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Key Points:

- B and C isotope analyses of the reef-building coral *S. siderea* reveal reef-scale patterns of pH, NEC, and NEP
- Declining coral extension is not associated with ocean acidification
- NEC was enhanced in the backreef in the 1920s–1940s, NEP was enhanced in the forereef since the 1980s

Supporting Information:

- Supporting Information S1
- Data Set S1

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Historical Trends in pH and Carbonate Biogeochemistry on the Belize Mesoamerican Barrier Reef System

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Abstract Coral reefs are important ecosystems that are increasingly negatively impacted by human activities. Understanding which anthropogenic stressors play the most significant role in their decline is vital for the accurate prediction of future trends in coral reef health and for effective mitigation of these threats. Here we present annually resolved boron and carbon isotope measurements of two cores capturing the past 90 years of growth of the tropical reef-building coral *Siderastrea siderea* from the Belize Mesoamerican Barrier Reef System. The pairing of these two isotope systems allows us to parse the reconstructed pH change into relative changes in net ecosystem productivity and net ecosystem calcification between the two locations. This approach reveals that the relationship between seawater pH and coral calcification, at both a colony and ecosystem level, is complex and cannot simply be modeled as linear or even positive. This study also underscores both the utility of coupled $\delta^{11}B-\delta^{13}C$ measurements in tracing past biogeochemical cycling in coral reefs and the complexity of this cycling relative to the open ocean.

Plain Language Summary Coral reefs worldwide are in decline due to multiple anthropogenic stressors (e.g., ocean warming and acidification in response to anthropogenic CO_2 release). Here we explore a new way to examine the relationship between these anthropogenic stressors and critical ecosystem-scale processes using paired annual-resolution boron and carbon isotope measurements of coral cores spanning the past century. This isotopic pairing allowed us to compare changes in seawater pH, net ecosystem productivity, and net ecosystem calcification between forereef and backreef locations of the southern Belize Mesoamerican Barrier Reef System. We specifically show that the forereef colony, which exhibits declining growth over the past century, recorded unexpectedly steady pH over the same interval. The backreef colony, in contrast, exhibited accelerating growth over this interval, while recording surprisingly rapid ocean acidification. This apparent contradiction is explained with carbon isotope data from the cores showing that increasing primary production on the forereef has offset anthropogenic acidification. Likewise, the increasing rate of coral calcification in the backreef (which generates acidity) has exacerbated anthropogenic acidification of backreef waters. Thus, although ocean acidification and warming influence calcification and primary production within reef systems, the resulting changes can drive biogeochemical feedbacks capable of eliminating or amplifying the anthropogenic CO_2 signal.

1. Introduction

1.1. Ocean Acidification and Coral Reefs

Ocean acidification (OA) in response to anthropogenic emissions of CO₂ has been shown to negatively impact marine calcifiers (e.g., Gattuso et al., 2015). Coral reefs account for around 50% of shallow water CaCO₃ production (~0.7 Pg of CaCO₃ per year), protect shorelines from storms and rising sea level, generate trillions of dollars in tourism worldwide, and the ecosystems that they support feed millions of people (Costanza et al., 2014). However, the effect of OA on CaCO₃ production and dissolution on coral reefs remains unclear. This is despite a conceptually simple positive relationship between ocean pH ($-\log_{10}$ [H+]) and the saturation state (Ω) of aragonitic CaCO₃, defined as

$$\Omega_{\text{aragonite}} = \frac{\left[\mathsf{Ca}^{2+}\right] \times \left[\mathsf{CO}_{3}^{2-}\right]}{K_{\text{aragonite}}^{*}}$$
(1a)

$$CO_{2(aq)} + H_2O \leftrightarrow HCO_3^- + H^+ \leftrightarrow CO_3^{2-} + 2H^+$$
(1b)

where $[Ca^{2+}]$ and $[CO_3^{2-}]$ are the calcium and carbonate ion concentration in seawater, respectively, and $K^*_{aragonite}$ is the solubility product of aragonite in seawater. Part of the uncertainty in the future response of coral reef calcification to OA is due to the high degree of natural variability in the ocean carbonate system that occurs in coral reefs (DeCarlo et al., 2017). In particular, it is recognized that the balance between net ecosystem production (NEP = gross primary production – autotrophic and heterotrophic respiration) and net ecosystem calcification (NEC = gross calcification – gross CaCO₃ dissolution) modulates the carbon chemistry of these complex ecosystems. CO₂ is both the product of calcification and the principal reactant in photosynthesis; therefore, these two processes can in principal be linked and feedbacks may exist such that anthropogenic or natural changes in CO₂ could alleviate or amplify the influence of OA on the inhabitants of a reef (e.g., Yeakel et al., 2015).

