Marginal Reefs Under Stress: Physiological Limits Render Galápagos Corals Susceptible to Ocean Acidification and Thermal Stress

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Abstract Ocean acidification (OA) and thermal stress may undermine corals’ ability to calcify and support diverse reef communities, particularly in marginal environments. Coral calcification depends on aragonite supersaturation (Ω > 1) of the calcifying fluid (cf) from which the skeleton precipitates. Corals actively upregulate pH\textsubscript{cf} relative to seawater to buffer against changes in temperature and dissolved inorganic carbon, which together control Ω\textsubscript{cf}. Here we assess the buffering capacity of modern and fossil corals from the Galápagos Islands that have been exposed to sub-optimal conditions, extreme thermal stress, and OA. We demonstrate a significant decline in pH\textsubscript{cf} and Ω\textsubscript{cf} since the pre-industrial era, trends which are exacerbated during extreme warm years. These results suggest that there are likely physiological limits to corals’ pH buffering capacity, and that these constraints render marginal reefs particularly susceptible to OA.

Plain Language Summary Reef-building corals regulate their internal environment to permit rapid growth, which is critical for creating the structure and function of coral reefs. However, we demonstrate that there are finite limits to the ability of corals to regulate their internal chemistry to optimize growth. This limitation will leave corals susceptible to ocean warming and acidification, particularly in sub-optimal environments. Galápagos corals already display signs of stress and an inability to maintain an optimal internal growth environment from the eighteenth century to today.

1. Introduction

The carbonate structures of coral reef ecosystems provide critical defenses against storm surge and sea-level rise, supporting billions of dollars of goods and services annually beyond their intrinsic value (Spalding et al., 2017) and highlighting the need to understand how changing ocean conditions impact coral calcification. Thermal stress and ocean acidification (OA) diminish coral calcification, as shown in both experimental systems and Free Ocean CO\textsubscript{2} Enrichment (FOCE) experiments on natural reefs (Gattuso et al., 2014). Analyses of coral density variations in cores of massive corals also reveal declining coral calcification through time (Lough, 2010). Collectively, these studies demonstrate spatially and temporally varying rates of calcification, with significant declines under recent extreme warming events and OA. Corals in the Galápagos Archipelago have been disproportionately impacted (Glynn et al., 2018), due to both extreme El Niño-related warming (Glynn, 2001; Glynn et al., 1988) and highly variable upwelling and pH/saturation state (Manzello, 2010; Manzello et al., 2008). These “marginal” reefs exhibit low species diversity and structural complexity (Cortés, 1997; Darwin & Bonney, 1889; Glynn, 2001; Glynn et al., 2017; Manzello et al., 2008), and have experienced acidification at rates of around −0.0026 (pH units, total scale) yr\textsuperscript{−1} over the last 1.5 decades (1997–2011, Sutton et al., 2014). Differential recovery rates along spatial pH gradients (Manzello et al., 2014) further demonstrate the importance of carbonate chemistry and calcification processes to reef health in this region. As CO\textsubscript{2} levels rise, changing patterns of OA and warming will increase the pressure on eastern equatorial Pacific and other marginal reef environments.
A critical question remains, however: do corals have the adaptive capacity to maintain sustainable calcification in the face of increasingly stressful environmental conditions? Here, we leverage advances in biominalization and boron isotope systematics to assess how changes in energy availability alter rates of calcification, the chemistry of the calcifying fluid, and the geochemistry of the carbonate skeleton (Table S1 in Supporting Information S1). We use this understanding of coral biominalization to elucidate the susceptibility of coral calcification to OA and thermal stress, and to assess the adaptive capacity of Galápagos (*Porites sp.*) corals to changing ocean conditions.

In reef-building corals, calcification varies in response to internal (physiological) and external (environmental) factors, and maintenance of aragonite supersaturation in their calcifying fluid ($\Omega_{AR} > 1$) is the ultimate factor that permits supercalcification and buffers against changes in seawater chemistry (McCulloch et al., 2012; Thompson, 2021). This state is achieved via upregulation of DIC and pH in response to changing environmental conditions. For example, during cooler seasons, corals upregulate the pH of their calcifying fluid ($pH_{sw}$) in response to a drop in metabolic (i.e., from zooxanthellar photosynthesis and coral respiration) DIC, resulting from reduced temperature and light (e.g., D’Olivo & McCulloch, 2017; McCulloch et al., 2017; Ross et al., 2017, 2019). Cool temperatures also slow calcification kinetics and reduce the buffering capacity of the coral calcifying fluid (hereafter “thermodynamic” factors, Guo, 2019; Georgiou et al., 2015). By upregulating $pH_{sw}$, corals maintain a stable aragonite saturation state, shifting the carbonate reactions to favor carbonate ion during the winter months and preserving their ability to calcify despite large seasonal changes in DIC availability and temperature (as reviewed by Thompson, 2021). If these processes operate across species and reef environments, corals may be able to withstand changes in seawater pH ($pH_{sw}$).

However, our understanding of coral biominalization processes largely depends on studies of modern massive corals from regions with relatively low interannual climate and geochemical variability (Figures 1a and 1b). Although a few studies have leveraged natural CO$_2$ seeps to study coral biominalization under extreme conditions (Wall et al., 2016, 2019), corals likely respond differently to sharp spatial gradients compared to temporal variations. In many marginal reef environments, strong oceanographic variability and low aragonite saturation states make reef-building corals particularly susceptible to changing ocean conditions. Further, such marginal reefs provide a potential analogue of future reef patterns, as OA broadens the coverage of sub-optimal to marginal conditions.

Here, we capitalize on the large natural gradients across the Pacific in SST variability (Figure 1a) and aragonite saturation state (Figure 1b) to understand the range of coral responses to ongoing warming and acidification. We apply a multi-proxy, multi-site synthesis of coral geochemistry, backed by a novel Earth system modelling framework, to reconstruct and contextualize the impact of environmental stresses on calcification and resiliency in Galápagos corals. We leverage geochemical tracers of coral biominalization (Table S1 in Supporting Information S1)—skeletal B/Ca ([CO$_3^-$]), $\delta^{11}$B ($pH_{sw}$), and U/Ca ([CO$_3^-$])—that constrain the calcifying fluid chemistry, including the aragonite saturation that governs calcification rate (DeCarlo et al., 2015, 2018). We combine these with paleo-environmental tracers that primarily reflect factors external to the coral calcification environment (Table S1 in Supporting Information S1): Sr/Ca (Beck et al., 1992; Corrège et al., 2000), Li/Mg (Hathorne, Felis, et al., 2013; Montagna et al., 2014), and $\delta^{18}$O (McConnaughey, 1989; Weber & Woodhead, 1972; all primarily controlled by SST); Ba/Ca (upwelling, Lea et al., 1989; G. T. Shen et al., 1992); and $\delta^{13}$C (upwelling, metabolic carbon/photosynthesis, respiration, and reproduction, G. T. Shen et al., 1992). These new recent (1976–2010) and fossil (1729–1733) Galápagos records (Wolf Island, 1°23.15’N, 91°49.90’W) significantly extend the multi-tracer data coverage prior to the industrial era, which allows us to assess the capacity of corals to buffer against changing environmental conditions. We compare our new Galápagos results with published data from the Great Barrier Reef (M. T. McCulloch et al., 2017) to contextualize results from the marginal Galápagos reef environment—a comparatively cold, low-saturation, and highly variable environment. Finally, we establish a comprehensive spatiotemporal framework for these results using simulations of ocean biogeochemistry that extend from pre-industrial to modern (Figure 1c), permitting the first cross-Pacific, multi-century synthesis of corals’ ability to buffer calcifying fluid chemistry in response to changing ocean conditions, including acidification, warming, and (internal and forced) climate variability.
2. Data and Methods

2.1. Coral Core Collection

We collected cores from modern (living) and underwater sub-fossil (i.e., deceased upon collection; hereafter “fossil”) *Porites lobata* colonies in Shark Bay, along the northeastern shore of Wolf Island, Galápagos (1°23.15’N, 91°49.90’W) in May–June 2010. Here, we analyze four cores from three colonies (two modern, and one fossil): (a) GW10-3 (modern), collected from 10 m depth; (b) GW10-10 (modern), collected from 12 m depth; and (c) GW10-4 and (d) GW10-5 collected from the same fossil colony at 13 m depth. We compare these geochemical records from Wolf to published data from Davies Reef, Great Barrier Reef (cores 13–2 and 13–3, M. T. McCulloch et al., 2017).

