REPORT

Multiple driving factors explain spatial and temporal variability in coral calcification rates on the Bermuda platform

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Received: 8 February 2014/Accepted: 11 July 2014/Published online: 22 July 2014 © The Author(s) 2014. This article is published with open access at Springerlink.com

Abstract Experimental studies have shown that coral calcification rates are dependent on light, nutrients, food availability, temperature, and seawater aragonite saturation (Ω_{arag}) , but the relative importance of each parameter in natural settings remains uncertain. In this study, we applied Calcein fluorescent dyes as time indicators within the skeleton of coral colonies (n = 3) of *Porites astreoides* and Diploria strigosa at three study sites distributed across the northern Bermuda coral reef platform. We evaluated the correlation between seasonal average growth rates based on coral density and extension rates with average temperature, light, and seawater Ω_{arag} in an effort to decipher the relative importance of each parameter. The results show significant seasonal differences among coral calcification rates ranging from summer maximums of 243 ± 58 and $274 \pm 57 \text{ mmol CaCO}_3 \text{ m}^{-2} \text{ d}^{-1}$ to winter minimums of 135 ± 39 and 101 ± 34 mmol CaCO₃ m⁻² d⁻¹ for *P*. astreoides and D. strigosa, respectively. We also placed small coral colonies (n = 10) in transparent chambers and measured the instantaneous rate of calcification under light and dark treatments at the same study sites. The results

Communicated by Biology Editor Dr. Simon Davy

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showed that the skeletal growth of D. strigosa and P. astreoides, whether hourly or seasonal, was highly sensitive to Ω_{arag} . We believe this high sensitivity, however, is misleading, due to covariance between light and $\Omega_{\rm arag}$, with the former being the strongest driver of calcification variability. For the seasonal data, we assessed the impact that the observed seasonal differences in temperature (4.0 °C), light (5.1 mol photons m⁻² d⁻¹), and Ω_{arag} (0.16 units) would have on coral growth rates based on established relationships derived from laboratory studies and found that they could account for approximately 44, 52, and 5 %, respectively, of the observed seasonal change of $81 \pm 14 \text{ mmol } \text{CaCO}_3 \text{ m}^{-2} \text{ d}^{-1}$. Using short-term light and dark incubations, we show how the covariance of light and $\Omega_{\rm arag}$ can lead to the false conclusion that calcification is more sensitive to Ω_{arag} than it really is.

Keywords Coral calcification · Ocean acidification · Bermuda · Aragonite saturation state

Introduction

The negative effects of ocean acidification (OA) on coral calcification rates and net ecosystem calcification (NEC) rates have been well documented with a strong correlation observed between seawater aragonite saturation state (Ω_{arag}) and rates of calcification (e.g., Langdon and Atkinson 2005; Hoegh-Guldberg et al. 2007; Andersson and Mackenzie 2011). These results, however, are heavily drawn from controlled aquarium and mesocosm studies, which isolate OA by manipulating seawater inorganic carbon chemistry while keeping other parameters such as light, temperature, salinity, nutrients, and flow rates constant. Previous research has shown that coral calcification

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rates are also a direct function of these parameters, as well as the seawater inorganic carbon chemistry (e.g., Chalker and Taylor 1975; Koop et al. 2001; Mass et al. 2010). In situ studies, which in general do not control these parameters, are limited in their ability to evaluate the relative importance of the controlling factors on the rate of calcification, but can be helpful in assessing the combined impact of these parameters.

Seasonal differences in light, temperature, and seawater inorganic carbon chemistry, including seawater pH and $\Omega_{\rm arag}$, are more pronounced in higher latitude reefs, such as the Bermuda coral reef. For example, seasonal differences in $\Omega_{\rm arag}$ of up to 1.2 units have been recorded on the northern reef platform of Bermuda (Bates et al. 2010), while lower latitude reefs such as Davies Reef (Great Barrier Reef, GBR) exhibit smaller seasonal ranges of 0.15 (Albright et al. 2013). On diurnal timescales, the same is true for shallow reef systems. For example, large diurnal fluctuations in Ω_{arag} , from 1.5 to 6 units, have been observed on Lady Eliot Island (GBR) where reef waters are typically less than 1 m deep (Shaw et al. 2012). These types of environments provide natural laboratories to investigate rates of calcification under a broad range of seawater Ω_{arag} .

Though field data are limited, correlation between calcification rates and temporal variability in seawater temperature, light, and Ω_{arag} have been observed on coral reef ecosystems at diurnal (e.g., Suzuki et al. 1995; Yates and Halley 2003, 2006; Price et al. 2012) and seasonal (e.g., Silverman et al. 2007; Manzello et al. 2008; Bates et al. 2010; Shamberger et al. 2011; Albright et al. 2013) timescales. However, the interdependence of these parameters makes deciphering the relative importance and the principle driving mechanism(s) difficult. For example, increased light levels cause warmer temperatures, which also directly affect Ω_{arag} (higher temperature increases Ω_{arag} by decreasing aragonite solubility). Some studies have found a significant relationship between coral calcification and temperature (e.g., Marshall and Clode 2004; Reynaud et al. 2004; Silverman et al. 2007), while others have not (e.g., Shaw et al. 2012). To some extent, a positive relationship between temperature and calcification rates may be due to the effect of temperature on seawater Ω_{arag} (Silverman et al. 2007; Shaw et al. 2012), but this effect is relatively small (0.03 units per 2 °C).

Conversely, the concept of light-enhanced calcification has been well established (Kawaguti and Sakumoto 1948; Gordeau 1959; Goreau and Goreau 1959) and can increase coral calcification rates by three to ten times compared with rates in the dark (Gattuso et al. 1999; Moya et al. 2006). Light availability may also determine the susceptibility of reef building corals to pressures from OA. Increased gross photosynthesis under high light conditions has been shown to effectively enhance calcification in *Aiphanes horrida* and *Parapercis cylindrica*, even under high CO₂ treatments (Suggett et al. 2012). At seasonal timescales, the net reef community metabolism may also affect the reef's susceptibility to pressures from OA. Strong net autotrophy from benthic components of the reef system may enhance Ω_{arag} due to photosynthetic uptake of CO₂, while Ω_{arag} may be suppressed by the release of CO₂ from reef metabolism during periods of strong net heterotrophy, as proposed by the coral reef ecosystem feedback (CREF) hypothesis (Bates et al. 2010).

The present study measured the calcification rates of two scleractinian coral species (Diploria strigosa and Porites astreoides) at three sites across the Bermuda coral reef platform at seasonal (winter/summer) and hourly (light/ dark) timescales to better understand how coral calcification rates vary across natural temporal and spatial differences in light, temperature, and seawater Ω_{arag} . Seasonal Calcein staining was conducted in March 2010, September 2010, and March 2011, and colonies were cored in September 2011, while hourly incubations were conducted in September 2010 and 2011. Though natural temporal and spatial differences in nutrients are also likely, these data were beyond the scope of the present study. We compare seasonal coral calcification rates from Bermuda with calcification rates measured by similar methods in other reef environments and discuss the relative importance of seawater carbonate chemistry (specifically Ω_{arag}) on coral calcification rates at hourly and seasonal timescales.

Materials and methods

Study site

Bermuda offers a unique environment to study the effects of OA on coral calcification rates. Bermuda lies on the northern limit of tropical coral reefs ecosystems and thus experiences threshold oceanic environmental and chemical conditions with respect to temperature and light, and possibly $\Omega_{\rm arag}$ and pH, for supporting this ecosystem. It has been hypothesized that Bermuda is likely to be affected by OA sooner than lower latitude reefs, making Bermuda an important location to study the effects of seawater inorganic carbon chemistry on coral calcification rates (Kleypas et al. 1999, 2001; Bates et al. 2010). Bermuda is also relatively spared from major human influences, such as pollution, run-off, overfishing, and dredging common to many US and Caribbean reefs and has been protected relatively successfully by local legislation. Therefore, observed long-term changes in coral calcification rates are likely to reflect large-scale pressures from global climate change (GCC) and OA rather than local factors.

Rim reef waters are rapidly flushed by offshore water (1-2 d), while longer residence times are found closer to shore (10-15 d; Venti et al. 2012). Consequently, the potential for a spatial gradient in carbon chemistry across the Bermuda reef platform seems likely. However, Andersson et al. (2013) have shown that despite a strong seasonal and spatial signal in the distributions of total alkalinity (TA) and dissolved inorganic carbon (DIC), only relatively small spatial gradients in pH and Ω_{arag} are observed across the platform due to the nearly proportional changes in DIC and TA owing to net reef metabolism (i.e., net ecosystem production, NEP = primaryproduction - autotrophic and heterotrophic respiration; net ecosystem calcification, NEC = calcification -CaCO₃ dissolution; Andersson and Gledhill 2013). Nevertheless, in an effort to capture the maximum range of environmental and chemical conditions experienced across the platform, three spatially separated sites were selected for the present study: East of North Rocks (NR; 32.473247°N, 64.74572°W), Whalebone Bay (WB: 32.364516°N, 64.71573°W), and Buoy 29 (B29; 32.32203°N, 64.811718°W; Fig. 1).