The balance between NEP and NEC, and its evolution through time, can be investigated through measurements of total alkalinity (ALK) and total dissolved inorganic carbon (DIC) in reef waters (e.g., Bates et al., 2010). Unfortunately, such monitoring of the "health" of coral reefs has only recently begun and instrumental records longer than ~5 years are rare. Those longer-term records that do exist tend to be characterized by considerable gaps in sampling (e.g., Silverman et al., 2012, 2014), with the longest continual record thus far restricted to the last 20 years (Bates, 2017). As a consequence, the full impact of the acidification that has occurred over the last 100 years (~ 0.1 pH units in the open ocean; Bates et al., 2014), and the ability (or lack thereof) of reef systems to modulate their carbonate biogeochemistry against this apparent threat, remains largely unknown.

1.2. Proxy Constraints on NEP and NEC Variability

Reliable observations of the carbonate chemistry of coral reefs prior to the 1980s do not exist. It is therefore unlikely that the ecosystem level response of coral reefs to historic anthropogenic oceanic change can be estimated from direct measurements. Instead, we must turn to geochemical proxies. One such proxy that has received much attention is the boron isotope pH proxy (e.g., Goodkin et al., 2015; Liu et al., 2014; Pelejero et al., 2005; Wei et al., 2009). There are two aqueous species of boron in seawater (boric acid and borate ion), and their relative abundance depends primarily on pH (Dickson, 1990). Boron has two stable isotopes, ¹¹B and ¹⁰B, and a strong isotopic fractionation exists between boric acid and borate ion (~27.2‰; Klochko et al., 2006). This causes the isotopic composition of the two aqueous species of boron to also vary with pH as their relative abundance changes (see Foster & Rae, 2016 for more details). The following relationship can thus be defined:

$$pH = pK_B^* - \log\left(-\frac{\delta^{11}B_{(sw)} - \delta^{11}B_{bor}}{\delta^{11}B_{(sw)} - \alpha_B\delta^{11}B_{bor} - \varepsilon_B}\right)$$
(2)

where $pK_B^* \approx 8.6$ is the $-\log_{10}$ of the stoichiometric equilibrium constant for boron dissociation in seawater ($pK_B^* \approx 8.6$ in seawater at 25°C, 35 psu salinity and atmospheric pressure; Dickson, 1990), $\delta^{11}B_{sw}$ is the boron isotopic composition of seawater (39.61‰; Foster et al., 2010), $\delta^{11}B_{bor}$ is the boron isotopic composition of seawater borate ion, and α_B and ε_B are expressions of the magnitude of isotopic fractionation between the two aqueous species of boron (1.0272 and 27.2, respectively; Klochko et al., 2006).

Recent experiments have shown that only the borate ion is incorporated into inorganic aragonite (Noireaux et al., 2015) and that this process occurs with little or no isotope fractionation, allowing substitution of aragonite $\delta^{11}B$ for $\delta^{11}B_{bor}$ in equation (2) and the calculation of the pH at which the aragonite precipitated. A complicating factor of using boron isotopes in coral aragonite, however, is the influence of coral physiology and the elevation of pH within the calcifying fluid (pH_{cf}) by the coral (Venn et al., 2013), such that $\delta^{11}B$ of the coral skeleton ($\delta^{11}B_{coral}$) represents $\delta^{11}B_{bor}$ of the calcifying fluid rather than $\delta^{11}B_{bor}$ of seawater (Holcomb et al., 2014; McCulloch et al., 2012). Nevertheless, because changes in seawater pH have been shown to cause changes in coral calcifying fluid pH (e.g., Ries, 2011), coral skeletal $\delta^{11}B$ tracks seawater $\delta^{11}B_{bor}$ and there is a consistent relationship between internal and external pH (e.g., Holcomb et al. (2014)). This permits the reconstruction of seawater pH at annual resolution from the boron isotope composition of coral skeletons using species specific $\delta^{11}B$ -pH calibrations (e.g., Liu et al., 2014; Trotter et al., 2011).