2.2. Sub-Sampling & Age Determination

All cores were milled for geochemical analysis at continuous 2 mm increments in 5mm-wide transects along the maximum growth axis; based on average modern extension rates (GW10-3 = 12.4 mm/year, GW10-10 = 20.3 mm/year), this sampling increment resolves sub-seasonal (bimonthly or better) variability of coral skeletal geochemistry and inferred environmental parameters. This resolution was selected based on the time-and sample-intensive nature of the ion exchange chromatography required for boron isotopic analysis; this work significantly extends the network of long, high-resolution, multi-proxy data. Modern corals were re-sampled adjacent to the original sampling transects (Jimenez et al., 2018) across intervals of known climatic extremes (e.g., large eastern Pacific El Niño events) and phases of Pacific decadal variability, while fossil cores were sampled prior to and following the depths sampled for U/Th age dating (to maximize precision of replicating and splicing these floating chronologies).

Figure 1. Map of study sites across tropical Pacific Ocean: (a) Interannual variability in sea-surface temperature (SST), calculated from standard deviation of CESM1 LME SST (see Figure S2 in Supporting Information S1 for validation against IGOSS SSTs, Reynolds et al., 2002); (b) aragonite saturation state Ω*sw* at 0 m, calculated using CO2SYS (Lewis et al., 1998) from CESM1 LME temperature, salinity, pH*sw*, and seawater dissolved inorganic carbon (DIC*sw*) over the climatological period (1970–2005); and (c) difference in CESM1 LME Ω*sw* between the modern and eighteenth century periods studied here. Simulated values for the Great Barrier Reef (Davies Reef) and Galápagos (Wolf Island) study sites are indicated by filled circles; validation of CESM1 against observational values can be found in Table S3 in Supporting Information S1.
Pre-industrial Wolf fossil cores (WLF10-04 and WLF10-5) were U/Th dated at the University of Minnesota following the procedures of (Cheng et al., 2013; Edwards et al., 1987; C. C. Shen et al., 2002). Wolf10-04 and WLF10-05 sample ages were 1732 ± 7 and 1738 ± 5 C.E., respectively (see Reed et al., 2021 for full U/Th results). These floating chronologies were tied to the complete Sr/Ca record from WLF10-4 (Reed et al., 2021) to optimize correlation among the series within the uncertainty of the U/Th dates. However, all Wolf fossil coral series are floating chronologies (i.e., they are not tied to overlapping modern records); thus, we estimate an absolute age error as ±5–7 years (based on the precision of the U/Th dates).

Age-depth models for all cores were developed using linear interpolation in MATLAB between seasonal Sr/Ca-SST tie points. Due to high interannual variability in the timing of the cool season minima, the age model relies only on warm-season tie points. Sr/Ca minima were tied to March SST maxima; tie points for modern Wolf cores (WLF10-3 and WLF10-10) are identical to those published in Jimenez et al. (2018). Data were linearly interpolated to obtain monthly records for time-series analysis. Although this approach may introduce sub-annual chronological errors, regressions among geochemical proxies that form the core of this study were performed on the raw data (prior to age modeling) and are not influenced by chronological errors or interpolation. Finally, we used Sr/Ca-SST reconstructions from GW10-3 (2010–1987; 1983–1940) and GW10-10 (2010–1985; 1982–1975) published by Jimenez et al. (2018) and Cheung et al. (2021) for comparison.

2.3. Trace Elemental Geochemistry

All trace elemental analyses were performed on a Quadrupole-ICP-MS (X-series II Q-ICP-MS, Thermo Fisher Scientific) at the University of Western Australia. First, sub-samples of 10 ± 0.2 mg of coral powder were weighed, dissolved in 500 μL of 0.51N HNO₃, agitated, and centrifuged for 1 min at 3500 rpm. A 38 μL aliquot of dissolved powder was diluted in 3 mL of 2% HNO₃ (100 ppm Ca) for trace elemental analysis; the remaining 400 μL of the dissolved powder was used for boron isotope analysis (see below). Analysis of 7Li, 25Mg, and 11B by Q-ICP-MS was performed on the 100 ppm Ca dilution, while an additional 300 μL sub-aliquot of the 100 ppm Ca solution was diluted (to 10 ppm Ca) in 2.7 mL of a 2% HNO₃ spike solution (containing ~19 ppb ⁸⁸Sr, 19 ppb ⁸⁹Y, 0.19 ppb ¹⁴¹Pr, and 0.095 ppb ²⁰⁸Bi) for analysis of ⁸⁸Sr, ⁸⁹Sr, and ²⁰⁸U. Although some recent work suggests that organic matter may bias TE/Ca values and increase analytical uncertainty (particularly for Li/Mg, Cuny-Guirriec et al., 2019), these issues were reported for green, organic-rich bands in the skeleton. As there were no green, organic-rich bands in these cores, we did not pre-treat the samples prior to geochemical analysis to avoid offsets and noise that can arise from oxidative cleaning under certain conditions (Holcomb et al., 2015; Sayani et al., 2021). Nevertheless, we note that the presence of organic matter in the samples could have caused small (1%–4%) variations in trace elemental ratios and add noise to our data. Reproducibility for the JCP-1 interlaboratory standard (2σ relative standard deviation, RSD; n = 19) was ±0.830% for Mg/Ca, ±0.636% for Sr/Ca, ±1.341% for U/Ca, ±3.649% for Li/Mg (N = 17), and ±3.651% for B/Mg (N = 17). The long-term laboratory values for JCP-1 are well within the robust standard deviation of reported values from Hathorne, Gagnon, et al. (2013): Mg/Ca 4.211 ± 0.024 mmol/mol (n = 173), Sr/Ca 8.848 ± 0.0194 mmol/mol (n = 173), Ba/Ca 7.297 ± 0.242 μmol/mol (n = 159), U/Ca 1.194 ± 0.0092 μmol/mol (n = 165), Li/Mg 1.441 ± 0.0325 mmol/mol (n = 144), and B/Ca 458.956 ± 11.790 μmol/mol (n = 144; see also D’Olivo et al., 2018).

We used published TE/Ca-SST calibrations to reconstruct SST from the (local) Sr/Ca-SST (M. T. McCulloch et al., 2017; Jimenez et al., 2018) and Li/Mg-SST (Montagna et al., 2014) relationships. For Wolf corals, we applied the Sr/Ca-SST calibration:

$$\text{Sr/Ca}_{\text{corr}}(\text{mmol/mol}) = -0.057(\pm0.001) \times \text{SST} + 10.658(\pm0.025)$$  

(1)

From weighted least squares regression of the WLF10-03 and WLF10-10 composite record against OISST between May 1987–March 2010 (Jimenez et al., 2018). The composite calibration was utilized to standardize the calibrations across cores; however, the same results were found when using core-specific calibrations for the modern corals, as the calibration equations were similar between cores (Jimenez et al., 2018). For the Davies Reef, GBR corals, we used the Sr/Ca-SST calibration obtained from local calibration with in-situ temperature data (M. T. McCulloch et al., 2017):

$$\text{Sr/Ca}_{\text{corr}}(\text{mmol/mol}) = -0.046 \times \text{SST} + 10.12.$$  

(2)
For both sites, the Li/Mg-SSTs were calculated using the calibration curve of Montagna et al. (2014). All new trace elemental geochemical data are shown in Figure 4 and S9–S10 in Supporting Information S1.

2.4. Determination of Calcifying Fluid pH and Carbonate Chemistry From Boron Systematics

The boron in the remaining 400 μL aliquot of dissolved powder (after trace elemental analysis, above) was purified by ion exchange chromatography (after M. T. McCulloch et al., 2014), and the δ11B was measured by MC-ICP-MS using a NU Plasma II at the University of Western Australia. The measured isotopic ratio of 11B and 10B of the carbonate samples were expressed relative to that of the NIST SRM 951 boric acid standard, in standard delta notation (in units of per mil or ‰):

\[
\delta^{11} B_{\text{carb}} = \left( \frac{^{11}B/^{10}B}{^{11}B/^{10}B_{\text{standard}}} \right) \times 1000.
\]  

Reproducibility for the JCP-1 interlaboratory standard across these runs (2σ; n = 29) was ±0.22‰. Further, the long-term laboratory JCp-1 value and reproducibility of 24.36 ± 0.34‰(2σ; n = 101; see also D’Olivo & McCulloch, 2017) agree well with reported values with and without oxidative pre-treatment (Gutjahr et al., 2021). Therefore, although the analytical uncertainty of our results may be slightly higher because the samples were not pre-cleaned (Gutjahr et al., 2021), the reported values are well within error of the pre-cleaned values from Gutjahr et al. (2021). The 2σ uncertainties in this study are on par with that of pre-cleaned samples (n = 29), while the long-term average falls between that of published values with and without cleaning (n = 101). Finally, previous work suggests that δ11B is relatively insensitive to sample cleaning methods (Holcomb et al., 2015).

