *P. obliquiloculata* showed the minimum symbiont density and photosynthetic activity (Takagi et al., 2019).

It is now accepted that the foraminifera $\delta^{11}B$ signature comes from the microenvironment pH (Jorgensen et al., 1985; Rink et al., 1998; Köhler-Rink and Kühl, 2000, Hönisch et al., 2003; Zeebe et el., 2003). Foraminifera with high photosynthetic activity and symbiont density like *G. ruber* and *T. sacculifer* present a pH of microenvironment higher than ambient seawater, $\delta^{11}B$ higher than 1:1 line (Foster et al., 2008, Henehan et al., 2013, Raitzsch et al., 2018). The opposite can also be true, from our study, species with lower photosynthetic activity and lower symbiont density present microenvironments lower than ambient seawater, $\delta^{11}B$ lower than 1:1 line (Martinez-Boti et al., 2015b; Henehan et al., 2016), this is the case in our data for *N. dutertrei*, *G. menardii* and *P. obliquiloculata* and likely *G. tumida*. Nevertheless, the low $\delta^{11}B$ of *O. universa* and *T. sacculifer* (*w/o sacc*) from the WEP are difficult to reconcile with a high photosynthetic activity compared to *T. sacculifer* et *G. ruber*.

The photosynthetic activity is also function of the light level they received which is, in the natural system, dependent of their depth in the water column, for the purpose of this study we will not consider turbidity which also influences the light penetration in the water column. In this case, the photosynthetically active for aminiferal iving close to the surface should see their microenvironment pH (thus δ^{11} B) more sensitive to water depth changes. A deeper depth habitat will change the light intensity they received and as a consequence may lower their photosynthetic activity reducing their microenvironment pH. This thought is supported by the significant trend observed between our Δ^{11} B and the calcification depth for G. ruber and T. sacculifer of our sites (Fig. S2). This trend basically supports the fact that the microenvironment pH decrease with calcification depth. We observe a decrease of δ^{11} B in the WEP for T. sacculifer (w/o sacc), significantly different from the other sites (p<0.05). The Δ^{11} B of G. ruber, T. sacculifer (w/o sacc and sacc) is also significantly lower in the WEP compared to the other sites (p<0.05). To test if the δ^{11} B signature was inferred to a light driven, we have been able to independently calculate the depth of the foraminifera based on various light insolation culture experiments (Jorgensen et al., 1985) and the Δmicroenvironment pH derived from our data (Fig. 8A and B). This exercise verified that this low δ^{11} B can be explained by the reduced light environment due to a deeper depth habitat in the WEP (Fig. 8B). Also, T. sacculifer has the potential to support more photosynthesis due to its higher symbiont density. Higher photosynthetic activity is observed compared to other species potentially supporting higher symbiont/host interactions. Those results could be in line with a greater sensitivity of T. sacculifer photosynthetic activity with changes in insolation/water depth. It can also be noted that this species presents the largest variations in symbiont density versus its test size. When applied to the other species O. universa data suggest a microenvironment pH 0.10 to 0.20 lower than ambient seawater pH which would be in line with species living deeper than 50m (light compensation point (Ec), Rink et al., 1998) also consistent with our calcification depth reconstructions. Δ microenvironment pH is higher in T. sacculifer > G. ruber > T. sacculifer (w/o sacc - WEP) > O. universa, N. dutertrei, G. menardii, G. tumida > P. obliquiloculata in line with photosymbiosis findings from Tagazaki et al., (2019). Also, the higher δ^{11} B data from the African upwelling published by Raitzsch et al., (2018) for *G. ruber* and *O. universa* might reflect the higher microenvironment pH due to a shallower depth habitat. This could highlight a potential issue with calibration when applied to sites with different oceanic regimes as the $\delta^{11}B$ specie-specific calibrations could be also location-specific for the mixed dweller species.

Microenvironment pH results for *N. dutertrei*, *G. menardii*, *G. tumida*, are similar to *O. universa* and suggest a threshold for respiration driven δ^{11} B signature. This threshold can be driven by a change of photosynthetic activity due to lower light intensity at deeper depth and/or a change in the symbiont assemblage with non-dinoflagellate symbionts at deeper depth. We can explain this threshold because deep dweller species do not experience important changes of insolation at those depths so their microenvironments should be respiration driven and relatively stable. We can also note that *P. obliquiloculata* which has the lowest symbiont density and photosynthetic activity has the lowest microenvironment pH compared to other deeper dweller species supporting this respiration driven microenvironment.

5.3 $\delta^{11}B$ sensitivity to $\delta^{11}B_{borate}$ and relationship with B/Ca signatures

 $\delta^{11}B_{carbonate}$ and B/Ca data have shown to be sensitive to precipitation rate with at higher precipitation rate increasing $\delta^{11}B_{carbonate}$ (Farmer et al., 2019) and B/Ca (Farmer et al., 2019; Gabitov et al., 2014; Kaczmarek et al., 2016; Mavromatis et al., 2015; Ushikawa et al., 2015). A recent study from Farmer et al, (2019) has proposed that in foraminifera at higher precipitation rates, more borate ion is incorporated into the carbonate mineral, while at lower precipitation rates, more boric acid is incorporated. They also suggest this may explain low sensitivities of culture experiments.

When combining all literature data, *T. sacculifer* and *G. ruber* have sensitivities of $\delta^{11}B_{carbonate}$ to $\delta^{11}B_{borate}$ of 0.83 ± 0.48 and 0.46 ± 0.34 respectively in line with previous literature and paleo-CO₂ reconstructions. Also, if we only take into account our data, the observation that the sensitivity of $\delta^{11}B_{carbonate}$ to $\delta^{11}B_{borate}$ are not statistically different from unity for most of the species investigated we can speculate that for these taxa, changes in precipitation rate and contributions of boric acid are not likely to be important. If considering only the data from this study, *G. ruber* (1.12 ± 1.67) and *T. sacculifer* (1.38± 1.35) present higher sensitivities of $\delta^{11}B_{carbonate}$ to $\delta^{11}B_{borate}$. We can then again speculate that the observed high values for $\delta^{11}B_{carbonate}$ at high seawater pH can be due to higher precipitation rates. We note this could also be consistent with the higher sensitivity of B/Ca signatures in these two surface dwelling species to ambient [B(OH)₄]/[HCO₃] relative to deeper dwelling species. Those interspecific differences still remain to be explained, however, part of this variability is likely due to changes in the carbonate chemistry of the microenvironment resulting in changing competition between borate and bicarbonate. A caveat is that we can not exclude specific biological processes, and that in taxa with a non respiration-driven microenvironment, changes in day/night calcification ratios also impacting observed values. As indicated by Farmer et al., (2019), studies of calcite precipitation rates in foraminifera may help to improve our understanding of the fundamental basis of boron-based proxies.

5.4 Evaluation of species for pH reconstructions and water depth pH reconstructions

This data set allows us to reassess the utility of boron-based proxies for the carbonate system. The main interest with utilizing boron-based proxies relates to the reconstruction of past oceanic conditions - specifically pH and pCO₂. Mixed-layer species (eg. *G. ruber* and *T. sacculifer*) are potential archives for atmospheric CO₂

reconstructions. Other species can shed light on other aspects of the carbon cycle including the physical and biological carbon pumps.

There are a few main inferences we can make. When compiled with data from the literature, sensitivities of $\delta^{11}B_{carbonate}$ to $\delta^{11}B_{borate}$ for *G. ruber* and *T. sacculifer* are similar to previous studies (Martinez-Boti et al., 2015b; Raitzsch et al., 2018) which is also supporting of previous paleo-reconstructions. Our data also support the observations of Henehan et al., (2016) for *O. universa*.

In order to derive accurate reconstructions of past ambient pH and pCO₂, accurate species-specific calibrations need to be used that are constrained by core-tops or samples from similar types of settings (Fig. 9, 10, S6). Lighter δ^{11} B signatures in *T. sacculifer* (w/o sacc) are observed in the WEP, which may be explained by the deeper depth habitat for these taxa, where lower light levels might reduce symbiont photosynthetic activity. Also, correction will be needed for *T. sacculifer* (w/o sacc) in the WEP. When applying the calibrations n°2 and 4 to *T. sacculifer* and *G. ruber* (compilation of all data, Table 3) our data show more variability, especially for *G. ruber* which lead to the larger mismatch compared to *in-situ* parameters. Henehan et al., (2013) reported a lighter δ^{11} B with smaller test size, our sample add a weight/shell of $11 \pm 4 \mu g$ (n=4, SD) which, despite a narrow range, could still explain this variability. The higher divergence of reconstructed values from *in-situ* measurements are observed at site WPO7-01 for both *T. sacculifer* (w/o sacc) and *G. ruber*. More data would be needed to determine a proper correction for both species, coretop study will be determinant for future downcore reconstructions, especially in the WEP.

We also find that for two species, the boron proxy is a relatively straightforward recorder of ambient pH, with sensitivities close to unity for and O. universa, and N. dutertrei. There is also promise in using multiple species in a sample from different hydrographic regimes to reconstruct vertical profiles of pH and pCO₂. We are able to reproduce pH and pCO₂ profiles from multiple sites with different water column structures (Fig. 9) with those reconstructions within error of the *in-situ* values, for most sites. In o der to avoid circularity, to validate these calibrations, we recalculated ambient pH and pCO₂ by first excluding site-specific data and then recalculating species-specific calibrations, followed by application to each specific site. The comparison of the two methods does not show significant differences and validates the robustness of the calibrations (Fig. S5). We utilized the calibrations derived from our data for G. ruber (calibration n°1 and 2, Table 3), T. sacculifer (calibration n°3 and 4, Table 3), O. universa (calibration n°8, Table 3), for P. obliquiloculata (calibration n°11, Table 3), and for N. dutertrei, G. tumida and G. menardii the calibration made on the compilation of the deepdweller (calibration n°13, Table 3). Results are shown in Fig. 9 and evaluated in Fig. 10. For G. menardii, more data would be helpful to provide additional constraints. Results for G. ruber are the sparsest, potentially due to difference in test sizes (Henehan et al., 2013) or undocumented diagenetic effects. Results reaffirm the importance of working with narrow size fractions (Henehan et al., 2013) and the importance of core-top study to determine corrections.

6. Conclusions and future implications

 Our study has extended the boron isotope proxy with data for new species and sites. The work supports previous work showing that depth habitats of foraminifera vary depending on the oceanic regime, and this impacts boron isotope signatures. Low $\delta^{11}B$ values in the WEP compared to other regions for *T. sacculifer* (w/o sacc) may be explained by a reduction in microenvironment pH due to a deeper depth habitat associated with

reduced irradiance and thus photosynthetic activity. Those results might also highlight a potential need for studying core-tops in order to establish what factors are important to accurately develop reconstructions in different areas.

The sensitivity of $\delta^{11}B$ to pH is in line with previously published data for *T. sacculifer*, *G. ruber*. The sensitivity of $\delta^{11}B$ to pH of *O. universa* (mixed-dweller), *N. dutertrei*. *G. menardii* and *G. tumida* (deep-dwellers) are similar but more data are needed to fully determine those sensitivities. The similarity of boron isotope calibrations for deep-dwelling taxa might be related to respiration-driven microenvironments.

Reconstruction of seawater pH and carbonate system parameters is achievable using foraminiferal $\delta^{11}B$ but additional core-top and down-core studies reconstructing depth profiles will be needed in order to further verify those calibrations. Past pH and pCO₂ water depth profiles can potentially be created by utilizing multiple foraminiferal species in concert with taxa-specific calibrations for similar settings. This approach has much potential for enhancing our understanding of the past workings of the oceanic carbon cycle, and the biological pump.

Author contribution

- R.E and A.T. wrote the proposals that funded the work. A.T. and F.C. provided the samples. M.G., S.M. and
- A.T. contributed to the experimental design. A.V. helped for sample preparation. M.G. and S.M contributed to
- developing the method of boron isotope analysis. M.G. performed the measurements with assistance from S.M.
- M.G conducted the data analysis. M.G. drafted the paper, which was edited by all authors. Interpretation was led
- by M.G., A.T., S.M. with input from R.E., A.V. and F.C.

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Competing interests

The authors declare that they have no conflict of interest.

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973 Figure caption

974

975 Figure 1: Reactions governing dissolved inorganic carbon equilibria.

976

- 977 Figure 2: (A) Speciation of H_3BO_3 and H_4BO_4 as function of seawater pH (total scale), (B) $\delta^{11}B$ of dissolved 978 inorganic boron species as a function of seawater pH, (C) sensitivity of δ¹¹B of H₄BO₄ for a pH ranging from
- 7.6 to 8.4. T=25°C, S=35, δ^{11} B=39.61 % (Foster et al., 2010), dissociation constant $\alpha = 1.0272$ (Klochko et al., 979
- 980 2006).

981

982 Figure 3: Map showing locations of the core-tops used in this study (white diamonds). Red open circles 983 represent the sites used for in-situ carbonate parameters from GLODAP database (Key et al., 2004).

984

985 Figure 4: Pre-industrial data versus depth of the sites used in this study. The figure shows seasonal temperatures 986 (extracted from World Ocean Database 2013), density anomaly (kg/m³), pre-industrial pH and pre-industrial δ¹¹B of H_PO₄ (calculated from the GLODAP database and corrected for anthropogenic inputs). 987

988 989

Figure 5: Boron isotopic measurements of mixed-layer foraminifera plotted against the δ^{11} B_{borate} were characterized by determination of the calcification depth of the foraminifera, A) G. ruber, B) T. sacculifer, C) O. universa. Mono-specific calibrations are summarized in Table 3.

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Figure 6: Boron isotopic measurements of deep-dwelling foraminifera ($\delta^{11}B_{carbonate}$) plot against $\delta^{11}B_{borate}$. 993 994 δ^{11} B_{borate} were characterized by determining the calcification depth of foraminifera, A) *P. obliquiloculata*, B) *G.* 995 menardii, C) N. dutertrei, D) G. tumida and E) Compilation of deep dweller species. Mono-specific calibrations 996 are summarized in Table 3.

997

998 Figure 7: Boxplots of B/Ca ratios for multiple species, T. sacculifer (this study; Foster et al., 2008; Ni et al; 999 2007; Seki et al., 2010), G. ruber (this study; Babila et al., 2014; Foster et al., 2008; Ni et al., 2007), G. inflata, 1000 G. bulloides (Yu et al., 2007), N. pachyderma (Hendry et al., 2009; Yu et al., 2013), N. dutertrei (this study; 1001 Foster et al., 2008), O. universa, P.obliquiloculata, G. menardii, G. tumida (this study).