Although surface ocean pH has declined by ~0.1 pH units over the last 100 years, pH on a coral reef is also intimately related to local physical and biogeochemical processes that change the relative concentrations



Figure 1. Plot of total alkalinity (ALK) versus total dissolved inorganic carbon (DIC) contoured with pH (solid) and $\Omega_{aragonite}$ (dashed) and calculated at 25°C, 35 psu, and atmospheric pressure. The red lines show the influence of NEP and NEC on ALK and DIC. The central red dot represents an arbitrary starting point with the arrows showing the direction of change for increase/ decrease (+/–) in NEC and NEP. One mole of CaCO₃ production on the reef removes 2 moles of ALK and 1 mole of DIC, whereas dissolution of CaCO₃ does the reverse. Reef primary production removes 1 mol of DIC as CO₂ and respiration does the opposite, with minimal effect on ALK (Zeebe & Wolf-Gladow, 2001). The NEP arrow has ticks that show the change in δ^{13} C of DIC caused by the removal/addition of organic carbon with a δ^{13} C of -24 ‰.

of ALK and DIC, the residence time of seawater on the reef, and water depth (e.g., Bates et al., 2010; Figure 1). Indeed, water column pH on a coral reef can vary up to ±0.5 pH units on daily, monthly, and seasonal time scales (e.g., DeCarlo et al., 2017; Drupp et al., 2013). Although CO2 outgassing (and, conversely, CO₂ invasion) and benthic processes can impact seawater pH on a reef, the main factors controlling this pH variability are temporal variations in NEC and NEP, which themselves reflect the addition/removal of DIC and ALK by reef biology, and also water residence time and water depth (Bates et al., 2010; DeCarlo et al., 2017; Gattuso et al., 1999). This strong biological control of coral reef pH (Figure 1) complicates the reconstruction of long-term open-ocean pH trends from the boron isotopic composition of long-lived tropical corals and likely accounts for the annual and decadal variability in existing δ^{11} B-derived seawater pH records (Figure S1 in the supporting information). Although the complex carbonate chemistry of coral reef systems complicates paleo-pH reconstruction from coral δ^{11} B, it affords an opportunity to reconstruct past changes in NEC and NEP within these critical ecosystems.

As shown in Figure 1, seawater pH can be affected by modifying NEC and/or NEP and, as noted by Yeakel et al. (2015), pH on its own cannot distinguish the process(es) responsible for any observed change in seawater carbonate chemistry. We assert that the carbon isotopic composition (δ^{13} C) of the coral skeleton can provide critical information in this regard, allowing a more nuanced view of the temporal evolution of the seawater carbonate system, because, unlike pH, the δ^{13} C of seawater DIC is only significantly altered by variations in NEP and is unaffected by NEC (Figure 1). Although it is unclear to what extent coral

skeleton δ^{13} C reflects δ^{13} C of seawater DIC, which may also be influenced by a number of "vital effects" (Allison et al., 1996; Gagan et al., 1994; Krief et al., 2010; Martin et al., 2016; McConnaughy, 1989; Meibom et al., 2006; Reynaud et al., 2004; Swart, 1983), Swart et al. (2010) showed that 78% of 37 colonies of several species of tropical corals exhibited a decline in δ^{13} C over the past century, consistent with their δ^{13} C being driven predominantly by the invasion of isotopically light anthropogenic carbon into the atmosphere and oceans—the so-called Suess effect (Keeling et al., 1979). That most tropical corals exhibit a steady decline in δ^{13} C over the past century implies an overriding sensitivity of coral δ^{13} C to the changing δ^{13} C of seawater DIC—regardless of whether that variability is driven by the Suess effect or changing NEP (Figure 1).