Seasonal coral calcification rates

Coral calcification rates (G) can be defined as the product of the coral's vertical extension during a given time interval (X) and its skeletal density (ρ) (Chalker et al. 1985):

$$X \operatorname{cm} \operatorname{d}^{-1} \cdot \rho \operatorname{gCaCO}_3 \operatorname{cm}^{-3} = G \operatorname{gCaCO}_3 \operatorname{cm}^{-2} \operatorname{d}^{-1} \qquad (1)$$

In this study, we combined a calcein dye staining method, which has not been shown to negatively impact coral growth, to assess seasonal vertical extension with X-ray imaging analysis of coral core slices to quantify coral density and thus seasonal calcification rates.

Seasonal calcification rates were measured for a total of eighteen coral colonies, three *P. astreoides* and three *D. strigosa* at three study sites. The staining process consisted of loosely "tenting" tagged coral colonies with an impermeable plastic tarp secured to the base of the coral with bungee cords. Diluted calcein dye (1 g L⁻¹ of sea water) was released under the tarp, which remained over the coral for 18–24 h before being removed. Despite longer than ideal tenting events, coral colonies did not exhibit any

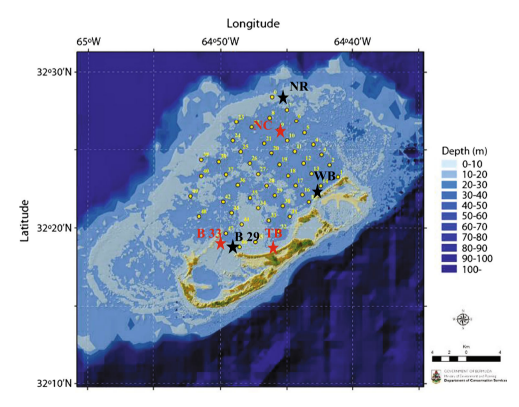


Fig. 1 Bermuda and the Bermuda reef platform. *Black stars* indicate three sites selected for the present study, North Rocks (NR), Whalebone Bay (WB), and Buoy 29 (B29). *Yellow dots* denote sampling stations for the surface water platform survey conducted in March (winter) and September (summer) 2012. *Red stars* indicate the

location of monthly environmental data collected between March 2010 and September 2011 by the BEACON project: Buoy 33 (B33), North Channel (NC), and Tyne's Bay (TB). Map credit: M. Shailer, Dept. of Conservation Services, Government of Bermuda

signs of stress (e.g., bleaching, paling) from being tented. Colonies were stained during seasonal sampling expeditions in March 2010, September 2011, and March 2011. In September 2011, stained corals were cored on SCUBA using a pneumatic hand drill attached with a diamond coring bit, capturing three complete growth periods: two periods from March to September, encompassing mostly "summer" months, and one period from September to March, designated as "winter" months. The distance between each seasonal calcein band, revealed by a fluorescent filter, represents the coral's vertical extension between sampling periods. Coral density was measured on coral cores sliced to a uniform thickness of 0.5 cm using X-ray imaging analysis and Coral XDS software (Chalker et al. 1985; Helmle et al. 2002; http://www.nova.edu/ ocean/coralxds/index.html).

Light/dark coral calcification rates

Though well suited for seasonal timescales, the calcein staining method is not applicable for diurnal resolution. Instead, we conducted short-term incubation experiments using the alkalinity depletion method to determine hourly calcification rates in light and dark conditions. This method is based on the principle that seawater TA is depleted in a mol ratio of 2:1 for every mol $CaCO_3$ deposited by the coral during the incubation period. This relationship is defined in Schneider and Erez (2006) as:

$$G(\mu \text{mol CaCO}_3 \text{cm}^{-2}\text{h}^{-1}) = \frac{(\Delta \text{TA}/2) \cdot (V_{\text{chamber}} - V_{\text{coral}}) \cdot \rho}{t \cdot \text{SA}}$$
(2)

where $\Delta TA \ (\mu mol \ kg^{-1})$ is the change in total alkalinity during incubation, $V_{chamber}$ and $V_{coral} \ (m^3)$ are the volumes of the chamber and displaced by the coral, respectively, ρ (kg m⁻²) is the density of sea water, t (h) is the duration of the incubation, and SA (m²) is the surface area of the coral nubbin.

In March 2010, ten coral "nubbins," five of each *P. astreoides* and *D. strigosa*, were collected at each site, epoxied onto PVC sleds and secured onto a platform at the site from which the corals were collected. The initial surface area of all corals was measured with calipers assuming simple geometric forms unique for each colony (i.e., half sphere, cone). In general, *D. strigosa* were classified as half sphere geometries, while *P. astreoides* were typically cone-shaped, though each colony's geometry was assessed independently for surface area analysis. In September 2010, after corals had a chance to recover from the collection process, and again in September 2011, light and dark calcification rates were measured in situ by collecting water samples from 2-1 incubation chambers after ~ 1.5 -h

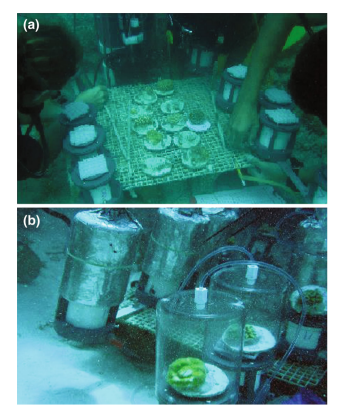


Fig. 2 Platforms used for a hourly incubation study and b hourly incubation analysis. Chambers in the background with the tin-foil represent dark incubations, and chambers in the foreground represent light incubations

incubation. All chambers were fitted with a battery-operated magnetic stirring bar to ensure sufficient water circulation within the chamber during the incubation process (Fig. 2). Dark incubations were achieved by covering chambers with aluminum foil. Water samples were collected from the water column above the platform at the start of incubations to quantify the initial conditions and also at the end from each chamber by syringe from a sampling hose secured with a clamp. Water samples from blank incubation chambers were used to correct for any net water column metabolism (i.e., to understand the change in TA over time in the water column). Incubations were typically conducted around noon, though exact starting times varied between stations and sampling year. Dark incubations were conducted first, after which aluminum foil was removed and chambers were flushed with fresh water before starting the light incubations, roughly 2.5 h after the dark incubation.

Platform surveys

To assess the spatial variability in seawater inorganic carbon chemistry across the Bermuda coral reef platform within the scope of this study, we conducted two platformwide surveys in March and September 2012 (Fig. 1). At each of 50 stations, surface seawater measurements of temperature, salinity, and dissolved oxygen were taken with a yellow springs instrument (YSI) analyzer and water samples were collected for TA and DIC analysis.

Sample analyses

Seawater samples for DIC and TA from the spatial surveys were collected in thrice rinsed 250-ml PETG bottles. Samples from the in situ chamber experiments were drawn into four 50-cc plastic syringes by divers and transferred through 0.45 µm cellulose acetate filters to 250-ml PETG bottles at the surface. A headspace of $\sim 1-2$ % of the sample volume was created to allow for water expansion, and 100 µl of saturated HgCl₂ was added to stop biological activity. All samples were analyzed within six weeks of collection. Dissolved inorganic carbon was measured by an acidification/gas extraction technique using an Apollo Scitech AS-C3 Total Dissolved Inorganic Carbon Analyzer. In this technique, 0.75 ml sample maintained at 25 °C is drawn up into a syringe pump, mixed with 1 ml of 10 % H₃PO₄ acid and injected into a gas stripping cell where UHP N_2 gas carries the CO₂ gas through a LICOR 7000 CO₂ Gas Analyzer. The LICOR integrates the absorbance versus time signal and yields a net result that is proportional to the total inorganic carbon content of the sample. Seawater-certified reference material (CRM's, prepared by A. Dickson, Scripps Institution of Oceanography) was analyzed at the beginning and end of each 15-20 sample run to ensure accuracy of the DIC measurements. Precision based on analysis of replicate offshore samples was 0.17 % (~3 μ mol kg⁻¹, n = 17).

Total alkalinity was determined by potentiometric titration with 0.1 M HCl. The second equivalence point was determined by iteratively fitting a modified Gran function (Hansson and Janger 1973) to the titration data using code originally written by D. Archer, University of Chicago. Certified reference materials were analyzed at the beginning and end of each set of 15 samples to ensure the accuracy of TA measurements. Precision based on analysis of replicate offshore samples was 0.19 % (~4 μ mol kg⁻¹, n = 17). All CRMs, DIC, and TA samples were run in duplicate. Total alkalinity and DIC samples were filtered through 0.45 µm cellulose acetate filters before analysis to remove any carbonate particulates that might be present in the sample. The pCO₂, pH (total scale), and Ω_{arag} were calculated as a function of the measured salinity, temperature, TA, and DIC using the program CO2SYS, dissociation constants for carbonate acid determined by Mehrbach et al. (1973) as refit by Dickson and Millero (1987), and the dissociation constant for boric acid determined by Dickson (1990).