1002

1003 Figure 8: A) Boxplet showing the calculated microenvironment pH difference (Δmicroenvironment pH) 1004 between microenvironment and external pH based on the δ^{11} B data. P) This figure shows that a decrease in insolation can explain the low δ^{11} B from the WEP. Light penetration profile in the Western Pacific, with E₀ in 1005 1006 the WEP of 220 J.s-1.m-2 (Weare et al., 1981) and a light attenuation coefficient of 0.028 (Wang et al., 2008). 1007 Theoretical depths were calculated for a decrease in microenvironment pH of $\Delta pH_1 = -0.02$ (e.g. WP07-a); 1008 $\Delta pH_1 = -0.04$ (e.g. A14), $\Delta pH_2 = -0.06$ (e.g. 806A). Light penetration corresponding to Ec is ~12%, $\Delta pH_0 \sim 7\%$, 1009 $\Delta pH_1 \sim 5\%$, $\Delta pH_2 \sim 1\%$ respective depth are 75m, 90m, 110m and 150m. Grey band is the calcification depth of 1010 T. sacculifer (w/o sacc) utilized in this study.

Figure 9: Water depth pH profiles reconstructed at every site applying the mono-specific calibrations derived from our results (Table 3). Figure is showing measured $\delta^{11}B_{calcite}$, $\delta^{11}B_{borate}$ calculated according to different calibrations (see Table 3 and text), calculated pH based on $\delta^{11}B$ (pH $_{\delta11B}$) and pCO $_2$ calculated from pH $_{\delta11B}$ and alkalinity.

Figure 10: Evaluation of the reconstructed parameters, $\delta^{11}B_{borate}$, pH and pCO₂ versus *in-situ* parameter. The recalculated parameters are consistent with *in-situ* data, except for *G. ruber*. This variability might be explained by the different test sizes.

1020	Table caption
1021	
1022	Table 1: Box-core information
1023	
1024	Table 2: Analytical results of δ^{13} C, δ^{18} O, δ^{11} B and elemental ratios $L_{\bullet}^{\bullet\prime}$ Ca and $\Delta_{\bullet}^{\bullet\prime}$ Ca
1025	
1026	Table 3: Species-specific $\delta^{11}B_{carbonate}$ to $\delta^{11}B_{borate}$ calibrations from literature and from our data

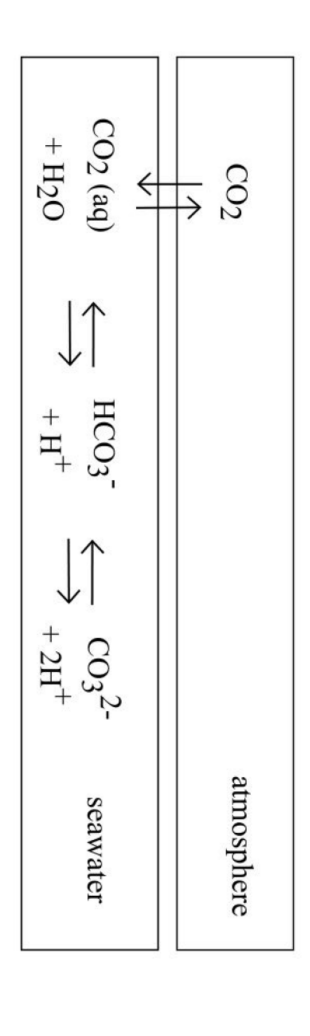


Figure 1

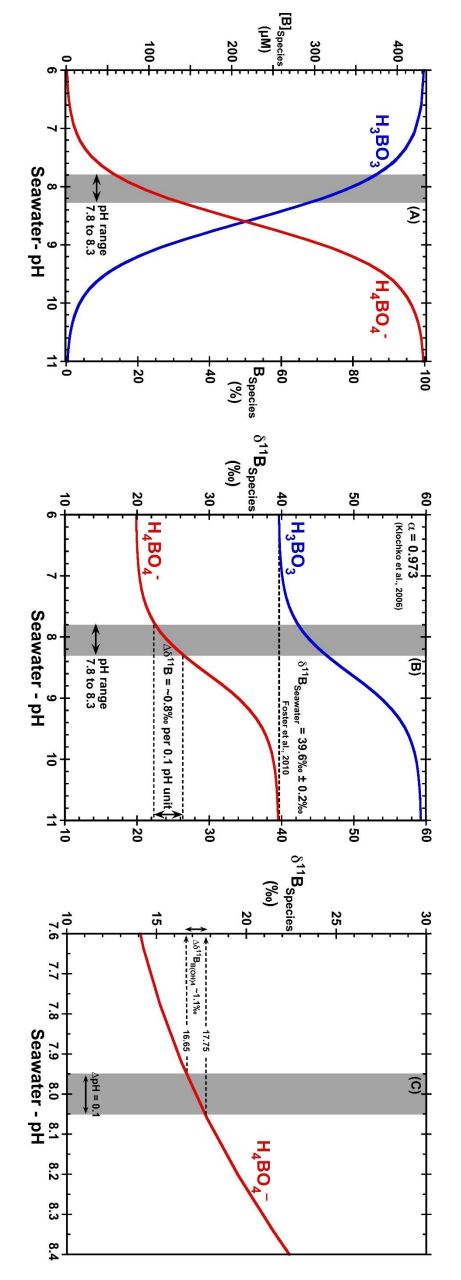


Figure 2

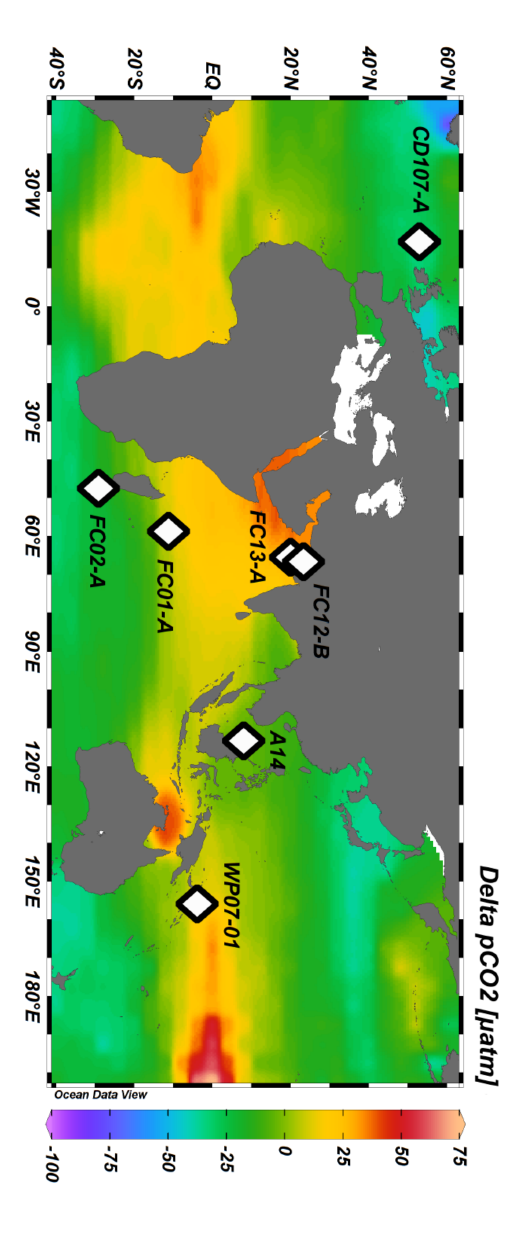
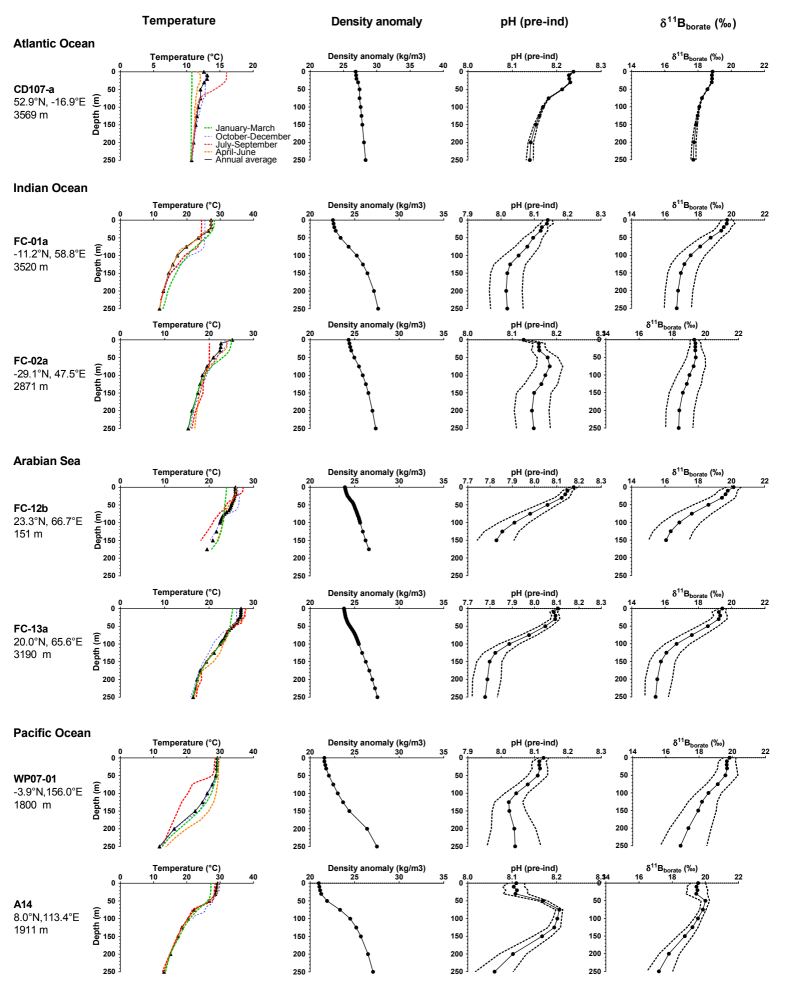
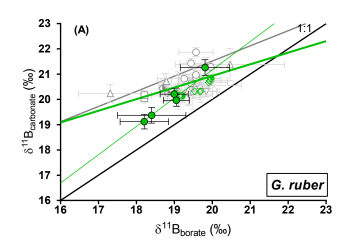
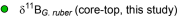


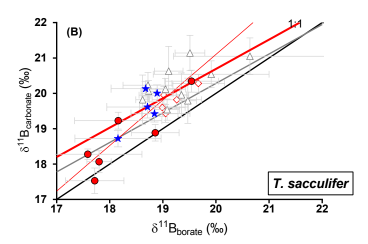
Figure 3



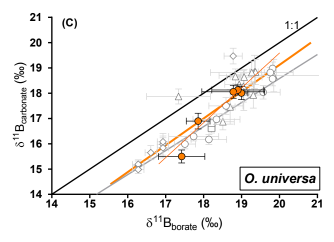




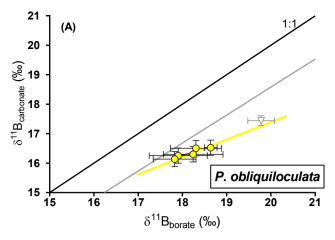
- δ $\delta^{11}B_{G, ruber}$ (core-top, Foster et al., 2008)
- $\delta^{11}B_{G. ruber}$ (core-top, 250-300µm, Henehan et al., 2013)
- δ^{11} B_{G. ruber} (core-top, Henehan et al., 2013)
- δ¹¹B_{G, ruber} (sediment trap, Henehan et al., 2013)
- $\delta^{11}B_{G, ruber}$ (tow, Henehan et al., 2013)
- ∇ $\delta^{11}B_{G, ruber}$ (grab sample, Henehan et al., 2013)
- \triangle $\delta^{11}B_{G. ruber}$ (Raizsch et al., 2018)
- G. ruber calibration line (All data, this study)
- G. ruber calibration line (Core-top, this study)
- G. ruber calibration line (Culture, Henehan et al., 2013)



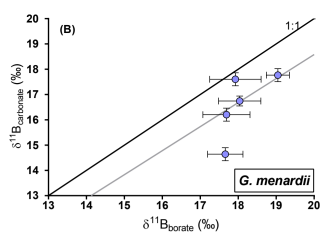
- δ¹¹B_{T.sacculifer (w/o sacc)} (core-top, this study)
- \triangle $\delta^{11}B_{T. sacculifer (w/o sacc)}$ (core-top, Raitzsch et al., 2018)
- ★ δ^{11} B_{T,sacculifer (sacc)} (core-top, this study)
- δ^{11} B_{T. sacculifer (sacc)} (core-top, Foster et al., 2008)
- T. sacculifer (w/o sacc and sacc) calibration line (All data, this study)
- T. sacculifer (w/o sacc and sacc) calibration line (Core-top, this study)
- T. sacculifer (s) calibration line (Martinez-Boti et al., 2015)



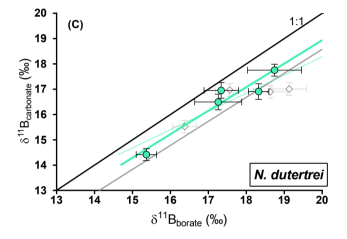
- $\delta^{11}B_{O. universa}$ (core-top, this study)
- \circ $\delta^{11}B_{O.\ universa}$ (core-top, Henehan et al., 2016)
- $\delta^{11}B_{O. universa}$ (sediment trap, Henehan et al., 2016)
- \diamond $\delta^{11}B_{O. universa}$ (tow, Henehan et al., 2016)
- \triangle $\delta^{11}B_{O. universa}$ (core-top, Raitzsch et al., 2018)
- O. universa calibration line (core-top, this study)
- O. universa calibration line (this study, Henehan et al., 2016, Raitzsch et al., 2018)
- O. universa calibration line (wild, Henehan et al., 2016)



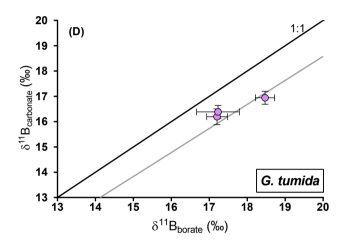
- δ¹¹B_{P.obliquiloculata} (Core-top, this study)
- ∇ δ^{11} B_{P.obliquiloculata}(Henehan et al., 2016)
- P. obliquiloculata calibration line (this study, Henehan et al., 2016)
- O. universa calibration curve (Henehan et al., 2016)