We used paired boron isotope and B/Ca ratios to determine the pH and carbonate ion concentration, leveraging three key features of boron isotope systematics (as reviewed by DeCarlo et al., 2018; Thompson, 2021). First, boron speciation in seawater depends strongly on pH, with borate ion (B(OH)4−) dominating at higher pH and boric acid (B(OH)3) dominating at lower pH (<~8.5). Second, boron isotopes are strongly fractionated between the two species, with a +27‰ offset between borate and boric acid. Taken together, as pH decreases, the fraction of boron as borate decreases and the δ11B increases. Third, as corals calcify from a semi-isolated calcifying fluid, borate may substitute for the carbonate ion (CO32−; Sen et al., 1994). Although there are multiple pathways by which this could occur, recent inorganic precipitation studies (Holcomb et al., 2016) suggest that it likely occurs via de-protonation and co-precipitation with CO32− (Noireaux et al., 2015), rather than via bicarbonate or some mixture of the two, as previously proposed (Allison et al., 2014).

The initial calcifying fluid δ11B and total boron concentrations are thought to be the same as that of seawater, as seawater serves as the source of boron; further, the boron isotopic composition and concentration remains relatively constant during calcification, due to low partitioning coefficient (Kw) of B/Ca between aragonite and seawater (i.e., B is strongly excluded from the skeleton during precipitation, Holcomb et al., 2016). We note that diffusion may violate these assumptions under certain conditions; for example, diffusion of isotopically distinct boric acid may alter the δ11B relative to seawater (Gagnon et al., 2021) or increase boron concentrations relative to seawater when pH is elevated. However, there is no experimental evidence for these confounding factors within tropical, symbiont-bearing coral species; as symbionts provide an additional critical source of DIC to the calcifying fluid, biomineralization processes in symbiont-bearing corals are markedly different from that of the cold-water species for which these limitations have been identified. We therefore follow the approach of other recently published studies in this regard (Chalk et al., 2021; DeCarlo et al., 2018; D’Olivo et al., 2019; Ross et al., 2017, 2019; M. T. McCulloch et al., 2017).

As a result of these processes, the skeletal δ11B reflects the pH of the calcifying fluid (pHcf), while the [B] reflects both pH and the [CO32−] (DeCarlo et al., 2018; Holcomb et al., 2016). We calculate pHcf from δ11B of the carbonate skeleton (after Zeebe & Wolf-Gladrow, 2001):

\[
pH_{cf} = pK_B - \log \left( \frac{\delta^{11}B_{sw} - \delta^{11}B_{carb}}{\alpha_{B(3−B4)}\delta^{11}B_{carb} - \delta^{11}B_{sw} + 1000(\alpha_{B(3−B4)} - 1)} \right),
\]

where the δ11B of seawater (δ11Bsw) was defined as 39.61‰ (Foster et al., 2010), the boron isotope equilibrium constant (α(B3−B4)) was set to 1.0272 (Klochko et al., 2006), and the dissociation constant of boric acid (pKb) was calculated from temperature, salinity and pressure (after Dickson, 1990). To standardize methods across cores
(as in situ data is not available for all sites or time periods), we used Li/Mg-derived SSTs and Simple Ocean Data Assimilation (SODA) sea-surface salinity (SSS). We used mean climatological SODA SSS (33.5 PSU) for fossil analyses (prior to the industrial era).

Empirical constraints on the B/Ca partitioning coefficient between aragonite and seawater and its dependency on pH$_f$ (Holcomb et al., 2016) permit reconstruction of carbonate ion concentration in the calcifying fluid from paired $\delta^{11}$B-pH$_f$ and B/Ca measurements (DeCarlo et al., 2018):

$$K_D \equiv (B/\text{Ca})_{CaCO_3} \times \frac{[CO_3^{2-}]_{cf}}{[B(OH)_4]_{cf}},$$  \hspace{1cm} (5a)

$$K_D = 0.00297 \exp(-0.0202[H^+]_{cf}),$$ \hspace{1cm} (5b)

and

$$[CO_3^{2-}]_{cf} = \frac{K_D \times [B(OH)_4]_{cf}}{(B/\text{Ca}_{CaCO_3})},$$  \hspace{1cm} (6)

where Equation 5b follows the formulation of M. T. McCulloch et al. (2017). Although there continues to be debate over the best $K_D$ formulation (DeCarlo et al., 2018), Equation 5b is likely to be most accurate for tropical reef-building corals as it does not include the (Mavromatis et al., 2015) experimental data, which was collected from NaCl solutions (rather than seawater) at very low [CO$_3^{2-}$] relative to that of coral CF.

As reviewed by DeCarlo et al. (2018), uncertainties still remain with regards to the most accurate formulation for $K_D$, and the degree to which Ca$^{2+}$ is upregulated within the cf. We evaluated the sensitivity of our results (see Figures S3 and S4 in Supporting Information S1) to the $K_D$ formulation, following the equations of Holcomb et al. (2016); M. T. McCulloch et al. (2017); DeCarlo et al. (2018) and the boron systematics package of DeCarlo et al. (2018), as well as using a constant $K_D$ of 0.002 (after Allison, 2017). Our sensitivity tests show that these uncertainties only marginally impact the absolute magnitude of inferred DIC$_{cf}$ and do not influence the relative changes across sites and time periods (the focus of this work). Further, the inferred DIC$_{cf}$ upregulation is higher using the $K_D$ formulation of M. T. McCulloch et al. (2017) (Figure S3 in Supporting Information S1); therefore, our chosen approach produces the most conservative change in DIC$_{cf}$ and $\Omega_{cf}$ under warming and acidification. We similarly test the impact of Ca$^{2+}$ upregulation relative to seawater on resulting $\Omega_{cf}$ calculations. For this, we use the mean and ±1 standard deviation from independent micro-sensor measurements (see Sevilgen et al., 2019, Table 1). These sensitivity analyses demonstrate that uncertainties Ca$^{2+}$ impact the absolute magnitude of $\Omega_{cf}$ within colonies (Figure S5 in Supporting Information S1), but not the relative differences among colonies, sites, or time periods (the focus of this study). We therefore utilize the most conservative approach, and report results using a Ca$^{2+}$ scaling factor of 1, which is the lower (-1σ) bound from from Sevilgen et al. (2019). Inferred trends in $\Omega_{cf}$ and calcification would be greater if a constant $K_D$ or higher Ca$^{2+}$ are assumed (Figures S4 and S5 in Supporting Information S1). Therefore, the results reported here are the most conservative estimate of inferred $\Omega$ and calcification changes from preindustrial to modern conditions.

Dissolved inorganic carbon is calculated from the pH$_f$ (Equation 4) and [CO$_3^{2-}$]$_f$ (Equation 6) using CO2SYS software (Lewis et al., 1998) and the following constants: carbonate species dissociation (Dickson & Millero, 1987; Mehrbach et al., 1973), borate and sulphate dissociation (Dickson, 1990), and aragonite solubility (Mucci, 1983). Finally, we explore the relationship between pH, DIC and $\Omega$ of the coral calcifying fluid and Sr/ Ca-SST (note: we utilize Sr/Ca-SST as a quasi-independent SST estimate rather than Li/Mg-SST, as the latter was used in Equation 4). Our findings are robust to the paleo-thermometer used to assess the impact of temperature on coral carbonate chemistry (e.g., Figure S8 in Supporting Information S1).

### 2.5. Stable Isotope Geochemistry

Stable oxygen and carbon isotope ratios ($\delta^{18}$O and $\delta^{13}$C) were analyzed on a Thermo Delta V Plus mass spectrometer, coupled to a Kiel IV carbonate preparation system, in the PACE lab, at the University of Michigan’s Earth and Environmental Sciences department. Analyses were performed on splits of the same powders analyzed for trace elemental chemistry and paired $\delta^{11}$B-B/Ca boron systematics. Long-term analytical precision (1 σ) of Luxor
internal carbonate standard was 0.08‰ for δ¹⁸O and 0.05‰ for δ¹³C. All new stable isotope data are shown in Figures 4 and S10 in Supporting Information S1.