Here we use paired measurements of δ^{13} C and δ^{11} B of the skeleton of the scleractinian coral *Siderastrea* siderea sampled at an annual resolution to provide an understanding of how reef pH and patterns of NEP and NEC varied across the Sapodilla Cayes region of the Mesoamerican Barrier Reef System (MBRS) in Belize over the last 90 years. These results provide key insights into the impact of OA on *S. siderea* and demonstrate the utility of these two proxies in tracing the "health" of coral reef systems beyond the reach of the instrumental record.

2. Materials and Methods

Two cores were obtained in December 2009 from forereef (FR-02; 16.13715°N, 88.252883°W) and backreef (BR-06; 16.14045°N, 88.26015°W) colonies of *S. siderea* growing at 3 to 5 m water depth within Sapodilla Cayes region of the MBRS (Castillo et al., 2011; supporting information Figure S2). Consistent with global trends, coral cover on the MBRS is in decline (Gardner et al., 2003), although the exact cause(s) is (are) currently widely debated (Aronson et al., 2002; Hughes, 1994; McClanahan et al., 1999). Methods for subsampling of coral cores to separate annual growth bands, X-raying, and age dating are described in Castillo et al. (2011). Further sampling protocols and analytical methodology for δ^{11} B and δ^{13} C are described further in the supporting information Text S1.



Figure 2. Geochemical data from forereef (FR-02; red) and backreef (BR-06, blue) cores of the scleractinian coral S. siderea from the southern MBRS, Belize, spanning 1910 to 2010. (a) Boron isotope (δ^{11} B) data with error bars representing analytical uncertainty. (b) The carbon isotope (δ^{13} C) composition of atmospheric CO₂ from Rubino et al. (2013) (gray line, left axis) and annually resolved $\delta^{13}\bar{C}$ of forereef (FR-02; red) and backreef (BR-06; blue) coral cores (right axis). Thick lines for the coral data are three-point running means. (c) Seawater pH derived from the coral δ^{11} B data plotted in panel (a) (see text for details). Gray line is seawater pH calculated for the past century from atmospheric pCO₂, the HadISST seawater temperature record, an assumed total of alkalinity of 2,300 $\mu \text{mol/kg},$ and an assumed salinity of 35 psu. Also shown are LOESS best fit lines and 95% confidence intervals, with degree of smoothing determined by generalized cross validation. (d) Linear extension of forereef (FR-02; red) and backreef (BR-06; blue) corals as reported by Castillo et al. (2011), with thick lines showing the three-point running mean.

The $\delta^{11}B$ compositions of the forereef and backreef corals were used to calculate pH_{cf} using a modified version of equation (2):

$$pH_{cf} = pK_B^* - \log\left(-\frac{\delta^{11}B_{(sw)} - \delta^{11}B_{coral}}{\delta^{11}B_{(sw)} - \alpha_B\delta^{11}B_{coral} - \varepsilon_B}\right)$$
(3)

 pK_{B}^{*} was determined for each sample based on the sea surface temperature from the HadISST1 (Hadley Centre Sea Ice and Sea Surface Temperature) data set (Rayner et al., 2003). Salinity has a very minor effect on calculated pH and so was assumed to be constant at 35 psu. All carbonate system calculations were made using the R package "seacarb" (Lavigne & Gattuso, 2010).

There is currently no δ^{11} B-pH calibration for *S. siderea*. Instead, we use equations developed for the coral species *Porites cylindrica* (Liu et al., 2014; McCulloch et al., 2012; Trotter et al., 2011) to estimate seawater pH (pH_{sw}) from the pH_{cf}:

$$pH_{\rm sw} = \frac{(pH_{\rm cf} - 4.72)}{0.466} \tag{4}$$

This choice of calibration determines the absolute pH but has little influence on relative pH change, which is the focus of the present work (see below and supporting information Figure S3; Foster & Rae, 2016). In order to isolate the long-term trends in the 1912–2008 time series, reconstructed pH_{sw} values were fitted with LOESS smoothers in R, with the degree of smoothing determined via generalized cross validation.