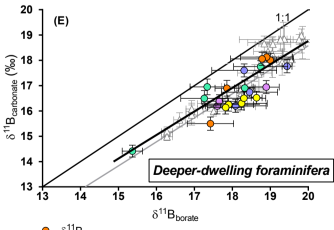
- δ¹¹B_{G. menardii} (this study)
 - O. universa calibration curve (Henehan et al., 2016)



- \bullet δ^{11} B_{N. dutertrei} (Core-top, this study)
- \diamond $\delta^{11}B_{N. dutertrei}$ (Core-top, Foster et al., 2008)
- O. universa calibration line (This study)
- O. universa calibration line (This study, Foster et al., 2008)
- O. universa calibration line (Henehan et al., 2016)

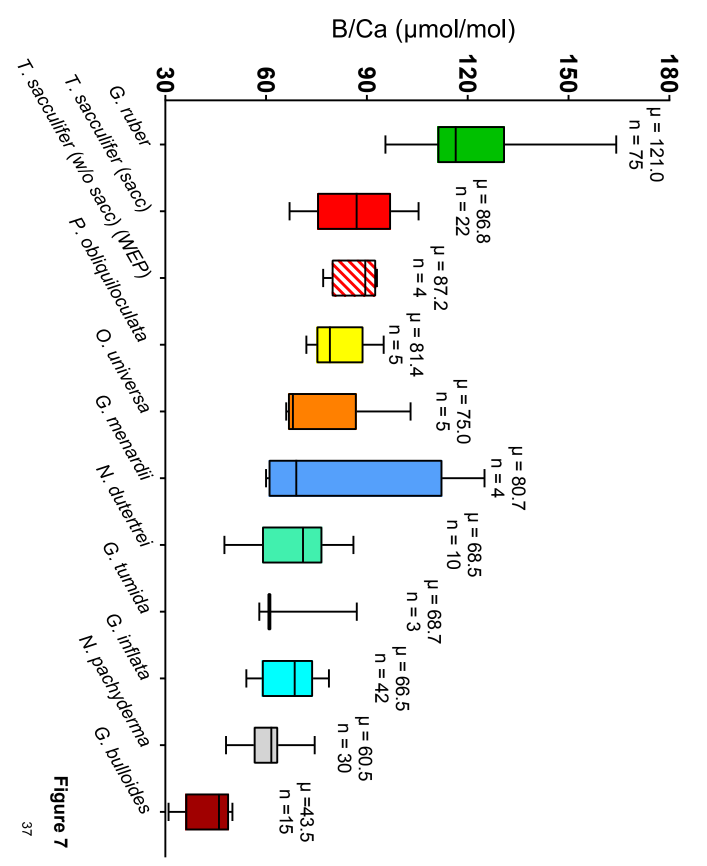


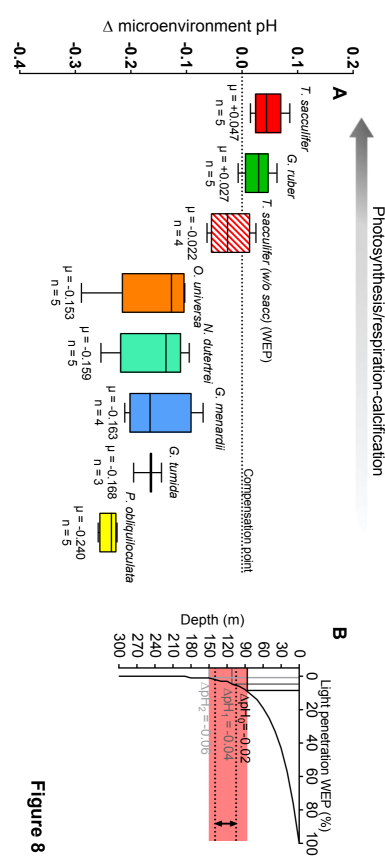
- \circ $\delta^{11}B_{G. tumida}$ (this study)
- O. universa calibration curve (Henehan et al., 2016)

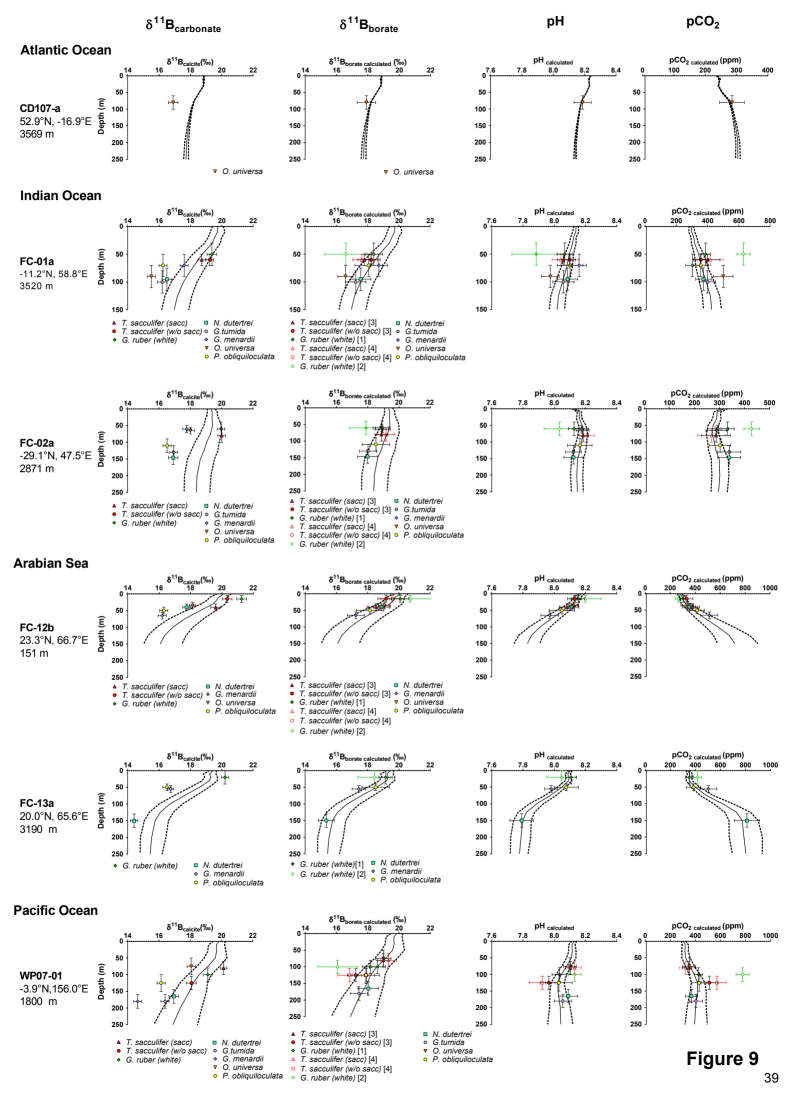


- δ¹¹B_{O.universa}
- \circ $\delta^{11}B_{P.obliquiloculata}$
- $\delta^{11}B_{N. dutertrei}$
- δ¹¹B_{G. menardii}
- O $\delta^{11}B_{G. tumida}$
- δ^{11} B_{deep-dweller} from literature
- Deep-dweller calibration line
- O. universa calibration line (Henehan et al., 2016)









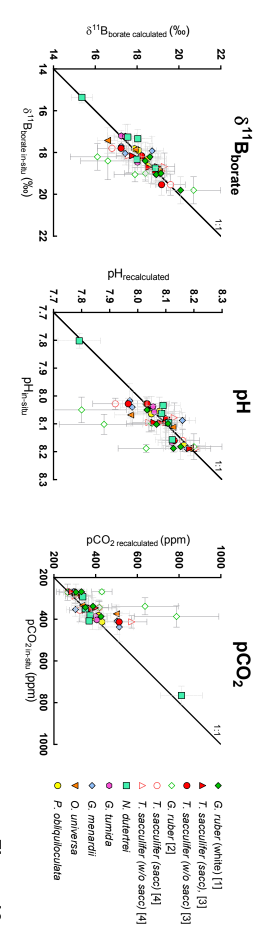


Figure 10₄₀

Table 1

Label	Box-Core	Site	Latitude (N)	Longitude (E)	(mbsl)	Oceanic Kegime	or Cage (year)
Atlantic Ocean	п						
CD107-a	CD107	Α	52.92	-16.92	3569	non-upwelling	<3000°a
Indian Ocean							
FC-01a	WIND-33B	Ι	-11.21	58.77	3520	non-upwelling	
FC-02a	WIND-10B	K	-29.12	47.55	2871	non-upwelling	7252 ± 27^b
Arabian Sea							
FC-12b	CD145	A150	23.30	66.70	151	seasonal upwelling	
FC-13a	CD145	A3200	20.00	65.58	3190	seasonal upwelling	
Pacific Ocean	7						
WP07-01			-3.93	156.00	1800	non-upwelling	$7.3-8.6^{\circ}$
A14			8.02	113.39	1911	non-upwelling	$7.3-8.6^{\circ}$
806		Α	0.32	159.36	2521	equatorial divergence	$7.3-8.6^{\circ}$
807		A	3.61	156.62	2804	equatorial divergence	$7.3-8.6^{\circ}$