Complementing environmental data

Though platform survey data were valuable in assessing spatial trends in carbon chemistry, it was limited in its temporal resolution. To assess the relationship between coral calcification over the 6-month growth periods of our three season study, we utilized monthly salinity, temperature, TA, and DIC data collected by BIOS' water quality monitoring program (WQMP) and the Bermuda Ocean Acidification and Coral Reef Investigation (BEACON) project from three sampling stations distributed across the Bermuda coral reef platform (Fig. 1). Though sampling sites were not exactly at the same location as our three study sites, similar environmental characteristics between north channel (NC) and NR, Tyne's Bay (TB) and WB, and Buoy 33 (B33) and B29, motivated the analysis of environmental data from the former sites with coral growth data from the later. What we lost in spatial resolution (i.e., these sampling sites were not exactly the same location as our three study sites) was compensated for by higher temporal resolution, as platform survey data from the present study was limited to 1 d during summer and winter. Light data were collected at the L. F. Wade Bermuda International Airport in St. George by BAS-Serco and the Bermuda Weather Service in 1-min intervals that were integrated over 24 h to yield daily PAR values for each day of the three 6-month sampling periods.

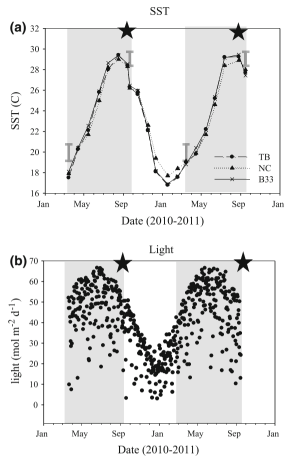
Results

Corals were stained at our three study sites, (NR, WB, and B29) in March 2010, September 2010, and March 2011, and cored in September 2011, marking three 6-month growth periods. The environmental characteristics of each of these growth periods, collected by the BEACON project, are reported below, followed by the corresponding growth data.

Environmental conditions

March 2010–Sept 2010 (summer I)

Monthly measurements from NC, TB, and B33 showed that surface water temperatures ranged from a minimum of 17.5 °C in late March to a maximum of 29.5 °C in late August, with an average temperature of 24.5 °C ($\sigma = 4.3$, n = 21) for the first 6-month growth period (Fig. 3). Light levels ranged from 7.4 mol m⁻² d⁻¹ in late March to 66.5 mol m⁻² d⁻¹ in June, with an average of 47.3 mol m⁻² d⁻¹ ($\sigma = 12$, n = 280; Fig. 3). The Ω_{arag} reached a minimum in April, with values of 3.35 and 3.36 for TB and B33, respectively (Fig. 3). At NC, Ω_{arag}



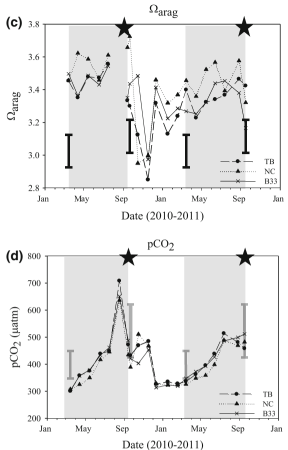


Fig. 3 Environmental data during the seasonal coral staining study. a SST, c Ω_{arag} and d pCO₂ data came from monthly sampling by the BEACON project. b Light data came from the BAS-Serco and the Bermuda Weather Service. *Light gray bars* represent the two

reached a minimum of 3.57 in both March and mid-June. The maximum Ω_{arag} was observed at all three study sites in mid-July with values of 3.56, 3.61, and 3.54, for TB, NC, and B33, respectively. The average Ω_{arag} during this period was 3.46 ($\sigma = 0.07$, n = 5), 3.46 ($\sigma = 0.07$, n = 5), and 3.55 ($\sigma = 0.08$, n = 5), at TB, B33, and NC, respectively. Seawater pCO₂, though not discussed in this paper, agrees with the Ω_{arag} results and is also shown in Fig. 3 for context. Though there were no significant differences in SST among the three study sites (1-way ANOVA, $F_{(1,12)} = 0.009$, p = 0.99), there was a significant difference in the Ω_{arag} among the three study sites (1-way ANOVA, $F_{(1,12)} = 4.43$, p = 0.03).

September 2010-March 2011 (winter)

Minimum temperatures reached 16.8 °C in January and a maximum temperature of 26.3 °C in late September, with an average of 20.0 °C ($\sigma = 4$, n = 21) for the second 6-month growth period (Fig. 3). Though daily light values

"summer" periods; the *white bar* represents the "winter" sampling period. *Stars* indicate time of hourly incubation study. *Dark gray bars* represent the range of the parameter observed during the platform surveys in March and September 2012

varied from a minimum of 3 mol m⁻² d⁻¹ to a maximum of 50.7 mol m⁻² d⁻¹ (Fig. 3), they were typically low, averaging 27.7 mol m⁻² d⁻¹ ($\sigma = 11.4$, n = 183). The Ω_{arag} was low with an average of 3.28 ($\sigma = 0.25$, n = 6) at NC, 3.27 ($\sigma = 0.048$, n = 6) at B33, and 3.22 ($\sigma = 0.048$, n = 6) at TB (Table 1). Spatial differences were minimal with SST and Ω_{arag} , showing no significant differences among the three study sites (1-way ANOVA, $F_{(1,12)} = 0.03$, p = 0.99 for SST; 1-way ANOVA, $F_{(1,12)} = 0.73$, p = 0.49 for Ω_{arag}).

March 2011-September 2011 (summer II)

Sea surface temperature ranged from 18.8 °C in March to a maximum of 29.5 °C in late August, with an average SST of 24.6 °C ($\sigma = 4.1, n = 6$). Light levels ranged from 10.3 to 66.5 mmol m⁻² d⁻¹ with an average light level of 47.8 mmol m⁻² d⁻¹ ($\sigma = 12.7, n = 281$). The Ω_{arag} displayed different trends among study sites during the third 6-month growth period. At B33, the Ω_{arag} reached a

Station (coral/ environmental)	Coral	Total extension $(mm \ yr^{-1})$	Average annual density G (g (g cm ⁻³) CaCC	G (g CaCO ₃ cm ⁻² yr ⁻¹)	Season	G (mmol CaCO ₃ m ⁻² d ⁻¹)	$\Omega_{ m arag}$	(C) ISS	$I ({ m mol} { m mol} { m m}^{-2}{ m d}^{-1})$
NR/NC	D. strigosa	54.4	1.191	6.48 (土0.23)	Summer	253 (土22)	3.53 (土0.2)	25.4 (±3.43)	48.26 (土12.06)
NR/NC	D. strigosa				Winter	144 (土30)	3.28 (土0.25)	20.48 (±3.29)	27.72 (土11.44)
B29/B33	D. strigosa	16.9	1.499	2.5 (土0.45)	Summer	187 (土41)	3.39 (土0.04)	25.83 (土3.4)	48.26 (土12.06)
B29/B33	P. astreoides	54.2	1.475	7.99 (土0.8)	Summer	222 (土77)	3.39 (土0.04)	25.83 (土3.4)	48.26 (土12.06)
B29/B33	P. astreoides				Winter	135 (土39)	3.27 (土0.05)	19.24 (土3.52)	27.72 (土11.44)
12.06	P. astreoides	45.5	1.167	5.31 (土0.57)	Summer	210 (土27)	3.39 (土0.04)	25.66 (土3.48)	48.26 (土12.06)
11.44	P. astreoides				Winter	145 (土13)	3.22 (土0.05)	19.79 (土3.56)	27.72 (土11.44)
12.06	D. strigosa	41.6	1.317	5.48 (土0.63)	Summer	203 (土57)	3.39 (土0.04)	25.66 (土3.48)	48.26 (土12.06)
11.44	D. strigosa				Winter	111 (±13)	3.22 (土0.05)	19.79 (土3.56)	27.72 (土11.44)

minimum of 3.17 in September and a maximum of 3.45 in mid-July, with an average of 3.31 ($\sigma = 0.13$, n = 7). At TB, the Ω_{arag} reached a minimum of 3.23 in mid-April and a maximum of 3.46 in August. The average Ω_{arag} at TB was 3.33 ($\sigma = 0.11$, n = 7). The Ω_{arag} at NC reached a minimum in September of 3.32, though the Ω_{arag} in April and July was also low (3.36 and 3.39, respectively). The maximum Ω_{arag} was observed in both June, 3.57 and again in August, 3.58 at NC. The average Ω_{arag} at NC during this period was 3.40 ($\sigma = 0.19$, n = 7). We found that the two summer (March–September)

growth periods were not significantly different in terms of SST (ANOVA, $F_{(1,12)} = 1.72$, p = 0.14), Ω_{arag} (ANOVA, $F_{(1,12)} = 0.40$, p = 0.126), and light levels (ANOVA, $F_{(1,361)} = 0.39, p = 0.46$). This allowed us to pool environmental data from the two summer (March-September) sampling periods. The average SST, Ω_{arag} , and light based on the pooled data for the two summer (March-September) periods were significantly greater than during the winter (September–March) period: 25.0 versus 21.0 °C $(F_{(1,58)} = 14.7, p < 0.0003),$ 3.43 versus 3.27 $(F_{(1.55)} = 14.0, p < 0.0004)$, and 47.9 versus 27.6 mol quanta $m^{-2} d^{-1}$ ($F_{(544)} = 348$, p < 0.0001), respectively. We compared these differences in seasonal environmental data with results from our bi-seasonal coral growth experiment to assess the impact of these parameters on seasonal coral calcification rates in Bermuda.