3. Results and Discussion

3.1. The $\delta^{11}\text{B-Derived}$ Seawater pH and $\delta^{13}\text{C}$ Time Series From the MBRS

Both the forereef (FR-02) and backreef (BR-06) core exhibit a decline in δ^{13} C of 1 to 1.5‰ over the last ~100 years, albeit with an offset between them of around 0.4‰ that increased since the 1980s (Figure 2). This broad temporal evolution is in good agreement with other coral studies (e.g., Böhm et al., 2002; Swart et al., 2010) and is similar to trends observed in the atmosphere (Rubino et al., 2013) over a comparable interval. Both Belize coral δ^{13} C records, however, display considerable inter-annual variability (Figure 2) with an apparent ~8-year cyclicity (supporting information Figure S4). This is consistent with the assertion that the long-term trend of

the δ^{13} C time series recorded in these two corals reflects the uptake of isotopically light anthropogenic CO₂, with short-term variability reflecting either local variability in DIC δ^{13} C or non-DIC influences on coral δ^{13} C (Figure 2).

The δ^{11} B time series for the two coral cores are surprisingly distinct (Figure 2), despite their geographic proximity (Figure S2). Specifically, the forereef coral (FR-02) exhibits no consistent long-term trend (p = 0.41), but exhibits the same quasi-decadal variability evident in the δ^{13} C records (Figure S4). In contrast, the backreef coral (BR-06) exhibits a consistent decline in δ^{11} B over the same interval (~ -1.2‰ per 100 years; Figure 2), while exhibiting less short-term variability than observed for FR-02 (Figure S4). Given the overriding influence of external pH on the δ^{11} B of coral aragonite [equations (2)–(4)], these disparate δ^{11} B records imply vastly different trends in seawater pH at the two core locations.

The δ^{11} B record within each core was converted to a pH time series using equations (3) and (4) (Figure 2). Although the lack of a species-specific δ^{11} B-pH calibration for *S. siderea* means that absolute values of the reconstructed seawater pH should be interpreted with caution, the observed trends in seawater pH can be interpreted with confidence (Figure S3). This analysis suggests that seawater pH at the site of the backreef



Figure 3. (a) The relative evolution of δ^{11} B-derived seawater pH and δ^{13} C in the forereef and backreef of the southern MBRS, Belize. The time series from the backreef and forereef cores plotted in Figure 2 were interpolated onto a common age scale, permitting calculation of the forereef-to-backreef difference in seawater pH and δ^{13} C (Δ pH and Δ^{13} C, respectively). Each datum is color coded for year, as shown on scale to the right. Error bars are quadratic sum of the uncertainty relating to the δ^{11} B measurement (±0.02 pH units) and an assumed but conservative uncertainty in the pH- δ^{11} B calibration (±0.04 pH units). The plot is contoured by assuming a starting condition of DIC = 2000 µmol/kg and pH = 8.16, with the removal or addition of DIC via changing NEP shown by black lines and the removal or addition of ALK:DIC at a 2:1 ratio via changing NEC shown by gray lines. Contours are labeled with µmol/kg change in either DIC or ALK. It is assumed that NEC has no effect on δ^{13} C of seawater DIC and that organic carbon has δ^{13} C = -24 % (Briand et al., 2015). Thick red line is best fit least squares regression (with associated 95% confidence interval) through Δ pH and Δ^{13} C data. (b) Estimated pH change caused by changing NEC only. Δ pH > 0 indicates higher NEP in the forereef and vice versa. Error bars in (b) and (c) are approximately ±0.05 pH units at 95% confidence as in (a).

coral decreased over the last ~100 years from 8.05 to 7.87, with an average decline of -0.016 pH units per decade (determined by linear regression). This rate of acidification accelerated to -0.019 ± 0.006 units per decade (p = 0.009) after 1970—similar to that of the open North Atlantic as measured by the Bermuda Atlantic Time Series for the interval 1983–2014 (-0.017 ± 0.001 per decade; Bates et al., 2014) and the nearby Cariaco Basin for the interval 1995–2014 (-0.025 ± 0.004 per decade; Table S1). Although the more recent portion of the backreef pH record matches seawater pH trends expected from records of atmospheric pCO_2 , the interval of marked acidification from 1920 to 1940 in the backreef core is much higher than expected given the atmospheric pCO_2 record (Figure 2). In contrast, the forereef core exhibits no significant long-term pH trend over the same interval (-0.004 ± 0.004 per decade, p = 0.36) or since 1970 (-0.002 ± 0.02 per decade; p = 0.946).