^a Thomson et al., 2000

^b Wilson et al., 2012

^c Age for core-top of site 806B from Lea et al., 2000

Table 2

A14 A14 A14 A14 A14	807A	806A	Pacific Ocean WP07-a WP07-a WP07-a WP07-a WP07-a WP07-a WP07-a WP07-a WP07-a	FC-13a FC-13a FC-13a FC-13a	Arabian Sea RC-12b	PC-02a PC-02a PC-02a PC-02a PC-02a PC-02a PC-02a PC-02a	Indian Ocean RC-01a RC-01a RC-01a RC-01a RC-01a RC-01a RC-01a RC-01a RC-01a	Atlantic Ocean CD107a
G. ruber (white ss) T. sacculifer (sacc) T. sacculifer (w/o sacc) O. universa N. dutertrei	T. sacculifer (w/o sacc)	T. sacculifer (w/o sacc)	G. ruber (white ss) T. sacculifer (sacc) T. sacculifer (woo sacc) O.universa P. obliquiloculata G. menardii N. dutertrei G. tunnida	G. ruber (white ss) T. sacculifer (w/o sacc) P. obliquiloculata G. menardii N. dutertrei	G. ruber (white ss) G. sacculifer (s) T. sacculifer (w/o sacc) O. universa P. obliquiloculata G. menardii N. dutertrei	G. ruber (white ss) T. sacculifer (sacc) T. sacculifer (wo sacc) O. universa P. obliquiloculata G. menardii N. dutertrei G. tumida	G. ruber (white ss) T. sacculifer (sacc) T. sacculifer (woo sacc) O. universa P. obliquiloculata G. menardii N. dutertrei G. tumida	O. universa
250-400 250-400 250-400 500-560 250-400	250-400	250-400	250-400 250-400 250-400 500-630 250-400 250-400 250-400 250-400	250-300 300-400 300-400 300-400 300-400	250-300 300-400 300-400 300-400 >500 300-400 300-400 300-400	250-300 300-400 300-400 >500 300-400 300-400 300-400 300-400	250-300 300-400 300-400 300-400 >500 300-400 300-400 300-400	>500 Ox-Red
Ox-Red Ox-Red Ox-Red Ox-Red Ox-Red	Ox-Red	Ox-Red	Ox-Red	Ox-Red Ox-Red Ox-Red Ox-Red Ox-Red	Ox-Red Ox-Red Ox-Red Ox-Red Ox-Red Ox-Red Ox-Red Ox-Red	Ox-Red	Ox-Red	Ox-Red
				$\begin{array}{c} 0.08\pm0.03 & \textbf{-3.71}\pm0.11 \\ 1.59\pm0.03 & \textbf{-2.46}\pm0.11 \\ 0.00\pm0.03 & \textbf{-0.97}\pm0.11 \\ 0.75\pm0.03 & \textbf{-1.07}\pm0.11 \\ 0.71\pm0.03 & \textbf{-1.41}\pm0.11 \end{array}$	$\begin{array}{c} 0.58 \pm 0.03 \ \text{-}2.82 \pm 0.11 \\ 1.76 \pm 0.03 \ \text{-}2.15 \pm 0.11 \\ 1.97 \pm 0.03 \ \text{-}2.19 \pm 0.11 \\ 1.89 \pm 0.03 \ \text{-}1.59 \pm 0.11 \\ 1.89 \pm 0.03 \ \text{-}1.58 \pm 0.11 \\ 1.05 \pm 0.03 \ \text{-}0.37 \pm 0.11 \\ 1.35 \pm 0.03 \ \text{-}1.57 \pm 0.11 \\ 1.35 \pm 0.03 \ \text{-}1.57 \pm 0.11 \\ 1.35 \pm 0.03 \ \text{-}0.37 $	$\begin{array}{c} 0.30\pm0.03 -1.40\pm0.11 \\ 1.43\pm0.03 -1.60\pm0.11 \\ 1.52\pm0.03 -1.40\pm0.11 \\ 1.79\pm0.03 0.02\pm0.11 \\ 0.34\pm0.03 0.05\pm0.11 \\ 1.73\pm0.03 0.51\pm0.11 \\ 1.73\pm0.03 -0.51\pm0.11 \\ 1.03\pm0.03 -0.55\pm0.11 \\ 1.64\pm0.03 -0.28\pm0.11 \end{array}$	$1.37 \pm 0.03 \cdot 1.32 \pm 0.11$ $1.88 \pm 0.03 \cdot 2.20 \pm 0.11$ $2.02 \pm 0.03 \cdot 1.05 \pm 0.11$ $1.00 \pm 0.03 \cdot 0.55 \pm 0.11$ $1.64 \pm 0.03 \cdot 0.43 \pm 0.11$ $1.28 \pm 0.03 \cdot 0.43 \pm 0.11$ $1.29 \pm 0.03 \cdot 0.53 \pm 0.11$	(%o) (%o) (80) $1.99 \pm 0.03 1.25 \pm 0.11$
18.91 ± 0.24 (2SD, nAE121=10) 19.53 ± 0.24 (2SD, nAE121=10) 19.59 ± 0.24 (2SD, nAE121=10) 18.99 ± 0.24 (2SD, nAE121=11) 17.33 ± 0.26 (2SD, nAE121=11)	18.38 ± 0.21 (2SD, nAE121=11)	17.53 ± 0.36 (2SD, nAE121=11)	19.12 ± 0.29 (2SD, nAE121=12) 20.13 ± 0.21 (2SD, nAE121=11) 18.10 ± 0.31 (2SD, nAE121=11) 18.13 ± 0.26 (2SD, nAE121=14) 16.08 ± 0.26 (2SD, nAE121=14) 14.74 ± 0.26 (2SD, nAE121=14) 16.91 ± 0.31 (2SD, nAE121=14) 16.45 ± 0.26 (2SD, nAE121=14)	20.27 ± 0.24 (2SD, nAE121=10) 17.85 ± 0.29 (2SD, nAE121=12) 16.51 ± 0.26 (2SD, nAE121=14) 16.74 ± 0.20 (2SD, nAE121=6) 14.43 ± 0.24 (2SD, nAE121=10)	21.30 ± 0.31 (2SD, nAE121=11) 19.65 ± 0.31 (2SD, nAE121=11) 20.32 ± 0.31 (2SD, nAE121=1) 18.13 ± 0.20 (2SD, nAE121=0) 16.45 ± 0.26 (2SD, nAE121=14) 16.2 ± 0.26 (2SD, nAE121=14) 17.77 ± 0.24 (2SD, nAE121=14)	20.02 ± 0.24 (2SD, nAE121=10) 20.07 ± 0.24 (2SD, nAE121=10) 23.23 ± 0.24 (2SD, nAE121=10) 18.05 ± 0.26 (2SD, nAE121=14) 16.35 ± 0.26 (2SD, nAE121=14) 17.77 ± 0.26 (2SD, nAE121=14) 16.78 ± 0.31 (2SD, nAE121=14) 16.93 ± 0.26 (2SD, nAE121=14)	19.33 ± 0.31 (2SD, nAE121=11) 18.71 ± 0.24 (2SD, nAE121=10) 19.13 ± 0.24 (2SD, nAE121=10) 15.50 ± 0.26 (2SD, nAE121=14) 16.40 ± 0.26 (2SD, nAE121=14) 17.52 ± 0.26 (2SD, nAE121=14) 16.40 ± 0.31 (2SD, nAE121=14) 16.21 ± 0.31 (2SD, nAE121=11)	(%a) 16.85 ± 0.31 (2SD, nAE121=11)
19.17 ± 0.24 (2SD, nAE121=10) 19.32 ± 0.24 (2SD, nAE121=10) 18.84 ± 0.24 (2SD, nAE121=10) 17.08 ± 0.26 (2SD, nAE121=14)	18.17 ± 0.21 (2SD, nAE121=11)		18.04 ± 0.31 (2SD, nAE121=11) 17.99 ± 0.26 (2SD, nAE121=14) 16.19 ± 0.26 (2SD, nAE121=14) 14.53 ± 0.26 (2SD, nAE121=14) 16.99 ± 0.31 (2SD, nAE121=14) 16.32 ± 0.26 (2SD, nAE121=11)	$20.15 \pm 0.24 \text{ (2SD, nAE121=10)}$ $16.50 \pm 0.26 \text{ (2SD, nAE121=14)}$ $14.40 \pm 0.24 \text{ (2SD, nAE121=10)}$		19.90 ± 0.24 (2SD, nAE121=10) 19.93 ± 0.24 (2SD, nAE121=10) 23.22 ± 0.24 (2SD, nAE121=10) 17.97 ± 0.26 (2SD, nAE121=14) 16.69 ± 0.26 (2SD, nAE121=14) 17.93 ± 0.31 (2SD, nAE121=11) 16.95 ± 0.26 (2SD, nAE121=14)	19.41 ± 0.31 (2SD, nAE121=11) 18.73 ± 0.24 (2SD, nAE121=10) 19.32 ± 0.24 (2SD, nAE121=10) 16.10 ± 0.26 (2SD, nAE121=14) 17.69 ± 0.26 (2SD, nAE121=14) 16.59 ± 0.31 (2SD, nAE121=11) 16.18 ± 0.31 (2SD, nAE121=11)	(%a) (%a) (%a) (%a) (%b) (%b) (%b) (%b) (%b) (%b) (%b) (%b
19.04 ± 0.17 19.42 ± 0.17 18.88 ± 0.17 17.20 ± 0.18 14.39 + 0.31	18.28 ± 0.15	17.53 ± 0.36	19.12 ± 0.29 20.13 ± 0.21 18.07 ± 0.22 18.06 ± 0.18 16.14 ± 0.18 14.64 ± 0.18 16.95 ± 0.22 16.39 ± 0.18	20.21 ± 0.17 17.85 ± 0.29 16.51 ± 0.18 16.74 ± 0.20 14.41 ± 0.17		$\begin{array}{c} 19.96 \pm 0.17 \\ 20.00 \pm 0.17 \\ 23.22 \pm 0.17 \\ 18.01 \pm 0.18 \\ 16.52 \pm 0.18 \\ 17.77 \pm 0.26 \\ 16.91 \pm 0.22 \\ 16.94 \pm 0.18 \end{array}$	19.37 ± 0.22 18.72 ± 0.17 19.23 ± 0.17 15.50 ± 0.26 16.25 ± 0.18 17.60 ± 0.18 16.50 ± 0.22 16.20 ± 0.22	(%o) 16.90 ± 0.22
12.0 ± 0.4 12.3 ± 0.4 11.3 ± 0.4 16.9 ± 0.4	12.54 ± 0.4	14.40 ± 0.4	14.5 ± 0.4 12.7 ± 0.4 12.3 ± 0.4 11.9 ± 0.4 13.4 ± 0.4 13.5 ± 0.4 21.7 ± 0.4 10.6 ± 0.4	16.4 ± 0.4 15.7 ± 0.4 18.7 ± 0.4 18.7 ± 0.4 9.2 ± 0.4 15.7 ± 0.4	19.5 ± 0.4 14.6 ± 0.4 16.7 ± 0.4 13.6 ± 0.4 16.7 ± 0.4 14.8 ± 0.4 17.1 ± 0.4	18.2 ± 0.4 14.2 ± 0.4 13.7 ± 0.4 14.8 ± 0.4 16.6 ± 0.4 15.8 ± 0.4 18.6 ± 0.4 15.6 ± 0.4	15.4 ± 0.4 12.1 ± 0.4 12.1 ± 0.4 12.1 ± 0.4 15.4 ± 0.4 12.7 ± 0.4 18.6 ± 0.4 10.0 ± 0.4	$(\mu \text{mol/mol})$ 13.9 ± 0.4
102 ± 7 93 ± 7 66 ± 7	87 ± 7	77 ± 7	144 ± 7 92 ± 7 192 ± 7 71 ± 7 72 ± 7 85 ± 7 86 ± 7 58 ± 7	147 ± 7 121 ± 7 79 ± 7 60 ± 7 69 ± 7	164 ± 7 101 ± 7 1116 ± 7 1103 ± 7 95 ± 7 75 ± 7	$125 \pm 7 \\ 106 \pm 7 \\ 106 \pm 7 \\ 106 \pm 7 \\ 67 \pm 7 \\ 83 \pm 7 \\ 125 \pm 7 \\ 82 \pm 7 \\ 87 \pm 7$	109 ± 7 87 ± 7 82 ± 7 78 ± 7 63 ± 7 61 ± 7	(μ mol/mol 68 ± 7
3.91 ± 0.01 3.76 ± 0.01 6.59 ± 0.01	4.24 ± 0.01	3.89 ± 0.01	$\begin{array}{c} 4.32 \pm 0.01 \\ 4.44 \pm 0.01 \\ 4.51 \pm 0.01 \\ 7.52 \pm 0.01 \\ 3.02 \pm 0.01 \\ 2.68 \pm 0.01 \\ 3.66 \pm 0.01 \\ 2.55 \pm 0.01 \end{array}$	4.52 ± 0.01 5.49 ± 0.01 4.43 ± 0.01 1.99 ± 0.01 1.98 ± 0.01	5.76 ± 0.01 4.28 ± 0.01 4.90 ± 0.01 6.91 ± 0.01 3.61 ± 0.01 3.44 ± 0.01 3.25 ± 0.01	3.47 ± 0.01 3.30 ± 0.01 3.34 ± 0.01 4.40 ± 0.01 2.33 ± 0.01 2.21 ± 0.01 2.13 ± 0.01 1.90 ± 0.01	3.98 ± 0.01 3.45 ± 0.01 3.42 ± 0.01 2.06 ± 0.01 2.26 ± 0.01 1.81 ± 0.01 1.79 ± 0.01	(µmol/mol) (µmol/mol) (mmol/mol) (µmol/mol) (µmol/mol) (nmol/mol) (13.9 ± 0.4 68 ± 7 3.60 ± 0.01 13 ± 7 0.16 ± 0.01
22 ± 7 25 ± 7 10 ± 7	17 ± 7	7 ± 7	15 ± 7 22 ± 7 21 ± 7 11 ± 7 7 ± 7 26 ± 7 42 ± 7 16 ± 7	13 ± 7 21 ± 7 30 ± 7 19 ± 7 15 ± 7	14 ± 7 17 ± 7 20 ± 7 10 ± 7 69 ± 7 52 ± 7 46 ± 7	10 ±7 10 ±7 10 ±7 110 ±7 111 ±7 7 ±7 17 ±7 13 ±7 17 ±7	10 ±7 9 ± 7 14 ± 7 14 ± 7 14 ± 7 8 ± 7 11 ± 7	(µ mol/mol) 13 ± 7
0.02 ± 0.01 0.06 ± 0.01 0.02 ± 0.01 0.02 ± 0.01	0.09 ± 0.01	0.15 ± 0.01	$\begin{array}{c} 0.16 \pm 0.01 \\ 0.05 \pm 0.01 \\ 0.08 \pm 0.01 \\ 0.02 \pm 0.01 \\ 0.03 \pm 0.01 \\ 0.08 \pm 0.01 \\ 0.63 \pm 0.01 \\ 0.10 \pm 0.01 \\ \end{array}$	$\begin{array}{c} 0.08 \pm 0.01 \\ 0.49 \pm 0.01 \\ 0.43 \pm 0.01 \\ 0.07 \pm 0.01 \\ 0.06 \pm 0.01 \end{array}$	$0.16 \pm 0.01 \\ 0.14 \pm 0.01 \\ 0.26 \pm 0.01 \\ 0.06 \pm 0.01 \\ 0.38 \pm 0.01 \\ 0.17 \pm 0.01 \\ 0.25 \pm 0.01$	$\begin{array}{c} 0.07 \pm 0.01 \\ 0.03 \pm 0.01 \\ 0.04 \pm 0.01 \\ 0.05 \pm 0.01 \\ 0.03 \pm 0.01 \\ 0.03 \pm 0.01 \\ 0.07 \pm 0.01 \\ 0.04 \pm 0.01 \end{array}$	$\begin{array}{c} 0.07 \pm 0.01 \\ 0.03 \pm 0.01 \\ 0.03 \pm 0.01 \\ 0.05 \pm 0.01 \\ 0.07 \pm 0.01 \\ 0.07 \pm 0.01 \\ 0.02 \pm 0.01 \\ 0.02 \pm 0.01 \end{array}$) (mmol/mol) 0.16 ± 0.01

^{*} uncertainties given in ISD (see text)

When two measurements were carried out uncertaity was calculated with $\Delta a = \sqrt{(1/\sum_i (1/\Delta_{ii})^2)}$; with only one measurement the error was determined on reproducibility of the AE121 standard *Uncertainty given in 2SD, calculated on the reproducibility of CamWuellestorfi (see text and table 83, ref in Misra et al., 2014)

Table 3

Species	Size fraction (µm)	n) Material	Instrument (original)	Regression methor	Instrument (original) Regression method $\delta^{11}B_{borate} f(\delta^{11}B_{caldit})$	n Calibra	Calibration number Reference
G. ruber	~380	Culture/core tops/plankton tows	MC-ICP-MS		$\delta^{11}B_{borate} = [\delta^{11}B_{calcite} - 9.52 (\pm 2.02)]/0.6 (\pm 0.11)$		Henehan et al., 2013
G. ruber	315-355	Core-tops	MC-ICP-MS		$\delta^{11} \mathbf{B}_{\text{bornte}} = [\delta^{11} \mathbf{B}_{\text{calcite}} - 11.78 (\pm 3.20)]/0.45 (\pm 0.16)$		Raitzsch et al., 2018
T. sacculifer	n.d.	Culture/artificial seawater enriched in B	N-TIMS		$\delta^{11}B_{\text{bornte}} = [\delta^{11}B_{\text{calcite}} - 3.94 (\pm 4.02)]/0.82 (\pm 0.22)$		Sanyal et al., 2001 refitted Martinez-Boti et al., 2015
T. sacculifer	315-355	Core-tops	MC-ICP-MS		$\delta^{11}B_{\text{borate}} = [\delta^{11}B_{\text{calcite}} - 8.86 \ (\pm 5.27)]/0.59 \ (\pm 0.21)$		Raitzsch et al., 2018
O. universa	no effect	Core-tops/plankton tows/sediment traps	MC-ICP-MS		$\delta^{11} B_{\text{bornte}} = [\delta^{11} B_{\text{calcite}} + 0.42 (\pm 2.85)]/0.95 (\pm 0.17)$		Henehan et al., 2016
O. universa	>425	Core-tops	MC-ICP-MS		$\delta^{11} B_{\text{bornte}} = [\delta^{11} B_{\text{calcite}} + 5.69 (\pm 7.51)]/1.26 (\pm 0.39)$		Raitzsch et al., 2018
G. bulloides	300-355	Core-top/sediment trap	MC-ICP-MS		$\delta^{11}B_{\text{bornte}} = [\delta^{11}B_{\text{calcite}} + 3.440 (\pm 4.584)]/1.074 (\pm 0.252)$		Martinez-Boti et al., 2015
G. bulloides	315-355	Core-tops	MC-ICP-MS		$\delta^{11} \mathbf{B}_{\text{borate}} = [\delta^{11} \mathbf{B}_{\text{calcite}} + 3.81 (\pm 13.17)]/1.13 (\pm 0.72)$		Raitzsch et al., 2018
N. pachyderma	150-200	Core-tops	MC-ICP-MS		$\delta^{11}B_{borate} = \delta^{11}B_{calcite} + 3.38$		Yu et al., 2013
G. ruber	250-400	Core-tops	MC-ICP-MS	Bootstrap	$\delta^{11}B_{borate} = [\delta^{11}B_{calcite} + 1.23\ (\pm0.59)]/1.12\ (\pm1.67)$	5	1 This study
G. ruber	250-400	Core-tops	MC-ICP-MS	Bootstrap	$\delta^{11}B_{bornte} = [\delta^{11}B_{calcite} - 11.73 (\pm 0.83)]/0.46 (\pm 0.34)$	40	2 This study; Foster et al., 2008; Henehan et al., 2016; Raitzsch et al., 2018
T. sacculifer (sacc and w/o sacc)	250-400	Core-tops	MC-ICP-MS	Bootstrap	$\delta^{11}B_{bornte} = [\delta^{11}B_{calcite} + 6.06 (\pm 0.25)/1.38 (\pm 1.33)$	Ξ	3 This study
T. sacculifer (sacc and w/o sacc)	250-400	Core-tops	MC-ICP-MS	Bootstrap	$\delta^{11}B_{borate} = [\delta^{11}B_{calcite} - 4.09 (\pm 0.86)]/0.83 (\pm 0.48)$	27	4 This study; Foster et al., 2008; Raitzsch et al., 2018
N. dutertrei	300-400	Core-tops	MC-ICP-MS	Bootstrap	$\delta^{11}B_{bornte} = [\delta^{11}B_{calcite} - 0.34 (\pm 1.83)]/0.93 (\pm 0.55)$	5	5 This study
N. dutertrei	300-400	Core-tops	MC-ICP-MS	Bootstrap	$\delta^{11}B_{borate} = [\delta^{11}B_{calcite} - 3.88 (\pm 0.65)]/0.72 (\pm 0.74)$	9	6 This study; Foster et al., 2008
O. universa	400-600	Core-tops	MC-ICP-MS	Bootstrap	$\delta^{11} B_{borate} = [\delta^{11} B_{calcite} + 8.01 (\pm 23)]/1.38 (\pm 2.67)$	5	7 This study
O. universa	400-600	Core-tops	MC-ICP-MS	Bootstrap	$\delta^{11}B_{bornte} = [\delta^{11}B_{calcite} + 2.08 (\pm 0.59)]/1.06 (\pm 0.13)$	36	8 This study; Henehan et al., 2016; Raitzsch et al., 2018
G. menardii	400-600	Core-tops	MC-ICP-MS	Bootstrap	$\delta^{11}B_{borate} = [\delta^{11}B_{calcite} - 5.36 (\pm 1.36)]/0.65 (\pm 0.76)$	5	9 This study
G. tumida	400-600	Core-tops	MC-ICP-MS	Bootstrap	$\delta^{11}B_{borate} = [\delta^{11}B_{calcite} - 6.33 (\pm 2.52)]/0.57 (\pm 1.2)$	သ	10 This study
P. obliquiloculata	300-400	Core-tops	MC-ICP-MS	Bootstrap	$\delta^{11}B_{borate} = [\delta^{11}B_{calcite} - 5.59 (\pm 4.16)]/0.59(\pm 0.65)$	6	11 This study; Henehan et al., 2016
Deep-dweller	300-600	Core-tops	MC-ICP-MS	Bootstrap	$\delta^{11}B_{\text{borate}} = [\delta^{11}B_{\text{calcite}} - 1.99 (\pm 0.13)]/0.82 (\pm 0.27)$	22	12 This study
Deep-dweller	300-600	Core-tops	MC-ICP-MS	Bootstrap	$\delta^{11}B_{bount} = [\delta^{11}B_{culpit} - 0.18 (\pm 0.6)]/0.95 (\pm 0.13)$	2	13 This study; Foster et al., 2008; Henehan et al., 2016; Raitzsch et al., 2018