2.6. Statistical Analysis

Ordinary least squares regressions (OLS) were used to assess relationships among geochemical parameters within and among coral colonies, and in upregulation with respect to seawater conditions. First, OLS regressions were performed among reconstructed calcifying fluid and skeletal geochemical parameters (Figures 2, S1, S3–S6, and S8 in Supporting Information S1). ANCOVA and multiple comparisons were then utilized to assess differences in the relationship among groups (i.e., among individual cores, or among fossil Wolf, modern Wolf, and GBR corals). Finally, OLS was utilized to assess the relationship between average upregulation of pHₙₐ, DICₑ, and Ωₑ and seawater chemistry and temperature (Tables S4 and S5 in Supporting Information S1). Confidence intervals (95% CI) were determined from the 5th and 95th percentiles of 1000 random draws of the distribution of upregulation estimates (based on the standard deviation and mean of each record).

2.7. Coral Densitometry and Calcification

Skeletal density was measured using a quantitative X-ray scanning method developed at the Australian Institute of Marine Science (Anderson et al., 2017, supplementary methods) alongside six compressed Porites sp. powder standards. These standards were used to calibrate X-ray grayscale values to known density, by applying a linear fit between known density (multiplied by thickness) and the natural log of each standard’s mean grayscale value. Grayscale values were measured from the background-corrected X-ray positives using Fiji software. Analytical precision of these X-ray density measurements was estimated using an additional standard with a known density (2.3977 g cm⁻²) and thickness (6.86 mm) with values within the typical range of massive Porites spp. coral slabs. The average density of this quality control standard across all five X-rays used in this study was 2.3655 g cm⁻²; thus, we report an uncertainty of 0.043 g cm⁻² or 1.8%.

For each core, grayscale values were measured along 4 mm-wide transects on either side of the geochemical transect. We report density values from each transect, as well as the average across both transects (to account for micro-scale variations in density associated with skeletal architecture). For each transect, density was calculated using the standard calibration curve, normalized by slab thickness. Thickness was measured at 0.125 cm increments along two transects, and the average thickness was interpolated to 0.005 cm (the sampling resolution of the X-ray density measurements).

Annual growth metrics (density, extension, and calcification) were calculated from warm season to warm season using annual tie points (Sr/Ca minima, SST maxima). This approach was utilized as the seasonal cycle was more clearly identifiable in the Sr/Ca series (relative to that of the growth series). Extension was calculated as the distance between successive Sr/Ca minima, and calcification as the product of extension and annual average skeletal density.

2.8. Seawater Carbonate System

Seawater carbonate chemistry (TCO₂, Total Alkalinity [TA], pCO₂, pH, and Ω₉₅) were obtained from (D. P. Manzello, 2010; Manzello et al., 2014; Humphreys et al., 2018). Briefly, seawater samples were collected during the cool (n = 24) and warm (n = 21) seasons over multiple years in 500 mL borosilicate glass bottles from 7 study sites throughout the archipelago: (a) Bartolomé, Santiago Island; (b) Santa Fe Island; (c) Punta Bassa, San Cristóbal Island; (d) Punta Pitt, San Cristóbal Island; (e) Devil’s Crown, Floreana Island; (f) Gardner Bay, Española Island; and (g) Darwin Island (N = 7; summary statistics obtained from Humphreys et al., 2018). Here, we utilize the mean (±standard error of the mean, SEM) values to assess the relationship between pHₑ and DICₑ (calculated from paired coral δ¹¹B and B/Ca) and regional changes in the seawater CO₂ system. However, available measurements are discrete, disjointed snapshots, and therefore lack temporal information with which to identify variability on interannual and longer timescales. Further, Ωₑ at Wolf Island is expected to display higher mean values and lower seasonal variability (see Manzello, 2009, Figure 1) than the seawater collection sites of (Manzello, 2010), as upwelling and equatorial undercurrent (EUC) strength and variability are weaker.
Figure 2.
at Wolf Island. As values from Wolf Island are not publicly available, analyses were performed using both the in situ data from Darwin Island (Manzello et al., 2014; Humphreys et al., 2018) and Community Earth System Model version 1 (CESM1).

### 2.9. Community Earth System Model Biogeochemistry

Given the sparse network of seawater inorganic carbon measurements (i.e., DIC, pH, alkalinity) with which to calculate seawater aragonite saturation state, we use the CESM1 Last Millennium Ensemble (LME, Otto-Bliesner et al., 2016) and Large Ensemble (LE, Kay et al., 2015) to compare the chemistry of the coral calcifying fluid to that of local seawater. This approach facilitates comparison across sites, as well as among eighteenth century (LME), twentieth century (LME and LE), and end of 21st century (LE) conditions. The CESM1 marine ecosystem-biogeochemical module (Hurrell et al., 2013) permits analysis of the entire carbonate systems across space and time, permitting the first multi-site, multi-century synthesis of coral calcifying fluid chemistry in response to changing ocean conditions.

The CESM1 LME simulation was validated against OISST SSTs (Reynolds et al., 2007, Figure S2 in Supporting Information S1), SODA SSS (Carton & Giese, 2008, not shown), buoy data (Sutton et al., 2019), seawater samples described above (Table S3 in Supporting Information S1), and the spatially interpolated climatology (1972–2013) from GLODAP version 2 (Lauvset et al., 2016, Table S3 in Supporting Information S1). CESM1 simulated pH and calculated $\Omega_{sw}$ compare well with the observations across the tropical Pacific, with differences of less than 0.05 and $<$0.5 (RSDs of $<$0.6% and 8%), respectively (Table S3 in Supporting Information S1). Further, these discrepancies may be at least partially attributed to the comparison of discrete in-situ snapshots of ocean pH with the climatological value over different baseline periods (over which there is a decreasing trend across the tropical Indo-Pacific).

We calculate $\Omega_{sw}$ from CESM1 LME (full forcing scenario) and LE (Representative Concentration Pathway; RCP8.5) simulated SST, SSS, pH, and DIC using CO2SYS (as described above). Combining the simulated seawater pH, DIC, and $\Omega$ with boron-derived estimates of coral calcifying fluid pH, DIC, and $\Omega$, we estimate the percentage upregulation of calcifying fluid geochemistry. For example, the percent change (henceforth “Pchange”) in aragonite saturation is calculated as:

$$\text{Pchange}_{\Omega} = \frac{\Omega_{cf} - \Omega_{sw}}{\Omega_{sw}} \times 100, \quad (7)$$

where $\Omega_{sw}$ represents the average over the time period overlapping each coral record from CESM1 LME and/or LE.

We perform sensitivity tests at the GBR site, where an in-situ seawater timeseries is available, to show that CESM1 LME and LE reproduce the observed Pchange$_{\Omega}$ (i.e., relative to seawater observations) to within ±26% (LME) and ±0.5% (LE), respectively (Table S8 in Supporting Information S1). Much of the discrepancy between LME and observed Pchange can be attributed to differences in the time periods of coverage. Therefore, two sensitivity tests were used to assess: (a) the impact of using the annual average, seasonal average (cold vs. warm season), or monthly seawater value, and (b) the impact of using the LME projected values versus using the LE values over the post-2005 interval (i.e., after the final year of the LME). Because the Pchange seasonal variability is dominated by the variability in the coral calcifying fluid (which is $\gg$ seawater variability), these sensitivity tests demonstrate that there is no difference in the mean Pchange if the average seawater value is used in place of the observed temporal evolution of in situ $\Omega_{sw}$ (M. T. McCulloch et al., 2017). Further, this approach generates the most conservative estimate of the Pchange variability at each site (i.e., 1 $\sigma$ = 23 & 32%; Table S8 in Supporting Information S1). The second sensitivity test demonstrated that LE-simulated seawater values displayed the best match with the in situ data over the post-2005 period ($\Delta$Pchange$_{\Omega} < 0.5\%$). Although there are

![Figure 2](https://example.com/figure2.png)

**Figure 2.** Comparison of the relationships among geochemical proxies, Wolf eighteenth-century fossil (red squares) and modern (twentieth century, orange circles) versus Great Barrier Reef modern (blue circles): (a) Sr/Ca-SST versus $p_{f}^{\text{Ca}}$, (b) Sr/Ca-SST versus $\delta^{13}C$, (c) Sr/Ca-SST versus $\Omega_{cf}$, (d) Sr/Ca-SST versus $\delta^{13}C$ and without flier outlined in gray, and (f) Sr/Ca versus Mg/Ca. Comparison of the pre-1997/98 thermal stress (blue), and post-1997/98 thermal stress (red) Sr/Ca-SSTs versus $p_{f}^{\text{Ca}}$ for all modern coral data (black) is shown in (d). In all panels, roman numerals (I-III) denote that the slope of the relationship is significantly different from other groups, based on ANCOVA and multiple comparisons (where a significant difference among groups was identified). Groups with the same roman numeral are not significantly different from one another.
no contemporaneous seawater samples collected near Wolf Island, Ω Pchange values using seawater data from nearby Darwin (collected in June 2012) are within the 1 σ range (±29%) of the CESM1-based estimates for WLF10-10a (ending in 2010, Table S9 in Supporting Information S1). We therefore conservatively reported an uncertainty of ~±30% for all Pchange estimates.