3.2. Relationship Between Linear Extension and Seawater pH for S. Siderea

Castillo et al. (2011) found that over the last 100 years, there was a marked regional-scale decline in linear extension for forereef *S. siderea* of the southern MBRS, averaging around -0.2 cm/year, while extension in

backreef corals increased by 0.1 cm/year. Given the close relationship between rates of linear extension and rates of calcification that exist in this species (Carricart-Ganivet et al., 2013), this provides the opportunity to assess how calcification of *S. siderea* has responded to the OA that it has experienced over the last 100 years (Figure S5). Contrary to expectation (e.g., McCulloch et al., 2012), we find no statistically significant relationship (p = 0.5) between extension rates and reconstructed seawater pH (Figure S5) for the forereef coral and a strong *inverse* correlation (p = <0.001) for the backreef coral (i.e., increasing extension with decreasing pH; Figure S5).

The contrasting relationship between seawater pH and extension rate for the backreef and forereef corals (Figures 2d and S5) suggests that OA has not had an obviously detrimental effect on the calcification of *S. siderea* over the last 100 years in the Sapodilla Cayes. This finding agrees with the recent culturing experiment of Castillo et al., (2014) that showed that for moderate rates of acidification, calcification in this species is enhanced, possibly due to increased availability of dissolved CO₂ for symbiont photosynthesis. Although this suggests that *S. siderea* on the MBRS is apparently resilient to moderate OA, the laboratory experiments also revealed that there is a limit to the degree to which increased CO₂ availability for photosynthesis (i.e., decreased pH) offsets the deleterious effects of CO₂-induced reductions in $\Omega_{aragonite}$ on calcification in *S. siderea*. Nonetheless, other factors need to be investigated to determine what is responsible for the decline in forereef (but not backreef) *S. siderea* extension reported by Castillo et al. (2011).

3.3. Spatial Patterns of pH and $\delta^{13}C$ on the MBRS

In order to compare the evolution of δ^{13} C and pH displayed by the two coral cores, we interpolate both cores to a common age scale and plot forereef-to-backreef pH difference (Δ pH) versus forereef-to-backreef δ^{13} C difference (Δ^{13} C; Figure 3). A linear regression fitted to Δ^{13} C and Δ pH is described by the following equation:

$$\Delta p \mathsf{H} = (0.16 \pm 0.03 \cdot \Delta^{13} C) + 0.08 \pm 0.17 \qquad (R^2 = 0.38, p = 0.00007) \tag{5}$$

This strong correlation between δ^{11} B-derived pH and δ^{13} C, as well as the apparent ~8-year cyclicity in both proxies (Figure S4), is consistent with trends previously reported for single coral colonies (e.g., Hemming et al., 1998). Examining the difference between the two cores in this way removes the inherent covariation in coral δ^{13} C and δ^{11} B driven by increasing anthropogenic CO₂: anthropogenic-CO₂-induced reductions in seawater DIC δ^{13} C (Swart et al., 2010) via the Suess effect and reductions in seawater borate δ^{11} B via ocean acidification (Sabine et al., 2004). The residual correlation between pH and δ^{13} C between the cores (i.e., the correlation between Δ^{13} C and Δ pH) should therefore represent relative variations in NEP and NEC at the two core locations (e.g., Figure 1), thereby allowing reconstruction of trends in NEP and NEC at those locations through time. For simplicity, we ignore the possible influence of seawater pH on coral δ^{13} C that has been observed in some studies (e.g., Krief et al., 2010; Martin et al., 2016; Spero et al., 1997) and assume that changes in coral δ^{13} C solely reflect changes in δ^{13} C of seawater DIC—which is reasonable given the relatively small change in pH experienced by the MBRS over the last 100 years (<0.2 pH units) and the relatively small impact that seawater pH and other non-DIC factors have on coral δ^{13} C (e.g., Martin et al., 2016).