Seasonal calcification rates

Calcification rates for *D. strigosa* were significantly higher during the summer (March–September) than the winter (September–March) period (218 ± 10 vs. 128 ± 14 mmol CaCO₃ m⁻² d⁻¹; 1-tailed *t* test, $t_{(11)} = 5.3$, p < 0.0001). *P. astreoides* also calcified faster during the summer (March– September) than in the winter (September–March) period, but the difference was not significantly different (267 ± 45 vs. 142 ± 64 mmol CaCO₃ m⁻² d⁻¹; 1-tailed $t_{(7)} = 1.6$, p = 0.076; Table 2; Fig. 4).

The stained corals in this study experienced a wide range of environmental conditions during the 1.5-yr-long study (Fig. 3). The seasonal ranges in SST, light, and Ω_{arag} were fairly similar between the summer (March–September) and winter (September–March) periods. However, despite the broad overlap of conditions, the average SST, light, and Ω_{arag} during the two seasons were significantly different as were the seasonal coral growth rates (preceding section). A regression of seasonal coral calcification rates of the two coral species at the three locations with the seasonally averaged environmental conditions showed strong correlations with light ($r^2 = 0.8$), SST ($r^2 = 0.79$), and Ω_{arag} ($r^2 = 0.75$; Fig. 5). The fact that the correlation coefficients for each of the factors are similar (0.75–0.8)

Site	Site Sample	Seasonal Calce	Seasonal Calcein dye experiment	rent						CoralXDS		
		March 2010– Sent 2010	March 2010- Sept 2010- Sent 2010 March 2011	March 2011–Sept 2011	spt 2011	Seasonal calci	Seasonal calcification (g cm^{-2} yr^{-1})	² yr ⁻¹)	Average annual $G(\alpha \text{ cm}^{-2} \text{ vr}^{-1})$		Density	Extension Density Average annual $(mm vr^{-1})$ (σcm^{-3}) G ($\sigma cm^{-2} vr^{-1}$)
		Extension (mm yr^{-1})	Extension Extension (mm yr^{-1}) (mm yr^{-1})	Extension (mm yr^{-1})	Density (g cm ^{-3})	March 2010- Sept 2010	Sept 2010- March 20 March 2011 Sept 2011	March 2011- Sept 2011			(III. 9)	
NR	NR D. strig.	86.4 (土13) 38.83 (土6)	38.83 (土6)	86.45 (土11)	1.2367	10.7 (土2)	4.8 (土1)	10.7 (土3)	7.81 (土0.6)	64	1.2367	7.91
NR	NR P. astr.			105.66 (土9)	1.8958			20.0 (土2)			1.8958	0.00
WB	WB D. stri.	62.7 (土8)	33.8 (土7)	62.7 (土5)	1.2668	7.9 (土2)	4.3 (土1)	7.9 (土3)	6.84 (土0.2)	53	1.2668	6.71
WB	WB P. astr.	68.44 (土3)	38.62 (土8)	34.9 (土7)	1.4109	9.7 (±3)	5.4 (土0.5)	4.9 (土1)	6.85 (土0.3)	51	1.4109	7.20
B29	B29 D. strig.			47.77 (土6)	1.4754			7.0 (土2)	7.05 (土0.3)	50	1.4754	7.38
B29	B29 P. astr.	69.3 (土7)	38.4 (土5)	57.33 (土5)	1.3195	9.1 (±1)	5.1 (土1)	7.6 (土2)	7.26 (±0.9)	58	1.3195	7.65

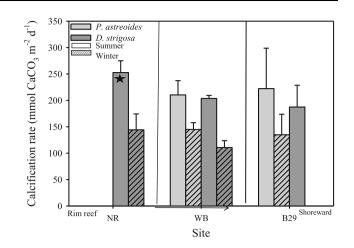


Fig. 4 Seasonal coral calcification rates averaged among three replicate samples of *Porites astreoides* (*light gray*) and *Diploria strigosa* (*dark gray*) measured from the seasonal calcein dye staining experiment. *Open bars* represent summertime rates (March–September); *hashed bars* represent winter rates (September–March). *Error bars* denote standard deviations among replicate species (n = 3). Study sites are shown as a function of their proximity to the rim reef, with NR closest to the rim reef, and progressing shoreward to WB and B29 along the *x*-axis. Calcification rates of *D. strigosa* at NR were statistically different from rates measured at WB or B29 (*star*). Calcein bands of *P. asreoides* at NR were not clear in all three replicate samples and thus are not shown. *D. strigosa* at B29 only displayed the final calcein sampling band (summer) in all three replicates, and thus, winter data are also not shown

does not mean that each factor contributed equally to the observed differences in calcification rate, but rather that the factors covary over time, i.e., temperature, light, and Ω_{arag} are highest during the summer and lowest during the winter. Increasing solar irradiance causes the water to warm. Warmer temperatures result in lower solubility of CO_2 in seawater and lower aragonite solubility, which contribute to an increase in seawater Ω_{arag} . Increasing solar irradiance also affects net community production and calcification, which could alter Ω_{arag} depending on the relative uptake of DIC and TA (Andersson and Gledhill 2013). We discuss the cause and effect between these changes and the calcification rates of the corals in the subsequent discussion.

Hourly light/dark calcification rates

Hourly incubation experiments were conducted just after maximum temperatures were reached at our three study sites, in September 2010 and 2011 (indicated by the stars in Fig. 3). Average hourly calcification rates for *D. strigosa* (n = 5) ranged from $-32 (\pm 20)$ mmol CaCO₃ m⁻² h⁻¹ in 2010 to 14 (± 6) mmol CaCO₃ m⁻² h⁻¹ in 2011 under dark treatments and light treatments ranged from 7 (± 6) mmol CaCO₃ m⁻² h⁻¹ in 2011 to 40 (± 26) mmol CaCO₃ m⁻² h⁻¹ in 2011 under light treatments. Average hourly

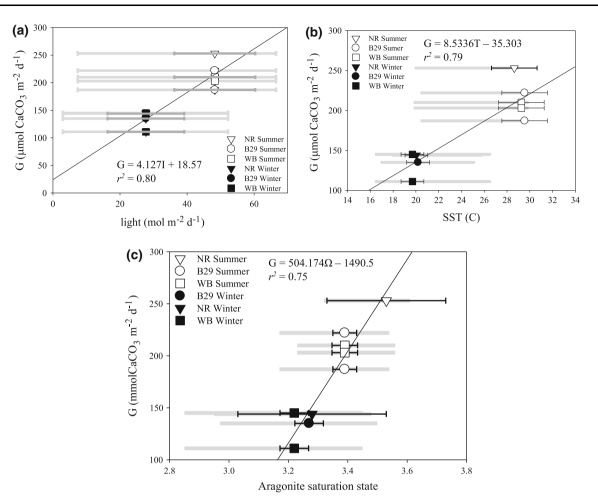


Fig. 5 Summer (*open*) and winter (*filled*) coral calcification rates averaged among replicate samples of *P. astreoides* and *D. strigosa* and corresponding **a** light, **b** sea surface temperature, and **c** aragonite saturation states at three study sites distributed across the northern Bermuda coral reef platform. *Gray bars* represent the observed seasonal range (Fig. 2); *black bars* represent average diurnal range;

calcification rates for P. astreoides (n = 5) ranged from $-34 (\pm 21) \text{ mmol CaCO}_3 \text{ m}^{-2} \text{ h}^{-1}$ in 2010 to 24 (±35) mmol $CaCO_3 m^{-2} h^{-1}$ in 2011 in dark treatments and from 14 (±10) mmol CaCO₃ m⁻² h⁻¹ in 2010 to 32 (±19) mmol CaCO₃ m⁻² h⁻¹ in 2011 in light treatments (Fig. 6). Average hourly calcification rates in light and dark treatments showed no significant difference between the three study sites for either species P. astreoides and D. strigosa (ANOVA, $F_{(2, 19)} = 0.06$, p > 0.05; Fig. 6). However, the calcification rates between light and dark treatments were significantly different (ANOVA, $F_{(1,19)} = 5.71$, p < 0.05; Fig. 6). Average seawater pH and Ω_{arag} were lower during dark treatments compared with light treatments in both 2010 and 2011 (Table 3). Light levels were similar during 2010 and 2011 sampling periods, while SST was about 2 °C cooler in 2011 (Table 3). Temperature and light did not change during the 1.5-h incubation period and were

dark gray bars in figure **a** represent the standard deviation of the seasonal average. Diurnal ranges in SST and Ω_{arag} were based on measurements from NOAA PMEL MAPCO₂ buoy systems at Hog Reef (32.46°N, 64.83°W) and Crescent Reef (32.40°N, 64.79°W) http://www.pmel.noaa.gov/co2/story/Coral+Reef+Moorings. Light data came from BAS-Serco and the Bermuda Weather Service

identical to water column values, but Ω_{arag} was altered by the coral's metabolism; thus, corals experienced a range of Ω_{arag} during the 1.5-h incubation. The values reported here are the average Ω_{arag} during the 1.5-h incubation; the complete range of Ω_{arag} each coral experienced during the 1.5-h incubation is shown in Fig. 7.