We also apply the method of (D’Olivo et al., 2019) to deconvolve the relative contribution of thermodynamics (i.e., SST-driven changes in calcification and/or buffering capacity, Guo, 2019) and pH<sub>sw</sub> in the observed pH<sub>cf</sub> trends and seasonal variability. Briefly, we performed a multivariate linear regression between CESM1 simulated temperature and pH<sub>sw</sub> (independent predictors) and pH<sub>cf</sub> (dependent predictand). The sensitivity of Wolf coral pH<sub>cf</sub> to SST and pH<sub>sw</sub> can be expressed as:

$$\text{pH}_{\text{cf}} = 0.26 \times \text{pH}_{\text{sw}} - 0.0019 \times \text{SST} + 6.34$$

Similar results were obtained when Sr/Ca-SSTs we used in place of CESM1 simulated SSTs. To quantify the role of SST and pH<sub>sw</sub> in the observed trends (WLF10-10 and fossil vs. modern) and seasonal variability, we model pH<sub>cf</sub> from Equation 8 using either (a) the average pH<sub>sw</sub> and simulated SST, or (b) the average SST and simulated pH<sub>sw</sub>, respectively.

### 2.10. Predicted Changes in Coral Calcification

Finally, we use the IpHRAC model from (M. McCulloch et al., 2012) to predict the changes in calcification rate (G) from Ω<sub>cf</sub> between time periods (i.e., eighteenth and twentieth):

$$G = k \times (\Omega_{\text{cf}} - 1)^n$$

where

$$k = -0.0177 \times \text{SST}^2$$

and

$$n = 0.0628 \times \text{SST} + 0.0985.$$  

Omega<sub>cf</sub> is calculated from simulated pH, Ω<sub>sw</sub>, SST, and SSS and the Pchange (%) upregulation, as described above. Calcification rates are reported as percent changes relative to the baseline period (1970–2005, unless otherwise noted).

### 3. Results and Discussion

#### 3.1. Seasonal pH, DIC and Ω of Coral Calcifying Fluid

Here we compare new reconstructions of SST and calcifying fluid geochemistry (Table S1 in Supporting Information S1) from modern and subfossil Galápagos coral cores with published reconstructions from the GBR (M. T. McCulloch et al., 2017; D’Olivo et al., 2019; Ross et al., 2017). Two eighteenth century coral cores collected at Wolf Island show that as SST increases, pH<sub>cf</sub> decreases (despite the regional pH<sub>sw</sub> increase; Figure 2a and Figure S1a in Supporting Information S1). The slope of this relationship (WLF04: −0.022 pH units per °C, N = 33, r² = 0.52; WLF05: −0.033 pH units per °C, N = 45, r² = 0.43; Figure S1a in Supporting Information S1) is nearly identical to that found among replicate modern corals from the GBR (Davies-02: −0.035 pH units per °C, N = 50, r² = 0.82; Davies-03: −0.020 pH units per °C, N = 54, r² = 0.80). The seasonal pH<sub>cf</sub> change is also similar among GBR modern corals and the Wolf fossil coral, with a −0.03 to −0.06 unit change between the average warm and cold seasons (Table S2 in Supporting Information S1) and a range of 0.2–0.3 pH units at each site. However, the SST-pH relationship weakens in the two modern (twentieth century) Wolf corals, which display a reduced seasonal pH range (ΔpH = −0.003 to −0.02, Table S2 in Supporting Information S1) and a weaker relationship with temperature (i.e., a shallower slope and lower r²) compared to fossil Wolf cores (Figure S1a in Supporting Information S1).

Comparing the modern and fossil data from Wolf, we demonstrate that the pH<sub>cf</sub>-SST relationship is significantly weaker in the modern corals than in the fossil corals. In contrast, the Wolf fossil and GBR modern corals are not
significantly different from one another (Figure 2a). The greater SST range in modern cores (Figure 2a, x-axis) would by itself strengthen this relationship (as in D’Olivo et al., 2019) and therefore cannot explain the observed patterns; we therefore infer that the weakening is likely driven by reduced pH upregulation (Figures 3c and 3d), due to the impacts of OA and/or thermal stress (rather than by temperature-induced changes in calcification or buffering capacity alone Guo, 2019). The difference in slope between the fossil and modern corals equates to 7%–40% difference in H+ ions in the calcifying fluid (with larger changes at lower temperatures, Figure 2a). As a result, $\Omega_{\text{cf}}$ displays a significant positive relationship with SST in modern Wolf corals (Figures 2c, 3c and 3d), with up to 5% lower saturation during the cold season (September-November; SON) relative to the warm season (Table S2 in Supporting Information S1). In contrast, there is no relationship between $\Omega_{\text{cf}}$ and temperature in the fossil coral (Figures 2c, 3a and 3b) and <1.5% change in $\Omega_{\text{cf}}$ seasonally (Table S2 in Supporting Information S1), though we note that a significant $\Omega_{\text{cf}}$-temperature relationship is observed in the WLF04 data alone (Figure S1d in Supporting Information S1). These results indicate that the fossil coral maintained a steady aragonite saturation state in its calcifying fluid across seasonally varying environmental conditions, while the modern Wolf corals did not. Put another way, modern Wolf corals appear to have partially lost their ability to buffer calcifying fluid chemistry against changes in seawater pH and $\Omega$. This loss of buffering capacity—shown here for the first time—implies a loss of resilience that is likely to lead to reduced calcification under continued environmental change.

Figure 3. Schematic overview of the main seawater and physiological controls (cold season a,c; warm season b,d) on calcifying fluid and skeletal geochemistry in Galapagos eighteenth-century fossil (a-b) and modern (c-d) corals, identified in the current study. Figure 3 modified from (Thompson, 2021), with polyp artwork by E. V. Reed. Arrows indicate the sign of the change, with the thickness indicating the relative magnitude of the change; “—” denote variables with limited to no change.
3.2. Reproducibility

The mean and seasonal-interannual variance in calcifying fluid geochemistry were broadly reproducible across cores from both periods (within and among cores at a single site; Table S2 in Supporting Information S1, Figure 4, S1 & S10 in Supporting Information S1). However, an anomalously low $\delta^{11}$B and B/Ca departure in core

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**Figure 4.** Time series of boron-derived calcifying fluid geochemistry, Sr/Ca-SSTs, and skeletal density: Wolf eighteenth-century fossil (red) and modern (twentieth century, orange) versus Great Barrier Reef modern (blue). (a) $\delta^{11}$B (permil), (b) B/Ca (μmol/mol), (c) pH$_{cf}$ (total scale), (d) DIC$_{cf}$ (μmol/kg), (e) $\Omega_{cf}$, (f) Percent upregulation of pH$_{cf}$ with respect to pH$_{sw}$ (%), (g) Percent upregulation of DIC$_{cf}$ with respect to DIC$_{sw}$ (%), (h) Percent upregulation of $\Omega_{cf}$ with respect to $\Omega_{sw}$ (%), (i) Sr/Ca-SSTs (°C), and (j) skeletal density (g/cm$^3$). See Methods for how these parameters were derived from proxy and model data. Gray shading depicts the range of eighteenth-century fossil geochemical values; red shading depicts warm anomalies associated with the 1997/98 El Niño event; mean geochemical values are denoted by dotted lines on each series.
WLF05 co-occuring with a low-density and high Sr/Ca-SST anomaly in 1731–1732 emphasizes the need for further work to assess the impact of skeletal density, microstructure (Chalk et al., 2021), and transect quality (Reed et al., 2019, 2021) on skeletal geochemistry within a single colony. Such within colony variations are likely to be more severe at marginal reef sites like the Galápagos Islands, where corals are susceptible to boring bivalves and display lobate growth structure and complex microscale growth features, such as convergent corallite fans, changes in growth direction, and corallites angled relative to the sampling plane (Reed et al., 2021). Nevertheless, outside this short-lived anomaly, the geochemical relationships reported here were reproducible within replicate cores from a single Galápagos fossil coral colony, with no significant differences in slope between the replicate fossil cores (Figure S1 in Supporting Information S1). The only exceptions were the slope of the relationship between δ13C and DIC cf (Figure S1e in Supporting Information S1) and the intercept of the DIC cf-SST relationship (Figure S1b in Supporting Information S1)—suggesting that proxies for metabolic activity may be most susceptible to non-environmental or physiological factors (e.g., skeletal microstructures, overall transect quality, symbiont density and composition, and/or shading within colonies with complex 3D structures), as previously observed for δ13C among transects. Nevertheless, the reproducibility of these relationships suggests that this approach can help expand our knowledge of calcifying fluid geochemistry prior to the industrial era.