Supplemental information

Seawater pH reconstruction using boron isotopes in multiple planktonic foraminifera species with different depth habitats and their potential to constrain pH and pCO₂ gradients

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Supplemental Figures

Figure S1: An example of the impact of seasonality on results. Based on data from GLODAP used for site FC13-a. Seasonality has less of an impact that a change in the depth habitats.

Figure S2: Figure showing the offset $\Delta^{11}B=\delta^{11}B_{carbonate}-\delta^{11}B_{borate}$ versus calcification depth, red symbols are from Arabian Sea, green from Indian Ocean and Blue from the WEP, blue with black line symbols are data from site A14. This figure highlights a decrease of $\delta^{11}B_{carbonate}$ for *T. sacculifer* and *G. ruber* with a deeper depth habitat.

Figure S3: Multi-panels figure showing the correlation between B/Ca and boron geochemistry and different variables. A to C show comparison of B/Ca and A) $[B(OH)_4^-]/[HCO_3^-]$, B) $\delta^{11}B_{carbonate}$ and C) temperature. Panel D) shows the correlation between $\delta^{11}B_{carbonate}$ and temperature. Symbol in brackets is high B/Ca; this point is included in the linear regressions. Linea regression (LR - black line) is when compiling *G. ruber* and *T. sacculifer*, LR of *G. ruber* (dotted line green), *T. sacculifer* (red line – p<0.05), *T. sacculifer* (w/o sacc – red dotted line).

Figure S4: Boron geochemistry against water depth. A) $\delta^{11}B_{carbonate}$ versus water depth, B) B/Ca against water depth and C) $\delta^{11}B_{carbonate}$ versus calcification depth and linear regressions for *G. ruber*, *T. sacculifer* (w/o sacc), *T. sacculifer* (sacc) and *O. universa*.

Figure S5: Figure evaluating the circularity of our reconstructions. It is showing in the y-axis the difference between reconstruction utilizing calibrations derive from the entire dataset and compare to *in-situ* values and in the x-axis the difference between the reconstruction utilizing the species-specific calibrations derived excluding the site of interest (no circularity) compared to *in-situ* values. Results show that difference is not significant between the two reconstruction methods (e.g. following the 1:1 line), validating the method and the calibrations.

Supplemental Tables

Table S1: Elemental ratios of multi-elemental standards utilized in this study.

Table S2: Reproducibility of boron isotope standards.

 Table S3:
 Reproducibility of elemental ratios for CamWuellestorfi standard.

Table S4: Seasonality of foraminifera utilized in this study.

Table S5: Mg/Ca-T calibrations used for reconstructions and $\delta^{18}O_w$ -T calibrations used for calcification depth reconstructions.

Table S6: Calcification depth (CD) calculations from δ^{18} O (CD1), Mg/Ca (CD2) and literature (CD3).

Table S7: Pre-industrial in-situ parameters estimated using calcification depths for each species and calculated parameters based on analytical results.

Trace element standards

A series of multi-element standards (Table S1) with fixed Ca concentration and variable B, Mg, Sr, Mn, Ba, Zn, Cd, U, Li, Al and Fe concentrations were prepared for elemental ratios analysis in Brest following the method developed by Yu et al., (2005). Multi-element stock standard mixtures were prepared gravimetrically by spiking a 10,000 ppm Ca standard with appropriate amounts of Li, B, Al, Mn, Zn, Sr, Cd and U mono-elemental 1,000 ppm (SCP Science). They were diluted with OPTIMA grade HNO₃ acid and 18.2 M Ω .cm-1water to reach a 0.28M HNO₃ final solution. The stock standards (1500 ppm Ca) were prepared in 500mL cleaned PFA bottles. Working standards were made by diluting the stock solutions to a final concentration of 100 ppm Ca. The multi-element standards were calibrated at the University of Cambridge, elemental ratios are presented in Table S1. An external standard CamWuellestorfi (Misra et al., 2014a) was used in Brest for cross-calibration and reproducibility (Table S3).

Potential contaminations

Possible contamination of samples due to presence of silicate minerals was monitored with the Fe/Mg ratio. Samples with Fe/Mg > 0.1 mol/mol would be rejected due to potential contamination by silicate minerals (Barker et al., 2003). Samples (site E035 excluded) have an average Fe/Mg of 0.03 ± 0.05 mol/mol (2SD, n=42), meaning that silicate minerals have been efficiently removed during our cleaning.

Contribution of Mn-Fe-oxide coatings to Mg/Ca ratio has been calculated to be 0.5 μ mol/mol Mg/Ca (change for 5 μ mol/mol Mn/Ca ratio Barker et al., 2003). The maximum Mn/Ca ratio in our samples is 89 μ mol/mol which can lead to a potential contribution of ~9 μ mol/mol Mg/Ca or in other words a decrease of 0.1°C in our reconstructed temperatures. However, the calibration error is much larger and is estimated to be ~1.4°C (Dekens et al., 2002). The range in Mn/Ca values in our samples is 0.021 \pm 0.033 mmol/mol (2SD, n=42) which allows us to not be ignore *Mn-Fe-oxide* coating related complications. Additionally, no correlations were found between Mg/Ca and Fe/Ca (R²=0.006) or with Mn/Ca (R²=0.008), and between B/Ca ratio and Mn/Ca (R²=0.003) or Fe/Ca (R²=0.062) ratios.

Contamination by clays was monitored with Ti/Ca calculated from blank corrected intensities. Al/Ca ratios were not reliable as we are using an alumina injector for HF matrix in our lab. A minor correlation was found between Ti/Ca and Mg/Ca (R^2 =0.1388) but none with B/Ca (R^2 =0.0887).

Calcification depth determination

The first approach involves comparing measured $\delta^{18}O_c$ with theoretical predictions of $\delta^{18}O_c$ based on vertical profiles of temperature and the $\delta^{18}O$ of seawater ($\delta^{18}O_w$). We assume $\delta^{18}O_c$ is in equilibrium with seawater. First, $\delta^{18}O_w$ was calculated using location-specific $\delta^{18}O_w$ -salinity relationships and salinity profiles. We used salinity values from the World Ocean Atlas database (Boyer et al., 2013). Oxygen isotopes may be affected by both temperature and salinity. As our sites present different hydrographic settings and freshwater inputs, location-specific $\delta^{18}O_w$ -salinity relationship relationships are utilized for accurate $\delta^{18}O_w$ reconstructions. For Site CD107-a, we used a $\delta^{18}O_w$ -salinity relationship of 0.56*S-19.3 (Duplessy et al., 1991). For FC01-a and FC02-a, we used a $\delta^{18}O_w$ -salinity relationship of 0.24*S-7.8 (Sime et al., 2005), and for FC13-a and FC12-b, we used a $\delta^{18}O_w$ -salinity relationship of 0.28*S-9.24-0.27 (Rosteket al., 1993). Then, we used the calculated ambient $\delta^{18}O_w$ in concert with: (1) temperature profiles from the World Ocean Atlas database (Boyer et al., 2013), and (2) published calcite-water oxygen isotope fractionation factors, to calculate theoretical values for $\delta^{18}O_c$. Species-

specific relationships were used when available, including for *T. sacculifer* (Mulitza et al., 2003), *G. ruber* (Mulitza et al., 2003), and *O. universa* (Bemis et al., 2002, medium light). For all other, species we used the calcite equation from Kim and O'Niel (1997), adapted to a quadratic form by Bemis et al., (1998) following the approach of Sime et al., (2005). To take into account the ecology of each species, theoretical $\delta^{18}O_c$ profiles were made for the season of maximum abundance (Table S4). Therefore spring and summer profiles were used for *T. sacculifer*, summer profiles used for *G. ruber*, and winter and annual average profiles was used for *N. dutertrei*. Annual average profiles were used for the other species.

For the two sites WP07-1 and A14, a different approach was necessary because $\delta^{18}O_c$ data is sparse. At these sites, and for our other sites, we utilized Mg/Ca-derived temperatures to estimate calcification depths (Table S6). $T_{Mg/Ca}$ was derived using species-specific Mg/Ca-temperature calibrations (Table S6) along with the Mg/Ca ratios determined in this study. Calcification depth was estimated by comparing $T_{Mg/Ca}$ to modern temperature profiles from the World Ocean atlas database 2013 (Boyer et al., 2013) in light of the ecology (seasonality of growth) of the species of interest. A caveat is that in certain cases $T_{Mg/Ca}$ may be partially biased by a carbonate ion effect or salinity effect (Russell et al., 2004; Elderfield et al., 2006; Ferguson et al., 2008; Arbuszewski et al., 2010; Martinez-Boti et al., 2011). These artifacts on $T_{Mg/Ca}$ may be significant at high-latitude sites such as CD107-a which is located in the North Atlantic Ocean.

Depth habitat

Planktonic foraminifera live in the upper 500 m of the water column. Their preferred depth habitat depends on their ecology, which in turn relies on the hydrographic conditions. For example, *G. ruber is* commonly found in the mixed layer (Fairbanks and Wiebe, 1980; Dekens et al., 2002; Farmer et al., 2007) during summer (Deuser et al., 1981) whereas *T. sacculifer* (n or ns) is present in the mixed layer until the midthermocline depth (Farmer et al., 2007) during spring and summer (Deuser et al., 1981, 1989). Specimens of *P. obliquiloculata* and *N. dutertrei* are found during winter (Deuser et al., 1989), in the mixed layer (~60m) for *P. obliquiloculata*, and at mid-thermocline depth for *N. dutertrei* (Farmer et al., 2007). Whereas, *O. universa* tends to record annual average conditions and is living within the mixed layer. Specimens of *G. menardii* calcify within the seasonal thermocline (Fairbanks et al., 1982, Farmer et al., 2007, Regenberg et al., 2009) even upper thermocline (Farmer et al., 2007) and records annual temperatures. And specimens of *G. tumida* are found at the lower thermocline or below the thermocline and record annual average conditions (Fairbanks and Wiebe, 1980; Farmer et al., 2007, Birch et al., 2013). Our calcification depth reconstructions are summarized in Table 3, also see Table S6 for comparison.

Atlantic Ocean

Farmer et al., (2007) determined the depth habitat for *O. universa* to be ranging from 0 to 60m (LL, Bemis et al., 1998). Our calculation, through δ^{18} O measurement, suggests a deeper habitat of around 70m (LL, Bemis et al., 1998), 80m (ML, Bemis et al., 2002). Whereas, the Mg/Ca method derived depth habitat calculation yields a depth habitat of 50 m. The lower habitat depth can also come from the different size fractions, as our size fraction is lower than Farmer's. *O.universa* is thought to migrate to shallower depth along its ontogeny (Emiliani et al., 1954) younger individuals are thus living deeper but smaller individuals might also have a

deeper habitat as already suggested by Hönisch and Heming, (2004). Since most of the published studies have used the δ^{18} O-based depth calcification, we will preferentially adopt this method.