Sea surface temperatures, measured in the water column and not inside the chambers throughout the incubation, were constant throughout the 1.5-h incubation period and did not vary between light and dark incubations. Temperature was only weakly correlated with hourly calcification rates ($r^2 = 0.06$). Light levels also remained constant throughout the incubation period and were, like SST, weakly correlated with calcification rates ($r^2 = 0.25$). Hourly calcification rates displayed greater correlation with the Ω_{arag} ($r^2 = 0.53$), which did vary throughout the 1.5-h incubation (Fig. 7).

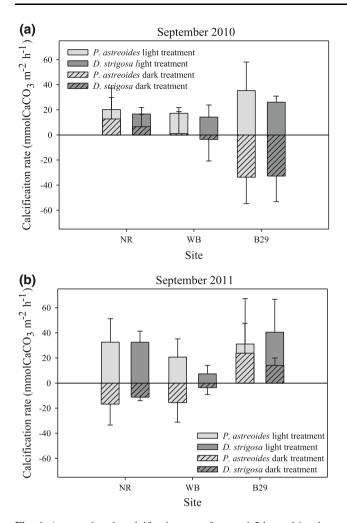


Fig. 6 Average hourly calcification rates from ~ 1.5 h coral incubations in **a** September 2010 and **b** September 2011 under light (*open*) and dark (*dashed*) treatments for the two sample species *Porites astreoides* (*light gray*) and *Diploria strigosa* (*dark gray*). Results are averaged from five coral nubbins of each *P. astreoides* and *D. strigosa* at each of the three study sites. *Error bars* denote standard deviations. Study sites are shown as a function of their proximity to the rim reef, with NR closest to the rim reef and moving further away progressing along the *x*-axis

Spatial surveys of carbonate chemistry

During the March 2012 platform spatial survey, distributions of DIC and TA were high and relatively uniform across the Bermuda coral reef platform, with no clear trends. The March seawater DIC ranged from 2,067 to 2,094 µmol kg⁻¹ (Δ DIC = 27 µmol kg⁻¹), and TA ranged from 2,346 to 2,361 µmol kg⁻¹ (Δ TA = 15 µmol kg⁻¹). The Ω_{arag} was relatively uniform across the platform with an average of 3.08 ($\sigma = 0.12$, n = 50). Seawater pCO₂ was also uniform with an average of 392 µatm ($\sigma = 24$, n = 50).

The September distributions of carbon chemistry from the 2012 platform showed a drawdown of both DIC and TA compared with offshore values, with a large shoreward decline in both parameters. Dissolved inorganic carbon ranged from 2,047 µmol kg⁻¹ at the rim reef to 1,938 µmol kg⁻¹ nearshore (Δ DIC = 109 µmol kg⁻¹), while TA ranged from 2,319 to 2,234 µmol kg⁻¹ (Δ TA = 85 µmol kg⁻¹; Table 1). The average Ω_{arag} was 3.29 ($\sigma = 0.24$, n = 50), and the average pCO₂ was 497.31 µatm ($\sigma = 37.29$, n = 50).

To evaluate spatial trends in these distributions, the 50 stations were divided into seven cross-platform transects extending from shore to the rim reef. Data from stations at similar distances from shore were averaged and evaluated in reference to the average distance from shore. The significance of this average trans-platform trend was determined by testing the null hypothesis that the slope equaled zero (H₀: $\beta = 0$). Based on this analysis, we observed no significant spatial trends in TA, DIC, or carbonate species pCO₂ or Ω_{arag} in the March sampling survey (Table 4).

Despite greater variability of both pCO₂ and Ω_{arag} in the September spatial survey, neither parameter displayed any statistically significant trend or gradient across the Bermuda coral reef platform (Table 4). When we averaged data from all rim reef stations and all nearshore stations, however, we found a significant difference of 0.14 units in the Ω_{arag} between these two regions of the platform (one sided *T* test, $t_{(16)} = 2.82$, p < 0.05). These results agree with the monthly data, which show elevated Ω_{arag} at the rim reef (NC) compared with nearshore sites (TB and B33).

Discussion

Evaluation of the calcein staining method

The objective of this study was to resolve seasonal changes in coral growth rates and evaluate these changes in the context of variations in environmental conditions. Historically, alizarin red has been the method of choice to stain corals because it leaves a clear, pink stain, visible to the eye. However, sometimes the corals do not take the stain up effectively and data may be lost. There has also been concern that alizarin may be toxic to some corals (Dodge et al. 1984; Holcomb et al. 2013). For these reasons, we decided to use calcein dye for this study. The drawback of this stain is that it is not visible to the naked eye; it fluoresces when excited by 475 nm light. The advantages are that corals seem to take it up more reliably, and there is no reported negative impact to corals.

Though experiments employing calcein dye to measure intra-annual extension are becoming increasingly common in laboratory experiments, application of this method to in situ experiments has remained relatively limited until recently (Tambutte et al. 2011; Holcomb et al. 2013). To

Table 3 Chemical and physical properties of water within the incubation chambers of the 2010 and 2011 hourly incubation study

Year	Site	Treatment	coral species	Temp C	Salinity	Light (μ mol photon m ⁻² s ⁻¹)	$TA \\ (\mu mol \ kg^{-1})$	DIC (µmol kg ⁻¹)	рН	$arOmega_{ m arag}$
2010	B29	Light	P. astreoides	27.78	36.86	486	2,233 (±20)	1,917 (±19)	7.97 (±0.01)	3.22 (±0.08)
	B29	Light	D. strigosa	27.78	36.86	486	2,272 (±8)	1,951 (±11)	7.97 (±0.01)	3.25 (±0.04)
	B29	Dark	P. astreoides	27.78	36.86	0	2,318 (±66)	2,061 (±63)	7.85 (±0.07)	2.7 (±0.37)
	B29	Dark	D. strigosa	27.78	36.86	0	2,333 (±30)	2,058 (±34)	7.88 (±0.02)	2.86 (±0.07)
	NR	Light	P. astreoides	27.08	36.832	243	2,304 (±27)	1,976 (±41)	7.99 (±0.03)	3.46 (±0.22)
	NR	Light	D. strigosa	27.08	36.832	243	2,273 (±34)	1,938 (±34)	7.98 (±0.02)	3.30 (±0.12)
	NR	Dark	P. astreoides	27.08	36.832	0	2,377 (±23)	2,107 (±43)	7.87 (±0.06)	2.84 (±0.28)
	NR	Dark	D. strigosa	27.08	36.832	0	2,371 (±30)	2,093 (±26)	7.87 (±0.02)	2.85 (±0.13)
	WB	Light	P. astreoides	27.69	36.767	87	2,283 (±16)	1,966 (±10)	7.98 (±0.01)	3.36 (±0.06)
	WB	Light	D. strigosa	27.69	36.767	87	2,272 (±24)	1,955 (±20)	7.98 (±0.02)	3.33 (±0.11)
	WB	Dark	P. astreoides	27.69	36.767	0	2,359 (±30)	2,075 (±45)	7.89 (±0.06)	2.95 (±0.32)
	WB	Dark	D. strigosa	27.69	36.767	0	2,338 (±45)	2,070 (±13)	7.88 (±0.05)	2.88 (±0.31)
2011	B29	Light	P. astreoides	25.56	36.273	441	2,266 (±31)	1,989 (±32)	7.96 (±0.06)	3.00 (±0.34)
	B29	Light	D. strigosa	25.56	36.273	441	2,245 (±14)	1,973 (±32)	8.03 (±0.05)	3.41 (±0.30)
	B29	Dark	P. astreoides	25.56	36.273	0	2,289 (±20)	2,052 (±38)	7.96 (±0.06)	3.04 (±0.31)
	B29	Dark	D. strigosa	25.56	36.273	0	2,299 (±9)	1,989 (±43)	7.98 (±0.07)	3.21 (±0.38)
	NR	Light	P. astreoides	25.56	36.323	206	2,266 (±19)	1,960 (±16)	8.04 (±0.01)	3.44 (±0.08)
	NR	Light	D. strigosa	25.56	36.323	206	2,252 (±21)	2,010 (±45)	7.96 (±0.09)	2.97 (±0.45)
	NR	Dark	P. astreoides	25.56	36.323	0	2,328 (±15)	2,033 (±16)	8.00 (±0.03)	3.30 (±0.14)
	NR	Dark	D. strigosa	25.56	36.323	0	2,324 (±11)	2,040 (±46)	7.95 (±0.02)	3.02 (±0.13)
	WB	Light	P. astreoides	26.11	36.227	58	2,231 (±16)	1,941 (±10)	8.04 (±0.05)	3.39 (±0.31)
	WB	Light	D. strigosa	26.11	36.227	58	2,240 (±21)	1,964 (±8)	8.02 (±0.07)	3.26 (±0.41)
	WB	Dark	P. astreoides	26.11	36.227	0	2,302 (±45)	2,028 (±56)	7.97 (±0.01)	3.08 (±0.09)
	WB	Dark	D. strigosa	26.11	36.227	0	2,286 (±31)	2,039 (±34)	7.93 (±0.02)	2.86 (±0.10)