Further, sensitivity tests demonstrate that the differences in pH and Ω upregulation across sites and time periods are robust regardless of the choice of Ksp and [Ca2+]sw (Figures S3–S5). Further, the values are within the range of those obtained through independent micro-sensor measurements (Sevilgen et al., 2019); recent work comparing δ11B- and microelectrode-based pHcf support the utility of δ11B as a proxy for diurnally averaged pHcf (Guillermic et al., 2020).

3.3. Dissolved Inorganic Carbon & δ13C Variability

The controls on pH upregulation and DICcf likely differ across sites. In Australia, seasonal upregulation of pHcf occurs in response to seasonal variations in temperature (Guo, 2019; D’Olivo et al., 2019), pHsw (D’Olivo et al., 2019), and metabolic DIC availability (M. T. McCulloch et al., 2017), with lower DIC during the winter months due to reduced light and cooler temperatures (M. T. McCulloch et al., 2017). This mechanism was proposed in the GBR and Ningaloo Reef, Australia, where both DICcf (Figures 2b, 3a, 3b, and S1b in Supporting Information S1) and DICcf/seawater dissolved inorganic carbon (DICsw) display a strong positive relationship with temperature (M. T. McCulloch et al., 2017). This pH seasonality is consistent amongst a wide range of reefs, including the GBR, Coral Sea, Western Australia, Caribbean, and Central Pacific (Chalk et al., 2021; D’Olivo & McCulloch, 2017; D’Olivo et al., 2019; Hemming et al., 1998; Knebel et al., 2021; M. T. McCulloch et al., 2017; Pelejero et al., 2005; Ross et al., 2019). However, all of these sites have fundamentally different dynamics than in the Galápagos, where the cool season experiences upwelling of DIC-rich waters (Kessler, 2006, Figures 3a and 3b) that impacts the seasonality of CF chemistry. As a result of these confounding DICsw and metabolic DIC signals, we find that DICcf is nearly independent of temperature in modern Wolf corals (Figures 2b, 3c and 3d; though note a very weak positive relationship observed in core WLF03, Figure S1b in Supporting Information S1) and displays a weak positive relationship with temperature in the Wolf fossil coral (Figures 2b, 3a and S1b in Supporting Information S1). Further, DICcf is upregulated by a near-constant factor of ~2 relative to DICsw in modern Wolf corals, compared with a stronger, seasonally varying DICcf enrichment in GBR corals (DICcf/DICsw = 2.2–3.2, Table S2 in Supporting Information S1) and Galápagos fossil corals.

Comparison of the carbon isotopic (δ13C) variability among cores may explain why pH regulation is weaker in modern Wolf corals (Figure 2e). First, a relationship between δ13C and the DICcf in Wolf modern and fossil corals is weak or absent, suggesting that metabolic processes and upwelling contribute approximately equally to the carbon pool at this site. During the cool season, both metabolic processes (which preferentially remove light carbon, enriching the carbon pool and increasing skeletal δ13C; as reviewed by Swart, 1983) and upwelling (which contributes isotopically light carbon, decreasing skeletal δ13C) contribute to DICcf, therefore the signals compensate, reducing δ13C variance relative to that of DICcf (e.g., Figure 3a and 3b). Nevertheless, we note a weak negative relationship between δ13C and DICcf in core WLF03 (Figure S1e in Supporting Information S1), in addition to the consistently more negative δ13C values in the modern samples from the burning of fossil fuels (“Suess effect”; Keeling, 1979). Although additional data are needed to assess the complex interplay of DIC variability at this site, these results suggest that the upwelling of isotopically light carbon is increasingly dominating the DICcf pool as the seawater DIC pool becomes isotopically lighter and the coral-algae symbiosis becomes increasingly
stressed. Indeed, a significant relationship between δ¹³C and DIC$_{sw}$ is only present in the post 97/98 data (Figure S6f in Supporting Information S1), driven primarily by large isotopically heavy, low DIC anomalies during and following thermal stress and bleaching.

To assess the strength of DIC upregulation, we use simulated values for seawater carbonate parameters that are unavailable from coral proxies (see Methods), but that compare reasonably well to the limited available direct seawater observations at nearby locations, collected over disparate time periods (Table S3 in Supporting Information S1). We find that Galápagos modern DIC$_{sw}$ never reaches above 2.2 times that of simulated DIC$_{sw}$ (DIC$_{sw}$ max = 4,654 μmol/kg vs. DIC$_{sw}$ max = 2,091 μmol/kg; Manzello, 2010), whereas the fossil coral DIC$_{sw}$ reaches as much as ∼2.8 times that of seawater (5,663 μmol/kg, which is within the range observed at the GBR, Figure 2b). These results are consistent with a larger contribution of metabolic carbon to the DIC pool in the fossil coral (values DIC$_{cf}$/DIC$_{sw}$ > 1), with large seasonal (Table S2 in Supporting Information S1) and interannual variability (Figure 4d) in DIC$_{sw}$ that reflects the relative strength of upwelling and photosynthetic carbon fixation in response to light and temperature. Further, the weak relationship between DIC$_{cf}$ upregulation and Ω$_{sw}$ across all Wolf corals (Table S4 in Supporting Information S1) suggests that this decrease in DIC$_{cf}$ variability from pre-industrial conditions is likely driven primarily by dysbiosis (i.e., bleaching or loss of healthy coral microbiome and thus a reduction in metabolic carbon) associated with thermal stress, rather than OA. This is consistent with DIC$_{cf}$/DIC$_{sw}$ departures of <1 (i.e., loss of metabolic carbon) during the 1997/98 thermal stress in both modern cores (equating to a 14%–34% reduction in DIC upregulation, Figure 4g). Similar reductions in DIC$_{cf}$ upregulation are observed during other warm extremes in the modern record, whereas DIC upregulation is highest during warm periods in the fossil record. Our results therefore add to the growing body of work identifying adverse effects of thermal stress and bleaching on coral CF chemistry under ocean warming (Cheung et al., 2021; Dishon et al., 2015; D’Olivo & McCulloch, 2017; D’Olivo et al., 2019; Schoepf et al., 2015, 2021). The changes in DIC upregulation identified here imply that extreme thermal stress undermines coral health via photosynthetic reductions that coincide with weak upwelling (and thus feeding capacity); together, these changes deprive the colony of the energy needed to drive the Ca-ATPase pump and/or other active pathways (e.g., other alkalinity pumps or paracellular transport) that upregulate pH$_{sw}$, leaving them more susceptible to regional changes in DIC$_{sw}$ and pH$_{sw}$.

Taken together, these results suggest that DIC$_{cf}$ variability in Wolf corals reflects a complex seasonal interplay between upwelling (cold, high DIC$_{sw}$, low δ¹³C$_{DIC}$; May-Nov cold season) and photosynthetic/metabolic (warm, high DIC$_{sw}$, high δ¹³C$_{cf}$; Dec-April warm season) processes, the latter of which contributes less to the carbon pool in modern Wolf corals. Regional upwelling elevates both concentrations and variability of DIC$_{sw}$; these combine with the coral's metabolic variations to produce fundamentally different DIC$_{cf}$ dynamics at this site relative to the GBR. In other words, in Galápagos corals, pH upregulation is partly driven by variations in the seawater carbon pool, rather than changes in metabolic pathways alone. We thus find that seasonal pH$_{cf}$ variations at Wolf (Table S2 in Supporting Information S1) are driven primarily by seasonal temperature and pH$_{sw}$ variability (e.g., 73% and 33%, respectively, in the longest core WLF10-10; after D’Olivo et al., 2019, see Methods). These results imply that Galápagos corals are more sensitive to environmental drivers, whereas metabolic processes can regulate cf chemistry more strongly in GBR corals.