Indian Ocean

Calcification depths for the Indian Ocean cores have already been determined for majority of the species by Sime et al., (2005). Additionally, Birch et al., (2013) have reconstructed the depth habitat of multiple species from a core collected in the offshore region of Tanzania (Glow 3). In Indian Ocean specimens of G. ruber is found in the top 50m (Birch et al., 2013) and until 60 m (Sime et al., 2005), T. sacculifer is found in the surface mixed layer (SML) but also in the upper thermocline between 50-70 m for Birch et al., (2013) and between 60 to 80 m for Sime et al., (2005). Our results are consistent with these reported depth habitats. We calculate that specimens of G. ruber, T. sacculifer with and without sac are living in the top 80m; O. universa lives between 50 to 90 m (Sime et al., 2005, Birch et al., 2013). For N. dutertrei we calculate a depth habitat of 90m at site FC01a, calcification depths derived from both δ^{18} O and Mg/Ca methods agree with the 93 m estimate by Sime et al., (2005). At site FC02a the calculated calcification depth based on δ^{18} O method is 65 m, and, the Mg/Ca derived depth is 100m; however, Sime et al., (2005) proposed a calcification depth of 146 m. A deeper depth habitat than site FC01a seems to be in line with the weaker stratification of the water column at site FCO2a. The depth habitat for P. obliquiloculata's was determined to be 106-120m by Sime et al., (2005); however, our calculations predict a lower and narrower depth habitat of 60 - 70m. Calcification depth for G.menardii has been calculated to be 60 - 70m, which is consistent with an upper thermocline depth habitat. Specimens of G.tumida is commonly found at the bottom of the thermocline which is around 200-250m at our Indian Ocean sites, Birch et al., (2013) found this species ranging from 100 to 200 m but our δ^{18} O based habitat reconstruction suggests a shallower depth of 70m at both sites. It seems that the calcification derived Mg/Ca for G. tumida is more realistic than with our $\delta^{18}O_c$ reconstructions, the Mg/Ca based calculation predicts a depth habitat of 100 - 130 m.

Arabian Sea

In the Arabian Sea foraminifera are affected by changes in local hydrology caused by the Indian summer and winter monsoons. The specimens of different species collectively record a shallower habitat during SW monsoon (e.g., upwelling) and a deeper habitat during NE monsoon (Peeters and Brummer, 2002). Except for *N. dutertrei*, our two methods of calcification depth reconstruction closely agree with each other within their respective uncertainties. The depth habitat reconstructions for *G. ruber* and for *T. sacculifer* are consistent with species living in the shallow mixed layer (SML) at a depth of 30m for *G. ruber* and 40 m for *T. sacculifer*. Specimens of *O. universa* are calculated to live at 30m depth; *N. dutertrei* at site FC12-b (water depth 151m) is living around 40m and at 150m at site FC13-b (water depth 3200m), which is consistent with the hydrography at this site (Fig. 4). The depth habitat of *P. obliquiloculata* is calculated to be 50m at both sites. Specimens of *G. menardii* is found at depths of 60m, consistent with Peeters and Brummer, (2002) estimate of 50 – 130m.

Pacific Ocean

For Pacific Ocean samples the Mg/Ca derived calcification depths were used in absence of δ^{18} Oc values. The Sites WP07-01 and A14 are located in the Western Equatorial Pacific with Site WP07-01

characterized by a deep thermocline. At these sites *G. ruber* and *T. sacculifer* have deep depth habitat of around 100m for *G. ruber* (Elderfield and Ganssen, 2000) and around 125m for *T. sacculifer* (Rickaby et al., 2005). The depth habitat for specimens of *O. universa* was determined to be 75m depth at site WP07-01 and at 55m depth at site A14. We calculate that *P. obliquiloculata* is living at 125m deep. Rickaby et al., (2005) estimated the living depth of *N. dutertrei* at 165m in agreement with our calculated depths (~125m). Specimens of *G. menardii* were determined to live at 180m like *G. tumida* (Rickaby et al., 2005).

The data are not consistent with a dissolution effect to explain the low $\delta^{11}B$ T. sacculifer in the WEP

Documented dissolution effects have been attributed to the preferential dissolution of ontogenic calcite relative to the light $\delta^{11}B$ of gametogenic calcite (Ni et al., 2007; Seki et al., 2010; Henehan et al., 2016). *T. sacculifer* (w/o sacc) should be less impacted compared to the *T. sacculifer* (sacc). Hönisch and Hemming, (2004) and Ni et al., (2007) reported a dissolution effect at site 806 (close to our WP07-01 site) for *T. sacculifer* (sacc), however, in our data the $\delta^{11}B_{carbonate}$ for *T. sacculifer* (sacc) is not decreasing with water depth when *T. sacculifer* (w/o sacc) is which suggests that at these sites and at the same size-fraction no dissolution is observed (eg. different water depth). $\Delta^{11}B$ for *T. sacculifer* (w/o sacc) and for *G. ruber* shift to lighter with higher calcification depth, a trend that does not support a dissolution effect. The lethal temperature for *T. sacculifer* is $14^{\circ}C$, which at site WP07-01 corresponds to a $\delta^{11}B_{borate}$ of 17.5 % which makes our low $\delta^{11}B_{carbonate}$ realistic. If no dissolution is observed, a deep depth habitat inducing a respiration-driven microenvironment might explain the low $\delta^{11}B$ of the measured carbonate.

Microenvironment calculations

We observe a trend between $\Delta^{11}B$ (eg. $\Delta^{11}B=\delta^{11}B_{carbonate}-\delta^{11}B_{borate}$) with derived calcification depth (Fig. S2). In order to verify why the WEP δ^{11} B_{carbonate} of *T. sacculifer* (w/o sacc) is low and to test the hypothesis of the depth habitat we try to recalculate independently the theoretical water depth habitat based on culture results from Jorgensen et al., (1985) and our microenvironment pH results. A change of microenvironment pH for T. sacculifer will change the theoretical light intensity needed to reach this microenvironment pH. The compensation light intensity (E_c) for T. sacculifer has been calculated by Jorgensen et al., (1985) to be ~30 μ Eistn.m⁻².s⁻¹, E_c corresponds to the energy where photosynthesis compensates respiration or where δ^{11} B_{carbonate} reaches the 1:1 theoretical line. We tested two microenvironment pH, $\Delta pH_1 = -0.04$ and $\Delta pH_2 = -0.06$ (Fig S3). We've recalculated the light energy needed to decrease the pH of ΔpH_1 and ΔpH_2 and apply these changes to the light penetration profile determined with an insolation E₀ in the WEP of 220 J.s⁻¹.m⁻² (Weare et al., 1981) and a light attenuation coefficient of 0.028 (Wang et al., 2008). A decrease of ΔpH₁ would lead to a decrease of 15 μEistn.m⁻².s⁻¹ and a decrease of ΔpH₂ would lead to a decrease of 24 μEistn.m⁻².s⁻¹ (Jorgensen et al., 1985). These results correspond in our case of a light penetration of 12% to reach Ec, 5% for a decrease of ΔpH₁ and 1% for a decrease of ΔpH_2 . This means that in the WEP if T. sacculifer calcifies below 75m where E_c is reached the $\delta^{11}B_{carbonate}$ is below the theoretical 1:1 line. T. sacculifer (w/o sacc) in the WEP is decreasing its pH of $\sim \Delta pH_1$ which would imply a calcification depth of 110m consistent with the reconstruction of Rickaby et al., (2005).

$$\Delta microenvironment~pH = -log(\frac{(\delta 11Bseawater - \delta 11Bcarbonate)~x~Kb*}{\epsilon - \delta 11Bseawater + \delta 11Bcarbonate}) - pHseawater$$

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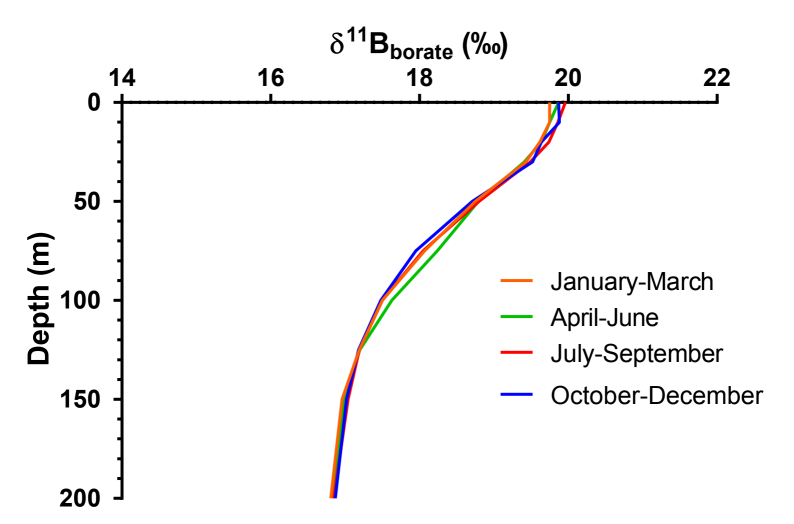


Figure S1

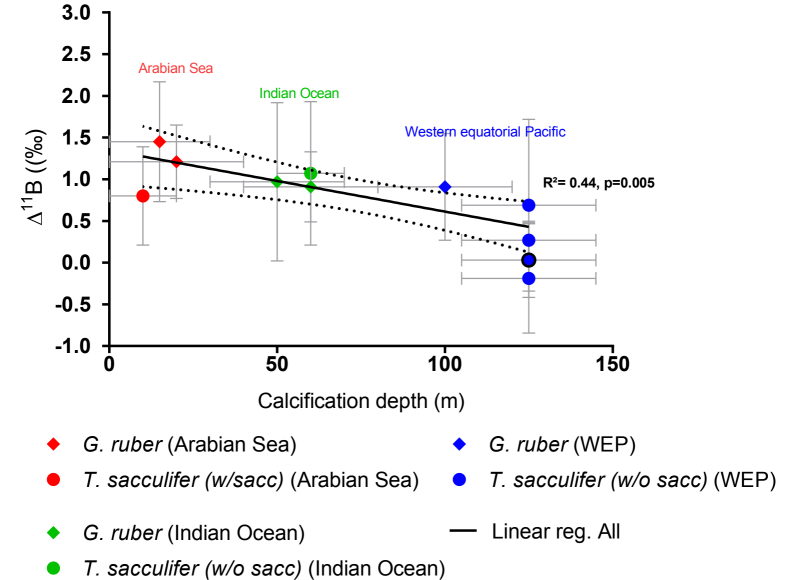
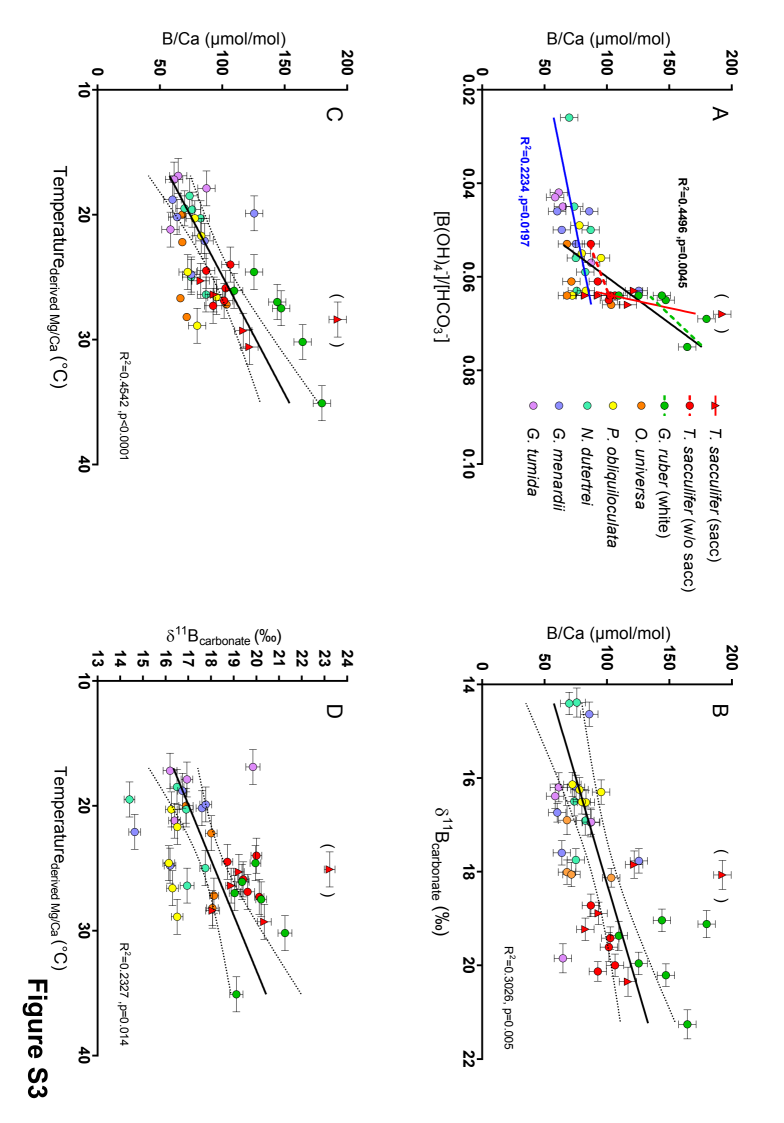
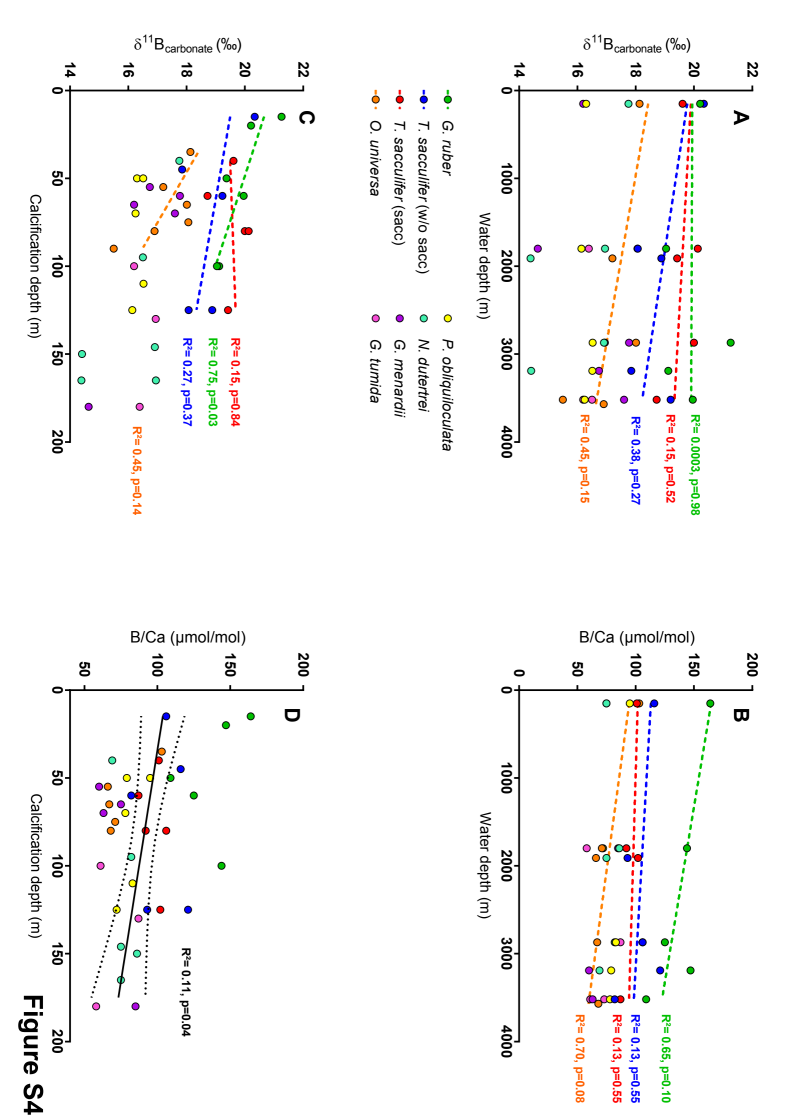