TA, DIC, pH, and Ω_{arag} change throughout the 1.5-h incubation; values reported here represent the average from the beginning and end of the incubation. Temperature, salinity, and light did not vary throughout the 1.5-h incubation. Values represent the average among replicate samples from each species (*P. astreoides* and *D. strigosa*, site (NR, WB, B29) and treatment (light and dark). Errors represent the standard deviation among replicates. Light values are given at depth (7 m for NR and WB, 3 m for B29), applying a diffusive attenuation coefficient of -0.27

validate the application of calcein dye staining in situ, we compared our results with coral calcification rates assessed using CoralXDS, which provides average annual calcification rates based on annual extension and average coral density (Helmle et al. 2002). Calcification rates from the calcein dye staining method used in this study provide seasonal resolution, and as expected, summer rates (March-September) were higher than the annual average results provided by CoralXDS, while winter rates (September-March) were lower (Fig. 8). In an effort to compare these two methods of measuring coral calcification rates, seasonal rates were converted to annual rates by assuming two 6-month growth periods with rates defined as the average winter and summer calcification rates determined from the seasonal staining study. These average annual calcification rates, scaled temporally from the seasonal staining study, were well correlated with average annual calcification rates obtained from CoralXDS $(r^2 = 0.85, p = 0.0004, n = 18;$ Fig. 8). The good agreement between these two methods suggests that calcein dye staining is an effective method for obtaining in situ seasonal coral extension rates.

Environmental controls on seasonal differences in growth rates

Previous studies investigating seasonal differences in coral calcification rates show a range of differences between summer and winter rates (Table 5; Gladfelter et al. 1978; Dodge and Brass 1984). Part of this difference is attributed to differences in environmental conditions associated with location and latitude. For example, low latitude reefs that experience small seasonal differences also show small differences between summer and winter calcification rates. However, part of the observed differences in seasonal calcification rates may be related to methodological challenges such as the aforementioned problems with alizarin staining. In contrast to low latitude locations, Bermuda

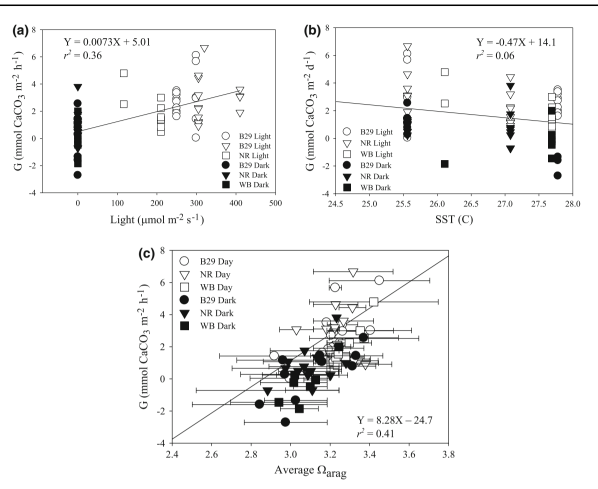


Fig. 7 Hourly coral calcification rates under light (*open*) and dark (*filled*) treatments and corresponding **a** light, **b** temperature, and **c** aragonite saturation states within each incubation chamber at three study sites distributed across the northern Bermuda reef platform. *Error bars* in **c** represent the range of aragonite saturation states

Table 4 Summary of the statistics of spatial trends across the Ber-muda northern reef platform from the March and September 2012spatial surveys

Parameter	March			Septe	mbe	r
	t	df	Significant at $p = 0.01$ Y/N	t	df	Significant at $p = 0.01$ Y/N
DIC	0.07	9	Ν	8.99	9	Y
ТА	1.67	9	Ν	6.09	9	Y
$arOmega_{ m arag}$	0.14	9	Ν	1.34	9	Ν
pCO ₂	-0.22	9	Ν	0.16	9	Ν

For this analysis, 50 sampling stations (Fig. 1) were divided into seven transects extending from the island of Bermuda out to the rim reef. Data from stations at similar distances from shore were then averaged to generate one average cross-platform transect. These data were then plotted as a function of their proximity to shore, testing the null hypothesis that the slope of this average cross-platform transect was zero (H₀: $\beta = 0$)

experience by each coral during the roughly 1.5-h incubation period. $\Omega_{\rm arag}$ is highest at the beginning and decrease throughout the incubation period. Light and temperature remained constant throughout the 1.5-h incubation

experiences large seasonality due to its relatively high latitude (32°N), which results in large differences in coral calcification rates between summer and winter, with an average ΔG between seasons of 77 (±29) mmol CaCO₃ m⁻² d⁻¹, compared with studies from lower latitude reefs, such as the Great Barrier Reef ($\Delta G = 49$, Barnes and Lough 1993; Table 5).

The large seasonal difference in coral calcification rates observed in the present study combined with monthly measurements of environmental parameters (temperature, light, and Ω_{arag}) make it possible to assess the correlation between these parameters and the calcification rates. Corals experienced significantly different average SST, light, and Ω_{arag} during the two growth periods (Fig. 3). The correlations shown in Fig. 5 suggest that each of the factors alone could conceivably explain the observed differences in seasonal growth. Of course, the strong correlations are

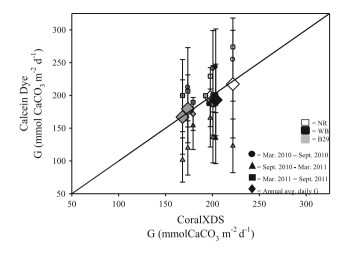


Fig. 8 Comparison of calcification rates measured using Coral XDS (annual rates) and calcein dye staining method (seasonal and annual average rates). Summer sampling periods are indicated by *circles* and *squares*; *triangles* indicate winter sampling period; and *diamonds* represent the average of winter and summer rates. The line designates a 1:1 ratio; points above this line indicate the calcein dying method overestimates calcification with respect to Coral XDS results; points below the line indicate the calcein dye method underestimates calcification. Average seasonal calcification from the calcein dye staining method agree well with results determined from CoralXDS ($r^2 = 0.85$, p = 0.0004, n = 18)

partly the result of the fact that we are working with only two sets of observations (March–September and September–March). We also know that light, temperature, and saturation state covary on both seasonal and hourly timescales. Therefore, we need to move beyond simple correlations in order to evaluate what is controlling the calcification rates measured in this study.

Silverman et al. (2009) and Anthony et al. (2011) took the approach of defining functions that attempted to describe the combined effects of temperature and Ω_{arag} on coral calcification. These functions did not include the effect of light, which certainly is important in explaining seasonal differences nor did they take into account that coral growth often has an asymmetric response to temperature (Houck et al. 1975; Reynaud et al. 1999). We make the assumption that the seasonal change in coral growth can be described by a partial differential equation of the form:

$$\Delta G_{(s-w)} = \frac{\partial G}{\partial I} \Delta I + \frac{\partial G}{\partial T} \Delta T + \frac{\partial G}{\partial \Omega_{\text{arag}}} \Delta \Omega_{\text{arag}}$$
(3)

where $\Delta G_{(s-w)}$ is the difference between summer and winter coral calcification rates, $\frac{\partial G}{\partial I}$, $\frac{\partial G}{\partial T}$, and $\frac{\partial G}{\partial \Omega_{\text{arag}}}$, are partial derivatives that describe the sensitivity of coral calcification (*G*) to a unit change in light, temperature and Ω_{arag} when the other factors are held constant and ΔI , ΔT , and $\Delta \Omega_{\text{arag}}$, are the measured seasonal change in light, temperature, and $\Omega_{\rm arag}$, respectively. Essentially, we are assuming that the growth rate of the corals (G) is a surface in 3-dimensional space (R3) and the three dimensions are light, temperature, and Ω_{arag} . The slope of the gradient in this R3 space is given by Eq. (3) where each of the terms is a vector and the sum of these vectors gives the change in G due to the change in the three environmental factors. We estimate the gradient along each of the three dimensions (axes) based on available laboratory studies. Obviously, this model is very simplistic and ignores covariance terms and the influence of other factors such as food availability, nutrients, and water flow, but it represents a starting point to evaluate whether we can come close to predicting the magnitude of the seasonal change in calcification by considering just three factors and laboratory data on how the rate of calcification changes when each of these factors is varied individually.

Light

Coral calcification rates are strongly dependent on light availability. While many studies have looked at the shortterm response of coral calcification to changes in light on an hourly or day/night time scale (Gattuso et al. 1999; Anthony et al. 2007), what is needed for the present analysis is how coral calcification varies in response to the slow change in daily integrated irradiance that occurs over many days to weeks. On this timescale, the coral-algal symbiosis has time to photo-acclimate to changing light conditions, thereby optimizing its performance through changes in pigment content per symbiont, symbiont density, and even changes in symbiont genotype. Marubini et al. (2001) incubated replicate colonies of Porites compressa at different light levels for 6 weeks and determined that the relationship between photo-acclimated growth rate, and the average daily PAR was well described by the same hyperbolic tangent function used to describe the instantaneous rate of light calcification (Chalker 1981) and ranged from 7.8 to 9.0 mol photons $m^{-2} d^{-1}$ per unit change in G and averaged 8.4 mol photons $m^{-2} d^{-1}$ per unit change in G. The change in G due to light (ΔG) is then given by:

$$\Delta G = G_{\max} * \left(\tanh\left(\frac{E_1}{I_k}\right) - \tanh\left(\frac{E_2}{I_k}\right) \right) \tag{4}$$

where G_{max} is the light-saturated rate of growth in mg cm⁻² d⁻¹, E_1 and E_2 are the daily integrated photosynthetically available light at time 1 and time 2, and I_k is the light saturation parameter in units of mol photons m⁻² d⁻¹.