3.4. Temporal Variability in pH$_{cf}$ & Impact of Thermal Stress

Comparing the temporal evolution of pH$_{cf}$ among GBR and Wolf corals over the late twentieth century supports our interpretation that corals experience difficulty upregulating pH$_{cf}$ as seawater conditions become less favorable. First, modern Wolf corals display an abrupt drop and subsequent rise in pH$_{cf}$ during and following the 1997/98 El Niño event (Figure 4c), respectively; this event was characterized by extreme temperature anomalies (Cheung et al., 2021; Jimenez et al., 2018; Figure 4i), stress and bleaching (Glynn, 2001). The decrease in pH$_{cf}$ (towards ambient values) likely resulted from a combination of the loss of metabolic DIC from symbiotic photosynthesis (weakening the ability of corals to regulate their internal pH via the Ca-ATPase or other alkalinity pumps), temperature-induced changes in buffering capacity, and the bleaching-related reduction in calcification rate. The latter is supported by the greater change in pH$_{cf}$ in core WLF-3, in which calcification rate declined by 26% in 1998 (Figure S7 in Supporting Information S1). In turn, these changes impact Ω$_{sw}$ regulation (Figures 4e and S6f in Supporting Information S1) and calcification, and thus the imprint of Rayleigh fractionation on the widely utilized Sr/Ca-SST proxy (with less fractionation following bleaching, suggesting a slowdown in calcification, Figure S6h in Supporting Information S1), though Cheung et al. (2021) demonstrate that the Sr/Ca-SST
To understand how corals will respond to ongoing and future environmental changes, it is critical to assess the limits to calcifying fluid homeostasis. Our synthesis of modern and fossil corals living under contrasting seawater conditions suggests that there may be a physiological limit to the capacity of corals to upregulate pH across sites and time periods with different baseline seawater chemistry. Here, we demonstrate that despite large changes in seawater chemistry between the eighteenth century and modern periods inferred from model simulations (Figure 1c), there is no relationship between Ω<sub>sw</sub> and the upregulation of Ω<sub>cf</sub> in Galápagos corals (Table S4 in Supporting Information S1). In other words, Ω<sub>sw</sub> has not had a detectable influence on upregulation capacity, implying that Galápagos corals have not adapted their capacity to regulate Ω<sub>cf</sub> in response to thermal extremes and OA since the pre-industrial era. Therefore, although they continue to regulate their internal growth environment at maximum capacity, the resulting calcifying fluid saturation levels are significantly lower in modern corals due to OA (Figures 3c and 3d vs. 3a and 3b).

Our results contrast with the apparent pH “homeostasis” observed in extreme environments near submarine seeps in Papua New Guinea (Wall et al., 2016) and Puerto Morelos, Mexico (Wall et al., 2019) and in the Heron Island (GBR) FOCE (Georgiou et al., 2015). At these pCO<sub>2</sub> extremes, Porites spp. corals show a strong relationship between Ω<sub>cf</sub> upregulation and seawater conditions (e.g., ΔΩ<sub>cf</sub> of 214% and 270% per unit change in Ω<sub>sw</sub>, respectively, Table S4 in Supporting Information S1). However, in both scenarios, Ω<sub>sw</sub> was 19%–82% lower than observed on any modern reefs studied here. Further, seep corals have persisted in these conditions for multiple generations and likely have acclimatized and/or adapted to low seawater saturation over long time periods. Therefore, such sites are unlikely to be good analogs for adaptation potential to current rates of OA, which can occur over the lifetime of an individual coral (100+ years). Therefore, despite the potential for acclimation indicated by such studies of extreme conditions, under the real-world environmental change and multivariate stressors, Galápagos Porites spp. corals have not demonstrated an ability to adapt to changing pH via pH<sub>cf</sub> upregulation.

Our synthesis of modern and fossil corals living under contrasting seawater conditions suggests that there may be a physiological limit to the capacity of corals to upregulate pH<sub>cf</sub> in response to changing ocean conditions and fluctuations in DIC<sub>cf</sub>. The capacity of corals to upregulate Ω<sub>cf</sub> is therefore likely to be dictated (to the first order) by their capacity to upregulate DIC<sub>cf</sub> via metabolic processes (e.g., GBR corals, particularly in the summer months), which we show is reduced both at marginal sites and following bleaching. Galápagos corals, which have low DIC<sub>cf</sub> despite high regional DIC<sub>sw</sub>, therefore require greater pH<sub>cf</sub> upregulation than modern GBR Porites spp. corals to maintain similar rates of calcification; the limited capacity to upregulate pH<sub>cf</sub> has therefore reduced Ω<sub>cf</sub> under modern conditions. Such a physiological limit, if it holds across future acidification (and across additional sites), is likely to leave corals in low-pH, high-DIC environments (i.e., in marginal environments) particularly susceptible to changing ocean saturation.
At both sites, the degree of pH\textsubscript{cf}, DIC\textsubscript{cf}, and Ω\textsubscript{cf} upregulation relative to seawater varied in concert with SST; warm seasons or years experience greater Ω\textsubscript{cf} and DIC\textsubscript{cf} upregulation, and weaker pH\textsubscript{cf} upregulation (Tables S2 and S5; Figures 4f–4h). These results agree with previous work showing a strong relationship between pH\textsubscript{cf} upregulation and temperature across a latitudinal gradient (Ross et al., 2019). Physicochemical modeling of coral cf chemistry suggests the temperature dependence of pH upregulation is driven primarily by calcification kinetics, and secondarily by seawater buffering capacity (i.e., the sensitivity of the pH\textsubscript{cf} to changes in TA; Guo, 2019). This dependence is particularly apparent during the 1997/98 El Niño in Wolf modern corals, with anomalously high pH\textsubscript{cf} and high Ω\textsubscript{cf} relative to seawater during and immediately following peak warming (January 1998 to September 1998), potentially due to increased buffering capacity at higher temperatures. However, the increase in pH\textsubscript{cf} upregulation following peak warming (i.e., during the stress recovery period) implies that other physiological mechanisms must also be at play, such as a change in the refresh rate of the cf or a change in the balance of bicarbonate and carbonate that is transported to the site of calcification (D’Olivo & McCulloch, 2017). Although uncertainties in the fidelity of the Sr/Ca-SST proxy across this thermal stress event may add uncertainty to the SST signal (D’Olivo & McCulloch, 2017), only ~2% of the pH\textsubscript{cf} anomaly can be explained by SST alone, and the Ω upregulation anomaly (i.e., 97/98 ∆Ω relative to the colony mean ∆Ω, Figure 4h) is robust between the replicate modern cores (23% and 31%) despite differences in calcification rate between colonies. Nevertheless, similar Ω upregulation anomalies do not preclude differences in the relative roles of DIC\textsubscript{cf} and pH\textsubscript{cf} in this saturation change (Figure 4). Our results suggest that although the response of metabolic carbon production and/or pH\textsubscript{cf} to thermal stress varies from colony to colony, the relative change in Ω\textsubscript{cf} with respect to seawater does not vary significantly among colonies. Again, these results demonstrate strong physiological limits to the corals’ ability to regulate their internal carbonate chemistry, and that this limit is likely an emergent property resulting from the interplay of numerous physiological processes or pathways.