The observed positive relationship between Δ^{13} C and Δ pH is likely indicative of enhanced NEP and reduced NEC at the forereef core location relative to the backreef core location. More precise insights into the causes of such changes can, however, be identified provided the δ^{13} C of primary production and respiration are known (vertical contours on Figure 3). Values for biological tissue in coral reef ecosystems range from -17 to -31% (Briand et al., 2015; Carvalho et al., 2015; Muscatine et al., 1989; Reynaud et al., 2004; Titlyanov et al., 2010) and here we use a value of -24%, the median value from Briand et al. (2015), allowing us to contour Figure 3 for the removal/addition of DIC alone (i.e., change in NEP; black lines), as well as for the removal/addition of ALK:DIC with a 2:1 ratio (i.e., change in NEC; gray lines).

Projecting Δ pH and Δ^{13} C data onto the zero contour of constant alkalinity effectively parses the pH change into that caused by NEC (only changes pH; Figure 3b) and that caused by NEP (changes pH and δ^{13} C of DIC; Figure 3c). This treatment reveals two important trends: (i) NEC was enhanced in the backreef, relative to the forereef, during the 1920s–1940s and (ii) NEP was enhanced in the forereef, relative to the backreef, since the 1980s. These findings are largely irrespective of the specific δ^{11} B-pH calibration used and the δ^{13} C composition assigned to organic matter in the system. This deconvolution of the pH signals into NEC and NEP variability also indicates that much of the quasi-decadal variability in pH at the forereef core is driven by differences in NEC between core locations—specifically, by ~8-year cycles in NEC in the forereef that cause corresponding cycles in forereef seawater pH (Figure S4 and Table S2). Cross-correlation analysis of the detrended time series using the Pearson product moment correlation reveals a significant positive correlation (cor = 0.44, p < 0.001; Table S2) between ΔpH from NEP and ΔpH from NEC, such that when NEC decreases in the forereef, NEP in that region also tends to increase. A similar pattern is revealed in long-term trends, albeit with a temporal offset: NEC was enhanced in the backreef compared to the forereef during the 1920s–1940s and NEP was enhanced in the forereef relative to the backreef since the 1980s.

It is important to recognize that these parameters can be influenced by both biological and physical mechanisms. For instance, the apparent increase in NEC in the backreef relative to the forereef could be caused by either an enhancement of calcification in the backreef (relative to the forereef) or an increase in the residence time of the water on the reef so the impact of calcification is proportionally enhanced (e.g., DeCarlo et al., 2017). It is therefore important to consider both biological and physical oceanographic processes when inferring changes in NEC and NEP from δ^{11} B and δ^{13} C data.

Castillo et al. (2011) observed that rates of linear extension in *S. siderea* decreased in the forereef and increased in the backreef of the southern MBRS since the early 1900s. These trends in linear extension are therefore consistent with the observed reduction in forereef NEC and enhancement in backreef NEC through time being primarily driven by biological processes. It is also likely that the enhanced NEP in the forereef since the 1980s contributes to negating the open ocean OA signal evident in the backreef (BR-06) pH record. Similarly, the larger-than-expected pH change recorded in the backreef (BR-06) pH record, given historical changes in atmospheric pCO_2 (Figure 2), appears to be largely a consequence of enhanced NEC in the backreef—as observed at other backreef localities on shorter time scales (e.g., Bates, 2017).

A quasi-decadal (~8 year) climate cycle in the Caribbean has been noted previously (Jury, 2009) in records of SST and rainfall. This mode of variability is considered distinct from the Pacific El Niño–Southern Oscillation (ENSO) and instead is related to slow oscillations of the North Atlantic Oscillation (NAO) driven by interactions between tropical Atlantic SST, the subtropical anticyclones, and the Hadley circulation (Jury, 2009). For this mode of variability, warm SST in the Caribbean is typically equated with high rainfall. The HadlSST record used here also shows a quasi-decadal ~8-year cycle when detrended (Figures S4 and S6 and Table S2) and is significantly positively correlated with detrended NEC (cor = 0.37, p < 0.001). These trends are consistent with Castillo et al.'s (2011) observation that linear extension of backreef *S. siderea* has been impaired by this warming—a dichotomy that they attribute to the increased thermal tolerance of the backreef populations that have historically experienced more stable temperature fluctuations relative to forereef populations that have historically experienced more stable temperatures.