The first step in determining the effect of the seasonal change in light is to estimate the amount of light that reaches the corals. The light at depth is given by $I_0 \exp(-k_z * z)$ where I_0 is the incident PAR at the water surface, k_z

Table 5 Seasonal coral calcification rates

Study	Study site	Method	Growth period	Coral species	$G \text{ (mmol} \text{CaCO}_3 \text{ m}^{-2} \text{ d}^{-1} \text{)}$	Seasonal ΔG (mmol CaCO ₃ m ⁻² d ⁻¹)
Gladfelter et al. (1978)	St Croix, USVI	X-ray	Summer	M. annularis	355 (±24)	73
Gladfelter et al. (1978)	St Croix, USVI	X-ray	Winter	M. annularis	282 (±21)	
Gladfelter et al. (1978)	St Croix, USVI	X-ray	Summer	M. annularis	299 (±18)	-81
Gladfelter et al. (1978)	St Croix, USVI	X-ray	Winter	M. annularis	380 (±18)	
Gladfelter et al. (1978)	St Croix, USVI	X-ray	Summer	P. astreoides	143 (±13)	7
Gladfelter et al. (1978)	St Croix, USVI	X-ray	Winter	P. astreoides	136 (±8)	
Barnes and Lough (1993)	Central BGR	X-ray	Summer	Porites	372 (±33)	49
Barnes and Lough (1993)	Central BGR	X-ray	Winter	Porites	323 (±32)	
Dodge and Brass (1984)	Gulf of Mexico	X-ray	Summer	M. annularis	291 (±33)	32
Dodge and Brass (1984)	Gulf of Mexico	X-ray	Winter	M. annularis	259 (±32)	
This study	Bermuda, WB	Staining/X-ray	Summer	P. astreoides	210 (±27)	65
This study	Bermuda, WB	Staining/X-ray	Winter	P. astreoides	145 (±13)	
This study	Bermuda, B29	Staining/X-ray	Summer	P. astreoides	222 (±77)	87
This study	Bermuda, B29	Staining/X-ray	Winter	P. astreoides	135 (±39)	
This study	Bermuda, NR	Staining/X-ray	Summer	D. strigosa	253 (±22)	109
This study	Bermuda, NR	Staining/X-ray	Winter	D. strigosa	144 (±30)	
This study	Bermuda, WB	Staining/X-ray	Summer	D. strigosa	204 (±51)	93
This study	Bermuda, WB	Staining/X-ray	Winter	D. strigosa	111 (±13)	

is the diffuse attenuation coefficient for PAR, and z is the depth in m. No data could be found for k_z on the Bermuda platform. We recognize that k_z varies both temporally and spatially due to changes in suspended particulates. The turbidity of the water in Bermuda varies between high in summer and low in winter. Certain areas of the reef platform (particularly B29) experience high turbidity following the passing of cruise ships. Given the lack of seasonal and spatial data on k_z on the Bermuda platform, we make the assumption that it is similar to the value we have measured in the upper Florida Keys and on Davies Reef in the Great Barrier Reef, i.e., -0.3 m^{-1} (C. Langdon unpublished data). Making this assumption, we believe that the attenuation coefficient is on the high end, and hence, our estimate of the effect of light on calcification will be on the conservative or low side. Based on the depth of our corals at approximately 5 m at our three study sites, we compute that the average light levels as 12.3 mol photons $m^{-2} d^{-1}$ for the summer (March-September) growth period and 7.2 mol photons $m^{-2} d^{-1}$ for winter (September-March). If we plug in 12.3 and 7.2 as E1 and E2, 8.4 as I_k , and 215 as G_{max} into Eq. (4), we find that the change in light could account for a 44 mmol $m^{-2} d^{-1}$ decrease in calcification rate between summer (March-September) and winter (September-March) or 54 % of the total seasonal change. If k_z is significantly greater than 0.3 m^{-1} during the winter (September-March) period, then light could play an even greater role in driving the seasonal change in G.

Temperature

Data on the effect of temperature on coral growth in wellcontrolled experimental studies is sparse, but based on what we have been able to find, the effect is symmetric about the temperature optimum for some species and asymmetric for others (Table 6). These temperature studies did not account for the corresponding change in Ω_{arag} , which also affects calcification. Though we recognize this as a limitation of our estimate of the effect of temperature on calcification, we argue that the corresponding change in $\Omega_{\rm arag}$, due to a change in temperature is small and will therefore have a minimal influence on calcification in comparison with temperature. A 2 °C change in temperature results in a 0.03 unit change in Ω_{arag} . If we assume a 15 % change in calcification for every unit change in $\Omega_{\rm arag}$ (Chan and Connolly 2013), this results in a 0.45 % change in G, which is small compared with the influence of temperature. We find that on average growth rate decreases by $12 \pm 4 \% \ ^{\circ}C^{-1}$ below the species temperature optimum and decreases by $24 \pm 17 \% \circ C^{-1}$ when temperature exceeds the temperature optimum. Utilizing this information in our study requires knowing the temperature optimum for our species. We could find nothing in the literature specific to the two species used in this study or for corals growing on the Bermuda platform. In general, temperature optima of tropical corals range from 25 to 29 °C (Vaughan and Wells 1943). However, the optimum is specific to the range of temperature that the corals

Species	Origin of colonies	<i>T</i> (°C) rang	% change in G per °C for $T < T_{opt.}$	% change in G per °C for $T > T_{opt}$.	Reference
M. capitata	Kaneohe Bay, HI	20-28	8	-8	Coles et al. (1976)
P. lobata	Kaneohe Bay, HI	22-28.5	11.1	-57	Houck et al. (1975)
G. fascicularis	Heron Reef, GBR	21–29	15.2	-18	Marshall and Clode (2004), Al Horani (2005)
S. pistilata	Gulf of Aqaba	21-29	8.1	-32	Reynaud et al. (1999)
Acropora sp.	Gulf of Aqaba	21-29	6.9	-9	Reynaud et al. (1999)
P. damicornis	Kaneohe Bay, HI Eniwetok Atoll, Marshall Islands	22.6–28	16.9	-18	Clausen and Roth (1975)
Average			12	-24	
SD			4	17	

Table 6 Examples of the effect of temperature both above and below the optimal temperature for coral growth (T_{opt}) on coral calcification rates

experience at a specific location. Jokiel and Coles (1974) found that the temperature optima for Pocillopora damicornis ranged from 31 °C for specimens collected at Eniwetok (11.5 N) to 27 °C for corals collected in Kaneohe Bay, Oahu (21 N) to 23 °C for corals collected at Midway Atoll (28 N). Based on these studies, it can be inferred that the optima is close to or slightly above the average annual temperature that the species encounters. The annual average SST for the Bermuda coral reef platform from 2010 to 2011 was 23.4 °C (Fig. 3) with a summer maximum of 29 °C. Even though Bermuda is located at a higher latitude (32°N) than Midway, it experiences summer temperatures that are approximately 2 °C warmer. Based on this, we assume that the temperature optimum of corals in Bermuda is in the 23.4–25.4 °C range, but we recognize that this range could be broader and very likely span a higher temperature range. The percentage change in calcification between two temperatures is given by:

If
$$T_{\rm L} < T_{\rm opt} < T_{\rm H}$$
:
 $\Delta\% G = (12\%^{\circ}{\rm C}^{-1})(T_{\rm opt} - T_{\rm L}) + (-24\%^{\circ}{\rm C}^{-1})(T_{\rm H} - T_{\rm opt})$
(5)

If
$$T_{\rm opt} > T_{\rm H}$$
: $\Delta\% G = (12\%^{\circ} {\rm C}^{-1})(T_{\rm H} - T_{\rm L})$ (6)

where $\Delta\% G$ is the percentage change in the rate of calcification, 12 and $-24 \% \,^{\circ}\text{C}^{-1}$ are the sensitivity of calcification to a unit change in temperature from Table 6, T_{opt} is the optimum temperature for growth, and T_{L} and T_{H} are the low and high temperatures of interest. T_{L} and T_{H} in this study are 21 and 25 °C, respectively. Based on Eq. (5) and a T_{opt} that ranges from 23.4 to 25.4 °C, we compute that $\Delta\% G$ ranges from -10 to +38 % and averages +24 %. If T_{opt} exceeds 25.0 °C (Eq. 6), then the $\Delta\% G$ asymptotes out at 48 %. A 24 % reduction in the summer (March–September) calcification rate amounts to a change of 52 mmol m⁻² d⁻¹. A 24 % decrease in the winter (September–March) rates amounts to a change of 32 mmol m⁻² d⁻¹. Seawater aragonite saturation state

Lastly, we consider the contribution from the 0.16 unit change in Ω_{arag} observed between summer and winter. Laboratory studies have found that the mean response of coral calcification is -15 ± 8 % per unit decrease in Ω_{arag} (Chan and Connolly 2013). Based on this sensitivity we estimate that the change in Ω_{arag} would cause a 2.4 % or 4 mmol m⁻² d⁻¹ change in coral calcification rate.