Figure S2





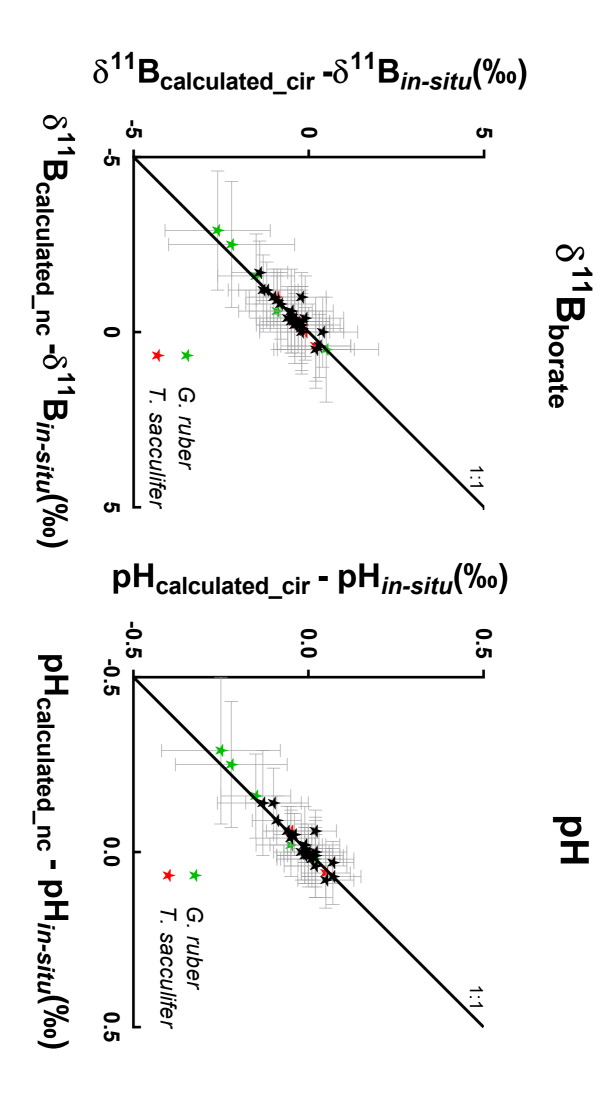


Figure S5

Table S1

Elemental ratios	Li/Ca	B/Ca	Mg/Ca	Al/Ca	Sr/Ca	Cd/Ca	Ba/Ca	U/Ca	7	In/Ca
	μmol/mol	μmol/mol	mmol/mol	mmol/mol	mmol/mol	μmol/mol	μmol/mol	nmol/mol		μmol/mol nmol/mol μmol/mol mmol/mol
Standard solution 0	0.8	9	0.10	0.131	0.00	0.03	0.6	31		1
Standard solution 1	2.3	38	0.31	0.112	0.49		1.9	38		12
Standard solution 3	6.8	108	1.31	0.177	1.06		3.0	53		39
Standard solution 5	14.6	216	3.17	0.223	1.57		5.1	62		129
Standard solution 6	19.0	278	5.23	0.352	1.97	0.28	5.5	74		196
Standard solution 8	25.0	281	6.07	0.602	2.99		20.1	390		501
Standard solution 9		408			4.89					
Standard solution 10		519			8.01					
Standard solution 11		607			9.93					

Table S2

JCP-1 JCP-1 JCP-1	NEP NEP	Standard	JCP-1-6	JCP-1-5	JCP-1-4	JCP-1-2	JCP-1-1	NEP22	NEP21	NEP20	NEP19	NEP18	NEP17	NEP16	NEP15	NEP14	NEP13	NEP12	NEP11	NEP10	NEP9	NEP8	NEP7	NEP6	NEP5	NEP4	NEP3	NEP2	NEP1	Standard
24.06 24.37 24.42	25.70 26.20 25.80	Average õ ¹¹ B	24.18	24.03	23.92	24.01	24.07	26.42	25.54	25.42	25.78	25.86	26.02	26.04	26.00	26.12	26.19	26.22	26.09	25.97	26.15	25.39	25.26	25.22	25.32	25.40	24.70	25.00	25 21	$\delta^{11}\mathbf{B}_{1}$ (‰)
0.20 0.32 0.28	0.93 0.88 0.89	2SD	0.36	0.26	0.26	0.11	0.10	0.22	0.22	0.15	0.26	0.29	0.29	0.26	0.26	0.26	0.26	0.26	0.26	0.26	0.26	0.26	0.26	0.21	0.21	0.21	0.30	0.30	0.25	$2SD_{\rm AE121}$
6 57 7	22 27 6	n						6	6	ယ	14	12	12	15	15	15	15	15	15	15	15	15	15	11	=	11	12	12		$\mathbf{n}_{\mathrm{AE121}}$
			24.16	24.05		£7.17	24 17		26.16	25.32		25.86				26.13	26.21	26.29	26.09						25.33				25 22	$\delta^{11}\mathbf{B}_{2}$ (%o)
			0.36	0.39		0.10	0 10		0.22	0.15		0.26				0.26	0.26	0.26	0.26						0.21			į	0.25	$2SD_{\rm AE121}$
									6	ω		14				15	15	15	15						=					n _{AE12}
This study Holcomb et al., 2015 Sutton et al., 2018	This study Holcomb et al., 2015 Sutton et al., 2018	Reference	This study	n _{AE121} Reference																										

Table S3

		10/7/17 10/7/17 10/7/17 10/7/17 10/7/17 10/7/17 11/7/17 11/7/17 11/7/17 11/7/17 11/7/17 11/7/17 11/7/17 11/7/17 11/7/17 11/7/17 11/7/17 12/7/17 12/7/17 12/7/17 12/7/17 12/7/17 12/7/17 12/7/17 12/7/17 12/7/17 12/7/17 12/7/17 12/7/17 12/7/17 12/7/17 12/7/17 12/7/2017 13/7/2017 13/7/2017 13/7/2017	Date of analysis
Published CamWuel 2SD n	Average CamWuel 2SD	CamWuel 1 CamWuel 2 CamWuel 3 CamWuel 4 CamWuel 6 CamWuel 7 CamWuel 7 CamWuel 1 CamWuel 2 CamWuel 3 CamWuel 5 CamWuel 6 CamWuel 6 CamWuel 7 CamWuel 7 CamWuel 7 CamWuel 7 CamWuel 7 CamWuel 8 CamWuel 3 CamWuel 4 CamWuel 4 CamWuel 5 CamWuel 5 CamWuel 6 CamWuel 7 CamWuel 7 CamWuel 7 CamWuel 8 CamWuel 9 CamWuel 9 CamWuel 1 CamWuel 9 CamWuel 1 CamWuel 2 CamWuel 3 CamWuel 3 CamWuel 6 CamWuel 6 CamWuel 7 CamWuel 7 CamWuel 7 CamWuel 8 CamWuel 9 CamWuel 1 CamWuel 1 CamWuel 2 CamWuel 3 CamWuel 3 CamWuel 3 CamWuel 4 CamWuel 5 CamWuel 6 CamWuel 6 CamWuel 7 CamWuel 7 CamWuel 7 CamWuel 8 CamWuel 9 CamWuel 9 CamWuel 9 CamWuel 1	Standard
16.4 0.8 180	16.2 0.4 31	16.0 16.3 16.0 16.0 16.0 16.0 16.0 16.0 16.0 16.0	Li/Ca µmol/mol
202 7 180	204 7 31	204 209 209 207 210 208 201 203 203 204 204 205 207 207 207 207 207 207 207 207 207 207	B/Ca μmol/mol
1.23 0.06 180	1.22 0.01 31		Mg/Ca mmol/mol
1.30 0.02 180	1.31 0.01 31	1 3 1 1 3 2 1 1 3 2 1 1 3 2 2 1 1 3 2 2 1 1 3 2 2 1 1 3 2 2 1 1 3 2 2 2 2	Sr/Ca mmol/mol
0.29 0.02 180	0.27 0.06 31	0.26 0.28 0.28 0.26 0.27 0.27 0.27 0.27 0.27 0.27 0.27 0.22 0.23 0.23 0.23 0.23 0.23 0.23 0.23	Cd/Ca µmol/mol
4.8 <i>0.2</i> 180	4.6 0.1 31	4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	Ba/Ca μmol/mol
44 2 180	42 2 31	\$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$	U/Ca nmol/mol
71 2 180	71 7 31	67 74 74 74 74 75 76 76 76 76 76 76 76 76 76 76 76 76 76	Mn/Ca μmol/mol
	0.04 0.01 31	0.04 0.04 0.04 0.04 0.04 0.04 0.04 0.04	Fe/Ca mmol/mol
Misra et al., 2014	This study	This study	Reference

Table S4

Specie	Seasonality
T. sacculifer (sacc)	Spring
T. sacculifer (w/o sacc)	Spring
G. ruber (white)	Summer
N. dutertrei	Winter
G. tumida	Annual average
G. menardii	Annual average
O. universa	Annual average
P. obliquiloculata	Winter

Table S5

Species O. universa O. universa O. universa O. universa T. sacculifer G. ruber All All	Species T. sacculifer (sacc) T. sacculifer (who sacc) G. ruber (white) N. dutertrei G. tumida G. menardii O. universa P. obliquiloculata
A 16.500 15.700 14.900 14.910 14.200 16.900 17.000	B 0.377 0.347 0.300 0.600 0.380 0.360 0.595 0.328
4.800 4.460 4.460 4.350 4.350 4.320 4.520 4.640	A 0.090 0.090 0.089 0.008 0.090 0.091 0.090 0.090
0.35 0.03 0.03 0.09	Reference Anand et al., 2003 Anand et al., 2003 Dekens et al., 2002 Dekens et al., 2002 Anand et al., 2002 Anand et al., 2003 Regenberg et al., 2009 Anand et al., 2003 Anand et al., 2003
dw correction -0.27 -0.27 -0.27 -0.27 -0.27 -0.27 -0.2 -0.2 -0.22 -0.27	
Condition LL ML HL	
Reference Bemis et al., 1998 Bemis et al., 2002 Bemis et al., 1998 Mulitza et al., 2003 Mulitza et al., 2003 Shackleton et al., 1974 Erez and Luz, 1983 Kim and O'Neil 1997	
Equation T=A-B* (\delta 180c-\delta 180w) T=A-B* (\delta 180c-\delta 180w)+C*(\delta 180c-\delta 180w)^2 T=A-B* (\delta 180c-\delta 180w) T=A-B* (\delta 180c-\delta 180w) T=A-B* (\delta 180c-\delta 180w) T=A-B* (\delta 180c-\delta 180w)+C*(\delta 180c-\delta 180w)^2 T=A-B* (\delta 180c-\delta 180w)+C*(\delta 180c-\delta 180w)^2 T=A-B* (\delta 180c-\delta 180w)+C*(\delta 180c-\delta 180w)^2 T=A-B* (\delta 180c-\delta 180w)+C*(\delta 180c-\delta 180w)^2	