Despite the strong correlation between NEP and NEC, the correlation between detrended NEP and detrended HadlSST is not significant (cor = 0.07, p = 0.56; Table S2)—probably a result of the relatively small variability in NEP at this 8-year cycle and the low sampling resolution (Figure S6). However, area-averaged annual precipitation rates over the western Caribbean are significantly correlated with NEP (cor = 0.30, p = 0.003) but, unlike SST, are not significantly correlated with NEC (cor = 0.15, p = 0.16; Figures S2 and S6 and Table S2). However, limiting the area-averaged annual precipitation to land-only precipitation reveals the opposite pattern: a significant correlation with NEC (cor = 0.38, p < 0.001) but not with NEP (cor = 0.14, p = 0.19; Table S2).

These correlations suggest that SST and rainfall either influence NEC and NEP on the MBRS directly at the ecosystem level or else indirectly by changing water residence time and/or water depth on the MBRS. Although we cannot identify the relative contributions of SST and rainfall to NEC and NEP with the data at hand, long-term anthropogenic processes may be forcing the reconstructed trends. For instance, the Caribbean experienced significant warming from 1910 to 1940, coincident with the first long-term increase in NEC preserved in the cores (Figures 3b and S7). Jury (2009) also noted a general increase in mean annual rainfall in the Caribbean since the 1960s and Stephenson et al. (2014) recorded a positive trend for the wider Caribbean in the interval of 1986–2010 for total annual precipitation, for average daily precipitation, and for heavy rainfall events. This secular increase in precipitation is coincident with the enhanced NEP evident in the forereef since the 1980s (Figure 3c). However, it is important to note that Stephenson et al. (2014) did not find

a significant precipitation trend for Belize itself, perhaps indicative of a more indirect, regional influence of precipitation on NEP, if not through residence time and water depth, possibly through sediment and nutrient supply via nearby rivers stimulating productivity.

The abundance of tropical corals has declined throughout the Caribbean and other reef systems around the world, with one regional study demonstrating that Caribbean coral cover declined from ~60% in the 1970s to ~10% by around 2000 (Gardner et al., 2003). Although there are potentially numerous causes of this dramatic decline in coral health and abundance, these declines tend to be associated with expansion of macroalgae (Schutte et al., 2010) and/or cyanobacterial mats into reef environments. It is therefore possible that the oscillations and long-term trends of enhanced precipitation and SST warming we observe here facilitated the expansion of noncalcifiers at the expense of calcifiers, thereby decreasing NEC and increasing NEP in some parts of the MBRS.

4. Conclusions

Boron isotopes measured within coral cores from the MBRS reveal that increasing atmospheric pCO_2 over the past 100 years has affected different parts of this reef system in very different ways: backreef environments have acidified in a manner consistent with, but in excess of, the open ocean, while forereef environments have maintained relatively stable pH. Comparison with trends in linear extension also reveals that the relationship between seawater pH and coral calcification is complex, with declining pH associated with increasing extension in the backreef and constant pH in the forereef associated with declining extension in the forereef. Although counterintuitive, this finding is generally consistent with recent culture experiments showing that calcification of symbiotic zooxanthellae (Castillo et al., 2014). Although the cause of declining calcification in the forereef of the Belize region of the MBRS remains uncertain, our results suggest that ocean acidification can be ruled out.

Examination of spatial patterns in coral δ^{11} B-derived seawater pH and δ^{13} C suggests that these systems can be used to reconstruct spatial variations in NEP and NEC through time. Although the complexity of vital effects for both isotope systems requires additional ground truthing to improve the quantitative rigor of the approach, the general trends derived from the present study suggest that NEP increased in the forereef of the MBRS since the 1980s and that NEC increased in the backreef during the 1920s–1940s. The increased NEP in the forereef since the 1980s effectively countered the ocean acidification expected from anthropogenic increases in atmospheric *p*CO₂ over this interval. Conversely, the increased NEC in the backreef during the 1920s–1940s enhanced the anthropogenic acidification signal. Time series analysis revealed a quasidecadal cycle in NEC and NEP that was correlated with oscillations in temperature and rainfall in the region. Our findings not only underscore the relatively untapped potential of using coupled analyses of δ^{11} B-derived pH and δ^{13} C to trace key biogeochemical processes in coral reefs, they also highlight that, in contrast to the open ocean, the evolution of the carbonate system on coral reefs over the last century has not been controlled solely by anthropogenic CO₂.

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