3.6. Implications for Calcification Under Warming & Acidification

Our results demonstrate that physiological limitations have already had a pronounced impact on the geochemistry of the calcifying fluid in Galápagos Porites sp. corals. The pH\textsubscript{cf} declined significantly between eighteenth century and modern Wolf corals (Z = 24.3, N = 108,277, p < 0.001), and from 1975 to 2010 in the long modern Wolf (GW10-10) record (with a trend of ~0.18 pH units per decade). Over 99.9% of this recent trend (between 1975 and 2010) can be attributed to pH\textsubscript{sw}, with warming contributing less than 0.3% (after D’Olivo et al., 2019, see methods). The mean pH\textsubscript{cf} was 8.57 in two eighteenth century fossil cores from one colony (N = 78) and 8.50 in the two modern corals (N = 203, Figure 4, Table S2 in Supporting Information S1). This pre-industrial to modern mean pH\textsubscript{cf} difference can be attributed some combination of pH\textsubscript{sw} or SST changes. A large model ensemble of simulated changes between these periods suggests that either pH\textsubscript{sw} or SST could produce pH\textsubscript{cf} changes of 0.06–0.07 (see methods). In contrast, the temporal change in DIC\textsubscript{cf} differs between cores, consistent with a varying role of photosynthesis (and thus metabolic carbon) among (and even within) colonies. The combined impact on cf saturation state was profound, with a significant decline of ~2.3 units between the eighteenth century and late-twentieth century corals (Z = 24.2, N\textsubscript{fossil} = 108, N\textsubscript{modern} = 277, p < 0.001). These results emphasize the importance of extending the existing boron reconstructions across time periods that experienced different seawater chemistry from today. This initial study focused on replicate cores from one colony, and it will be critical to further replicate and extend these analyses to other fossil colonies to confirm these findings (given the potential for within and among colony differences in boron geochemistry, e.g., Chalk et al., 2021). Nevertheless, the first such application of boron systematics to pre-industrial fossil coral samples, presented here, paints a potentially stark future under projected acidification, with limited adaptive capacity in the upregulation of the coral calcifying fluid.

Despite this reduction in pH\textsubscript{cf} between the eighteenth and twentieth century Galápagos corals, there was no significant change in calcification or skeletal density among cores (or between modern and fossil colonies; see section “Coral densitometry and calcification” for description of methods). This is in contrast to previous work that demonstrates a strong relationship between calcification and pH\textsubscript{cf} (Guillermic et al., 2020; Ross et al., 2019). Rather, we find large interannual changes in calcification rate within (15%–27%) and among (24%–27%) cores (Table S6 and Figure S7 in Supporting Information S1). The predicted change in calcification between the eighteenth and twentieth centuries (of ~10%), using simulated Ω\textsubscript{sw} from Figure 1c, the Ω\textsubscript{cf} Pchange from Table S4 in Supporting Information S1 and the model of M. McCulloch et al. (2012), therefore falls within the range of interannual calcification variability at this site. Thus, despite large declines in Ω\textsubscript{cf}, the impact on coral calcification is not yet detectable at Wolf Island, Galápagos given the high interannual calcification variability.
However, these results should not be interpreted as evidence that Galápagos corals are robust to changing ocean chemistry, for five reasons. First, monthly skeletal density data is strongly related to both CF saturation state and temperature in both fossil and modern Galápagos corals (Figure 4). Although the nature of these relationships vary across cores (see Table S7 in Supporting Information S1; e.g., as a function of colony-to-colony variations in bleaching susceptibility), the relationships indicate declining density with warming and lower CF saturation (except in core WLF-3) and an increasing importance of warming in recent decades (becoming the dominant predictor in core WLF-10a, ending in 2010). Second, the corals studied here are likely to represent the “best-case-scenario,” as these long-lived corals targeted for paleoclimate reconstructions are the “winners” that were able to maintain rapid upward extension and calcification despite thermal stress (1997/98) and acidification (Figure S7 in Supporting Information S1). In smaller P. lobata colonies at nearby Darwin Island (Manzello et al., 2014), calcification rates were less than half those measured in our longer Wolf cores, despite similar density values among colonies from both sites (Table S6 in Supporting Information S1). Further, the modern Wolf colonies regrew in 3.4 (WLF10-10) and 5 (WLF10-03) years following the very strong 1982/83 El Niño event that devastated reefs across the Galápagos (Glynn et al., 1988), suggesting they experienced only partial mortality during this extreme event. Both colonies also displayed only modest reductions in extension and calcification during or following the 1997/98 event (Figure S7 in Supporting Information S1). Because paleoclimate records are biased towards corals that survive, they likely yield a conservative (i.e., too-stable) estimates of past calcification changes. Third, observed and simulated ocean pH at Galápagos remained above 8.0 over this period (mean CESM1 = 8.08–8.11 over intervals of coral coverage; Darwin = 8.07, Humphreys et al., 2018), a critical tipping point below which corals across the archipelago suffer reduced calcification and structural persistence (Manzello et al., 2014). High nutrients (Manzello et al., 2014) and variable seawater conditions exacerbate the stressful impacts of acidification in upwelling regions, resulting in tipping points at higher pH values (Manzello et al., 2014). Fourth, the temperature dependence of calcification kinetics does not appear to compensate for the impacts of saturation-state changes at Wolf (unlike in more optimal environments; Burton & Walter, 1987; Lough & Barnes, 2000). Lastly, and critically, we demonstrate that as oceans acidify, Wolf corals have not intensified their upregulation of pH or Ω, suggesting that continued OA is likely to have significant impacts on calcification at this site.

Finally, our results support the potential to reconstruct changes in paleo-pH from the geochemistry of coral calcifying fluid. Consistent with recent studies (Guo, 2019; D’Olivo et al., 2019), the narrow range in pH$_w$ upregulation of Porites spp. across sites and time periods (Table S4 in Supporting Information S1) suggests that within this paleo-relevant genus, long-term pH$_w$ trends are primarily driven by pH$_{cf}$ and not physiological controls (which regulate calcifying fluid chemistry on seasonal timescales, in response to temperature-related changes in DIC, calcification, and buffering capacity). Physiological limits in this capacity to regulate pH$_w$—identified here for the first time—suggest that as seawater saturation shifts to lower values (as observed with OA, or across spatial gradients, Manzello et al., 2014), so will the distribution of carbonate saturation in the calcifying fluid (as observed between eighteenth and twentieth corals). Corals’ capacity to buffer against OA may therefore be more limited than predicted from experimental manipulations and extreme environments (CO$_2$ seeps), with particularly severe consequences for corals at marginal sites characterized by reduced metabolic carbon production, low seawater pH, and frequent or severe thermal stress.

4. Summary

In presenting the first analysis of calcifying fluid geochemistry in pre-industrial and modern corals from a marginal environment, we are able to reconcile two seemingly competing truths about the capacity for corals to buffer against changing environmental conditions. On the one hand, we provide further evidence that corals are able to strongly upregulate the pH of their internal growth medium to maintain supersaturation in response to seasonal changes in DIC and temperature. This physiologically driven seasonal upregulation of pH$_w$ precludes the use of boron isotope geochemistry for reconstructing short-term variations in paleo-pH$_w$, but suggests that corals may be able to buffer against changing ocean conditions and maintain calcification under future warming and OA. On the other hand, recent work suggests that long-term trends in pH$_w$ inferred from boron isotope geochemistry are driven primarily by pH$_{cf}$ (D’Olivo et al., 2019), suggesting at least some sensitivity to environmental conditions. However, the resource-intensive nature of boron isotope geochemistry has limited the production of long reconstructions with which to assess corals’ buffering capacity under changing ocean conditions and therefore corals’ resilience to future warming and acidification.
Using cores from a pre-industrial fossil and two modern coral colonies from the Galápagos Islands—a marginal environment characterized by high environmental variability, low seawater pH, and frequent thermal stress—we identify significant declines in pH and Ω with warming and OA since the pre-industrial period. These trends are exacerbated during and after thermal stress events observed in the modern corals, likely due to the impact of bleaching on metabolic DIC production and the energy-intensive active transport that concentrates alkalinity against the electrochemical gradients. Critically, we demonstrate that these changes may be attributed to a remarkably narrow range of pH upregulation across sites and time periods, suggesting a strict physiological limit in corals’ ability to regulate their internal carbonate chemistry. We therefore find that the capacity of corals to maintain stable Ω supersaturation is dictated (to the first order) by their capacity to upregulate DIC via metabolic processes, which is reduced both at marginal sites and following bleaching. Such physiological limits in this capacity to regulate pH—identified here for the first time—suggest that corals’ capacity to buffer against OA may be more limited than predicted from experimental manipulations and extreme environments (e.g., CO2 seeps). These findings have particularly severe consequences for coral calcification and thus reef structure and function at marginal sites.

Conflict of Interest

The authors declare no conflicts of interest relevant to this study.

Data Availability Statement

All geochemical data is publicly available on the National Center for Environmental Information (formerly the National Climatic Data Center) paleoclimatology database: https://www.nci.noaa.gov/access/paleo-search/study/35193.

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