Table S6

FC-01a G. ruber (white ss) 83 ± 20 30 ± 10 50 ± 20 Sime thesis 2006 FC-02a G. ruber (white ss) 56 ± 10 15 ± 10 60 ± 20 Sime thesis 2006 FC-12b G. ruber (white ss) Surface ± 10 -0.30 60 ± 10 Pectrs and Brumer, 2012 (non upwelling station Arabian sea) FC-13a G. ruber (white ss) Surface ± 10 20 ± 20 60 ± 10 Pectrs and Brumer, 2012 (non upwelling station Arabian sea) FC-13a G. ruber (white ss) Go ± 10 100±20 Elderfield and Ganssen, 2000 A14 G. ruber (white ss) 48 ± 10 50 ± 10 60 ± 10 Sime et al., 2005 FC-01a T. sacculifer (sacc) 7 ± 10 30 ± 10 80 ± 20 Elderfield and Ganssen, 2000 FC-02b T. sacculifer (sacc) 7 ± 10 30 ± 10 80 ± 20 Elderfield and Ganssen, 2000 FC-01a T. sacculifer (sacc) 7 ± 10 30 ± 10 80 ± 20 Sime tests 2006 FC-02a T. sacculifer (wo sacc) 32 ± 10 10 ± 10 80 ± 20 Sime tests, 2006 FC-01a T.	Core	Species	CD ₁	CD ₂	CD ₃	Reference
FC-12b G. ruber (white ss)	FC-01a	G. ruber (white ss)	83 ± 20	30 ± 10	50 ± 20	Sime thesis 2006
FC-13a G. ruber (white ss)	FC-02a	G. ruber (white ss)	56 ± 10	15 ± 10	60 ± 20	Sime thesis 2006
NP7-01 G. ruber (white ss)	FC-12b	G. ruber (white ss)	Surface ± 10	0-30	60 ± 10	Peeters and Brumer, 2012 (non upwelling station Arabian sea)
A14 G. ruber (white ss)	FC-13a	G. ruber (white ss)	Surface ± 10	20 ± 20	60 ± 10	Peeters and Brumer, 2012 (non upwelling station Arabian sea)
FC-01a	WP7-01	G. ruber (white ss)			100±20	Elderfield and Ganssen, 2000
FC-02a	A14	G. ruber (white ss)		60 ± 10	100±20	Elderfield and Ganssen, 2000
FC-12b	FC-01a	T. sacculifer (sacc)	48 ± 10	50 ± 10	60 ± 10	Sime et al., 2005
WP7-01 T. sacculifer (sacc) S0 ± 20 125 ± 15 Rickaby et al., 2005	FC-02a	T. sacculifer (sacc)	7 ± 10	30 ± 10	80 ± 20	Sime et al., 2006
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	FC-12b	T. sacculifer (sacc)	15 ± 10	40 ± 10		
FC-01a	WP7-01	T. sacculifer (sacc)		80 ± 20	125 ± 15	Rickaby et al., 2005
FC-02a	A14	T. sacculifer (sacc)		60 ± 10	125 ± 15	
FC-12b	FC-01a	T. sacculifer (w/o sacc)		50 ± 10		Sime thesis 2006 (Wind22-b)
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	FC-02a	T. sacculifer (w/o sacc)	32 ± 10	10 ± 10		Sime thesis 2006
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	FC-12b		$0-15 \pm 10$	30 ± 10	45 ± 20	Peeters and Brumer, 2012 (non upwelling station Arabian sea)
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	WP7-01	T. sacculifer (w/o sacc)		80 ± 20	125 ± 15	Rickaby et al., 2005
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	A14	T. sacculifer (w/o sacc)		60 ± 10	125 ± 15	Rickaby et al., 2005
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$						
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$						
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$					65 ± 10	Birshe et al., 2013
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$			35 ± 10			
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$						
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	A14	O. universa		55 ± 15		
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$					106 ± 20	Sime et al., 2005
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$		•				
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$		P. obliquiloculata				
FC-01a N. dutertrei 95 ± 20 90 ± 20 146 ± 20 Sime et al., 2005 FC-12b N. dutertrei 40 ± 10 50 ± 10 FC-13a N. dutertrei 45 ± 10 150 ± 20		4	65 ± 10			
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	WP7-01	P. obliquiloculata		125 ± 25		
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	FC-01a	N. dutertrei		90 ± 20		Sime et al., 2005
FC-13a N. dutertrei 45 ± 10 150 ± 20 Rickaby et al., 2005 MP7-01 N. dutertrei 125 ± 25 165 Rickaby et al., 2005 FC-01a G. menardii 135 ± 20 70 ± 20 FC-02a G. menardii 60 ± 10 60 ± 10 FC-12b G. menardii 65 ± 10 55 ± 15 60 ± 10 Peeters and Brumer, 2012 (non upwelling station Arabian sea) FC-13a G. menardii 55 ± 10 70 ± 10 60 ± 10 Peeters and Brumer, 2012 (non upwelling station Arabian sea) WP7-01 G. menardii 180 ± 20 Peeters and Brumer, 2012 (non upwelling station Arabian sea) FC-01a G. tumida 70 ± 20 100 ± 10 160 ± 20 Birshe et al., 2013 FC-02a G. tumida 70 ± 20 130 ± 20 160 ± 20 Birshe et al., 2013					146 ± 20	Sime et al., 2005
WP7-01 N. dutertrei 125 ± 25 165 Rickaby et al., 2005 FC-01a G. menardii 135 ± 20 70 ± 20 FC-02a G. menardii 60 ± 10 60 ± 10 FC-12b G. menardii 65 ± 10 55 ± 15 60 ± 10 Peeters and Brumer, 2012 (non upwelling station Arabian sea) FC-13a G. menardii 55 ± 10 70 ± 10 60 ± 10 Peeters and Brumer, 2012 (non upwelling station Arabian sea) WP7-01 G. menardii 180 ± 20 Peeters and Brumer, 2012 (non upwelling station Arabian sea) FC-01a G. tumida 70 ± 20 100 ± 10 160 ± 20 Birshe et al., 2013 FC-02a G. tumida 70 ± 20 130 ± 20 160 ± 20 Birshe et al., 2013						
A14 N. dutertrei 110 ± 20 165 Rickaby et al., 2005 FC-01a G. menardii 135 ± 20 70 ± 20 FC-02a G. menardii 60 ± 10 60 ± 10 FC-12b G. menardii 65 ± 10 55 ± 15 60 ± 10 Peeters and Brumer, 2012 (non upwelling station Arabian sea) FC-13a G. menardii 55 ± 10 70 ± 10 60 ± 10 Peeters and Brumer, 2012 (non upwelling station Arabian sea) WP7-01 G. menardii 180 ± 20 FC-01a G. tumida 70 ± 20 100 ± 10 160 ± 20 Birshe et al., 2013 FC-02a G. tumida 70 ± 20 130 ± 20 160 ± 20 Birshe et al., 2013			45 ± 10			
FC-01a						
FC-02a G. menardii 60 ± 10 60 ± 10 60 ± 10 Peeters and Brumer, 2012 (non upwelling station Arabian sea) FC-12b G. menardii 55 ± 10 70 ± 10 60 ± 10 Peeters and Brumer, 2012 (non upwelling station Arabian sea) FC-13a G. menardii 180 ± 20 Peeters and Brumer, 2012 (non upwelling station Arabian sea) FC-01a G. tumida 70 ± 20 100 ± 10 160 ± 20 Birshe et al., 2013 FC-02a G. tumida 70 ± 20 130 ± 20 160 ± 20 Birshe et al., 2013	A14	N. dutertrei		110 ± 20	165	Rickaby et al., 2005
FC-12b G. menardii 65 ± 10 55 ± 15 60 ± 10 Peeters and Brumer, 2012 (non upwelling station Arabian sea) FC-13a G. menardii 55 ± 10 70 ± 10 60 ± 10 Peeters and Brumer, 2012 (non upwelling station Arabian sea) WP7-01 G. menardii 180 ± 20 Birshe et al., 2013 FC-01a G. tumida 70 ± 20 130 ± 20 Birshe et al., 2013 FC-02a G. tumida 70 ± 20 130 ± 20 Birshe et al., 2013						
FC-13a						
WP7-01						
FC-01a G. tumida 70 ± 20 100 ± 10 160 ± 20 Birshe et al., 2013 FC-02a G. tumida 70 ± 20 130 ± 20 160 ± 20 Birshe et al., 2013			55 ± 10		60 ± 10	Peeters and Brumer, 2012 (non upwelling station Arabian sea)
FC-02a G. tumida 70 ± 20 130 ± 20 Birshe et al., 2013	WP7-01	G. menardii		180 ± 20		
						Birshe et al., 2013
WP7-01 <i>G. tumida</i> 180 ± 20 $210 - 240$ Rickaby et al., 2005	FC-02a		70 ± 20	130 ± 20	160 ± 20	Birshe et al., 2013
	WP7-01	G. tumida		180 ± 20	210 - 240	Rickaby et al., 2005

 $[\]text{CD}_1\text{:}$ Depth habitat estimated from $\delta^{18}\text{O}_c$

CD₂: Depth habitat estimated from Mg/Ca derived temperature

 $CD_{3:}$ Depth habitat from literature

 Pacific Ocean

 WP07-a
 G.

 WP07-a
 T.

 WP07-a
 T.

 WP07-a
 P.

 WP07-a
 P.

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 N.

 WP07-a
 C.

 Indian Ocean

 FC-01a
 G.

 FC-01a
 T.

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FC-12b
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FC-12b FC-13a FC-13a FC-13a FC-13a FC-02a FC-02a FC-02a FC-02a FC-02a FC-02a FC-02a Atlantic Ocean
CD107a O. universa G. ruber (white ss)
T. sacculifer (sacc)
T. sacculifer (woo sacc)
O. universa
P. obliquiloculata
G. menardii
N. dutertrei Sea G. ruber (white ss)
T. sacculifer (sacc)
T. sacculifer (wo sacc)
O. universa
P. obliquiloculata
N. duterirei
G. tumida G. ruber (white ss)
T. sacculifer (sacc)
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O. universa
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N. dutertrei
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T. sacculifer (wo sacculifer (wo sacculifer (wo sacculifer)
O. universa
P. obliquiloculata
G. menardii
N. dutertrei
G. tumida G. ruber (whn T. sacculifer P. obliquiloci G. menardii N. dutertrei G. ruber (white ss) T. sacculifer (w/o sacc) P. obliquiloculata Depth habitat (m) Temperature 60 ± 20 80 ± 20 80 ± 20 80 ± 20 65 ± 10 60 ± 10 60 ± 10 146 ± 20 130 ± 20 50 ± 20 60 ± 10 60 ± 10 90 ± 20 106 ± 20 70 ± 20 95 ± 20 100 ± 20 $100\pm20 \\ 80\pm20 \\ 125\pm20 \\ 75\pm25 \\ 125\pm25 \\ 165\pm20 \\ 180\pm20 \\ 180\pm20 \\$ 20 ± 20 45 ± 20 50 ± 10 55 ± 10 150 ± 20 0-30 40 ± 10 15 ± 10 35 ± 10 50 ± 10 65 ± 10 40 ± 10 80 ± 20 PRE INDUSTRIAL IN-SITU PARAMETERS 26.1 27.4 24.7 27.7 27.7 24.7 20.6 18.7 27.1 25.7 25.4 25.4 25.1 19.4 23.4 21.9 21.9 18.2 20.5 20.5 17.7 17.1 25.9 25.3 25.9 25.9 25.5 25.0 24.1 25.3 20.4 19.3 19.3 20.1 18.3 20.4 17.5 12.0 2sd*0.3 2.4 2.2 1.8 2.6 3.6 3.8 0.6 0.6 0.0 0.9 1.1 0.6 1.2 1.3 5.7 5.7 5.7 4.0 4.6 4.6 4.6 4.6 4.6 1.9 1.9 1.9 1.9 1.9 1.9 1.9 Salinity 35.2 35.0 35.4 34.9 35.4 35.5 36.7 36.6 36.6 36.6 36.5 36.5 36.5 36.5 36.5 35.5 35.6 35.6 35.6 35.6 35.6 35.0 35.1 35.1 35.2 35.2 35.2 35.2 35.6 0.01 2sd* pH (pre-ind) 0.5 $\begin{array}{cccc} 0.1 & 0.$ 8.05 8.08 8.03 8.09 8.03 8.03 8.10 8.07 8.06 8.04 7.80 8.18 8.10 8.16 8.11 8.06 8.06 8.02 8.19 2sd*0.03 0.05 0.05 0.00 0.07 0.07 0.01 0.03 0.00 pCO_2 387 357 413 349 413 408 274 271 350 290 334 332 438 350 350 343 343 343 343 343 343 2sd*21 51 56 30 68 30 10 39 64 34 56 69 72 72 72 34 45 HCO₃ 1753 1708 1797 1797 1696 1797 1855 1724 1782 1797 1815 2056 1655 1756 1681 1738 1738 1794 1724 1741 1741 1728 1728 1738 1743 1773 1773 1856 1862 1802 1802 1868 1856 2sd*81 89 31 110 80 80 58 53 75 45 68 75 75 75 75 75 75 75 75 75 75 75 75 26 38 38 15 17 17 19 31 1138 1126 1126 1126 94 99 99 29 CO_3^2 292 249 281 281 257 257 209 209 249 263 233 233 233 233 233 246 240 240 245 245 228 228 216 216 224 242 209 209 247 247 209 189 189 226 215 215 218 184 204 204 180 175 192 2sd*23 31 19 28 32 32 31 31 17 16 29 29 10 15 15 6 6 7 7 7 7 13 31 33 9 39 26 18 24 35 36 36 37 36 12 2059 DIC 1996 2032 2041 2052 2199 1954 2015 1970 2004 2008 2072 2015 1979 1989 1989 1981 1981 2007 2024 2015 1979 1998 1998 2053 2018 2018 2061 2070 1988 1960 2017 2017 1953 2017 2057 2sd* 93 62 65 65 67 18 52 58 23 26 35 23 4 4 23 25 17 24 24 9 9 10 12 18 ALK 2371 2371 2370 2369 2348 2374 2369 2374 2369 2368 2365 2365 2332 2333 2333 2333 2332 2334 2332 2333 2333 2299 2303 2303 2311 2316 2306 2306 2313 2313 2333 2305 2303 2311 2311 2303 2311 2322 2322 2sd* 7 10 10 15 18 0 0 22 18 18 11 11 3 _ · δ11Bborate 17.86 18.21 18.68 17.80 18.78 17.82 17.34 17.23 19.81 18.71 19.54 18.90 18.24 17.69 18.75 19.06 118.89 118.90 119.00 118.64 118.64 119.05 118.33 19.00 18.86 18.31 18.04 15.37 18.40 18.15 18.16 17.42 17.91 17.93 17.26 0.31 2sd* 0.64 0.65 0.23 0.83 0.48 0.450.65 0.77 0.50 0.69 0.67 0.62 0.71 0.37 0.45 0.56 0.34 0.30 0.30 0.17 0.14 0.30 0.27 0.90 0.81 0.82 0.61 0.66 0.68 0.68 T_{Mg/Ca} 2sd pH 811B 2sd** pCO₂ 2sd*** CALCULATED PARAMETERS 27.0 27.3 28.4 28.2 24.6 26.4 21.2 27.5 30.6 28.9 18.8 19.5 30.2 26.9 29.3 27.2 26.6 24.8 24.6 24.0 25.1 25.1 22.2 21.7 19.9 20.3 17.9 20.3 20.2 18.5 17.2 26.1 24.5 25.3 20.0 1111111 1.4 1.4 1.4 1 1 4 1 4 1 1 1 1 1 1 1 1 4 4 4 4 4 4 4 14 4 4 8.19 8.11 8.18 8.16 8.17 8.12 8.13 8.03 7.99 8.03 8.10 8.03 8.09 8.08 7.98 7.79 8.15 8.09 8.13 8.13 8.05 7.97 8.06 8.10 8.10 7.98 8.10 8.10 8.09 0.06 0.04 0.05 0.04 0.05 0.04 0.05 0.040.08 0.04 0.08 0.04 0.04 0.03 0.03 0.08 0.06 0.04 0.03 0.04 0.08 0.08 0.05 0.05 0.04 0.04 0.06 0.08 0.08 0.05 0.05 285 305 374 332 336 415 514 514 514 514 350 350 351 426 478 427 427 352 430 430 406 387 369 357 369 373 373 400 332 286 286 287 332 287 332 293 28 54 50 50 50 58 68 52 29 29 55 68 252324552 22 4445 39 Calibration [11] [13] [13] 8 pH 611B 2sd** pCO2 2sd*** Calibration 8.05 8.20 8.10 8.16 8.03 8.21 7.80 8.13 7.92 7.88 8.04 8.10 0.09 0.04 0.10 0.06 0.06 $0.10 \\ 0.05$ 0.16 0.06 0.05 417 630 417 356 778 324 573 265 366 302 428 267 52 49 73 32 26 53 2 2 484 777 7 777 77 777

 ^{*} uncertainties calculated using Henehan's 2016 R code
 ** propagated uncertainty on pH including d11B, temperature, salinity uncertainties
 *** propagated uncertainty on pCO2 including d11B, temperature, salinity, Alk uncertainties