In order to compare the sensitivities (slope of calcification vs. Ω_{arag} relationship) measured in different studies, it is first necessary to normalize the data because the absolute rates may be different. A simple way to normalize the rates is to express them relative to a rate at a reference seawater aragonite saturation state. While any Ω_{arag} can be used, it is advantageous to use the pre-industrial average seawater $\Omega_{\rm arag}$ of 4.6 because the rates then reflect the change since the onset of modern anthropogenic ocean acidification. For data sets where an equation of the form y = mx + b has been provided, multiplying m by [100/ $(m^*4.6 + b)$] yields a normalized slope or sensitivity that reflects the percentage change in calcification per unit change in Ω_{arag} . This is how the single organism sensitivities were compared in Chan and Connolly (2013), which showed an average of 15 ± 8 % change in G per unit change in Ω_{arag} . The stronger sensitivity of 63 % change in G per unit change in Ω_{arag} observed in this study for both hourly and seasonal calcification rates is much greater than that observed in controlled experiments (Fig. 9). A stronger sensitivity of community calcification to a change in Ω_{arag} has been previously noted (Pandolfi et al. 2011). This strong sensitivity results in a threshold Ω_{arag} of 2.97, below which we should expect dissolution; however, we know that to be incorrect as corals on the Bermuda platform continue to calcify at $\Omega_{\text{arag}} < 3.0$. With regard to the seasonal data, this strong sensitivity may be a result of the covariance of Ω_{arag} with other driving factors like

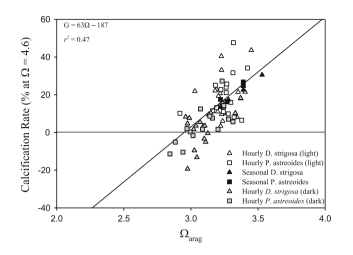


Fig. 9 Calcification rates from *Diploria strigosa* (*triangles*) and *Porites astreoides* (*squares*) from the dark (*gray*) and light (*white*) hourly incubation study and seasonal staining study (*black*) normalized to the calcification rate at $\Omega_{arag} = 4.6$ and corresponding Ω_{arag} . Hourly Ω_{arag} shown are the average values from the ~1.5-h incubation period. Seasonal average Ω_{arag} were determined from monthly water sampling by the Bermuda Ocean Acidification and Coral Reef Investigation (BEACON) project

temperature and light. With regard to the hourly incubation data, the strong sensitivity may be a result of the effects of respiration or dissolution on Ω_{arag} within the chambers. Alternatively, if environmental conditions (e.g., light, temperature, food availability) in the field were sub-optimal, corals may be more sensitive to Ω_{arag} compared with corals reared in controlled environments where these parameters are maintained at optimal levels.

Controlling factors of seasonal coral calcification in Bermuda

In summary, our analysis suggests that light and temperature contribute roughly equally to the seasonal difference in calcification, i.e., 44–54 mmol m⁻² d⁻¹ and the change in Ω_{arag} contributes about 4 mmol m⁻² d⁻¹. Thus, the total seasonal difference in calcification based on this assessment is roughly 102 mmol m⁻² d⁻¹, which is more than the observed change of 81 ± 14 mmol m⁻² d⁻¹, suggesting that the cumulative effects of changing light, temperature, and saturation state are less than the individual effects based on experimental studies.

It is encouraging that the prediction is not significantly different from the observations, but the uncertainty of the prediction is undoubtedly large. Future work is required to better constrain the I_k and T_{opt} of the Bermuda corals and determine if there are important covariance terms that we are ignoring.

Environmental controls on hourly differences in growth rates

During our short-term incubation experiments, we found the response of calcification to changes in light to be 0.007 mmol $m^{-2} h^{-1}$, which is similar to that reported by Langdon and Atkinson (2005; 0.01 mmol $m^{-2} h^{-1}$) and Gattuso et al. (1996; 0.005 mmol $m^{-2} h^{-1}$). During the light incubations, Ω_{arag} increased by 0.13 ± 0.06 (SD) units, and during the dark incubation, it decreased by 0.18 ± 0.07 units. Calcification was similarly correlated with both light ($r^2 = 0.36$) and Ω_{arag} ($r^2 = 0.41$; Fig. 7a,c). the slope of G versus While light plot (0.007 mmol $m^{-2} h^{-1}$ compares well with other studies, the slope of the G versus Ω_{arag} implies a sensitivity that is much greater than observed in other single organism laboratory studies. This can be seen by normalizing the hourly data as described above for the seasonal data (Fig. 9). The slope of the normalized hourly data is 63 % per unit Ω_{arag} , which is similar to that reported for the seasonal data and likewise four times greater than the average coral sensitivity (Chan and Connolly 2013). We think that this high sensitivity to $\Omega_{\rm arag}$ is overestimated. Photosynthesis and respiration have a direct impact on Ω_{arag} (i.e., photosynthesis results in an increase in Ω_{arag} , while respiration decreases Ω_{arag}), which in turn affect calcification rates. In this study, photosynthesis is causing $\varOmega_{\rm arag}$ to rise in the light incubations, and respiration is causing it to decline in the dark incubations. This net change in Ω_{arag} between the light and dark incubations is 0.13 units. Calcification rates of the corals do respond to these changes in Ω_{arag} , but it is also responding to the light. The effect of light is much greater than the effect of Ω_{arag} . The average light received is sufficient to increase calcification rates by 0.0073 * (200-400) or 1.5-3 mmol m⁻² h⁻¹, while the 0.13 unit increase in Ω_{arag} probably increases G by only 0.04 mmol m⁻² h⁻¹. In the dark incubation, the effect of the removal of light causes G to abruptly decrease to the dark rate, while the decrease in Ω_{arag} probably only contributes a decrease of only 0.05 mmol $m^{-2} h^{-1}$. The decrease in $\Omega_{\rm arag}$ in the dark incubation may also cause the rate of dissolution to increase. This increase in dissolution may be why we see net dissolution in some of the chambers (Fig. 6).

The combination of decreased calcification and increased dissolution would result in an even larger decrease in G. The net effect of light and dark is to cause a large change in G that looks like a very strong sensitivity to Ω_{arag} if the full change in G is mapped onto the 0.31 unit change in Ω_{arag} , but is actually largely due to light and perhaps also the stimulation of dissolution in the dark incubations.

TA:DIC property–property plots and lack of a spatial gradient in Ω_{arag} across the platform

To understand why we are not seeing gradients in Ω_{arag} across the platform, which is a common feature of many reefs (Kawahata et al. 1997, 2000; Suzuki and Kawahata 2003; Watanabe et al. 2006; Albright et al. 2013), we turn to the plot of salinity-normalized TA versus DIC, which shows data points from the 50 station platform-wide surveys conducted in March and September 2012 (Fig. 10). The points tend to fall along a fairly tight line with a slope of 0.93 ± 0.04 that closely approximates the slope of the Ω_{arag} isopleths. This is not significantly different from the 1.05 ± 0.04 reported by Andersson et al. (2013). This means that the biota on the platform is removing DIC and TA from the water in a fairly constant ratio both temporally and spatially and that this ratio balances the effect of photosynthesis to elevate pH (reduce pCO₂) and calcification to reduce pH (elevate pCO₂; Andersson and Gledhill 2013). When the two processes combine in this ratio, the DIC and TA can be drawn far down from their source water values without altering the pH, pCO₂, or Ω_{arag} significantly. One consequence of this is that you do not find spatial gradients in pH, pCO₂, or Ω_{arag} as you do on many other reefs where the slope of the TA:DIC line tends to be lower.

Understanding the relationship of temperature, light, and Ω_{arag} on coral calcification rates requires high temporal and spatial resolution of environmental data. The monthly data showed sustained differences in Ω_{arag} and pCO₂ at the rim reef (NC) compared with nearshore sites (TB and B33). The platform surveys in March and September, however, showed that despite significant differences between the rim reef and nearshore sites, there was no consistent or uniform spatial gradient across the platform. This highlights the importance of obtaining environmental data at both high spatial and temporal resolutions in order to understand how these parameters are influencing coral calcification rates.

Seasonal coral calcification rates were measured at three study sites across the Bermuda coral reef platform and showed a difference between seasons that was reasonably well explained by a simple linear model that combined the known sensitivity of coral growth to temperature, light, and $\Omega_{\rm arag}$ from laboratory studies with the measured differences in temperature, light, and Ω_{arag} . We have presented a new way to assess the impact of different driving parameters on coral calcification rates, based on laboratory relationships between G and environmental driving factors, and measured differences of these environmental factors. Based on these assessments, seasonal differences in coral calcification rates in Bermuda are driven mostly by the seasonal differences in temperature and light. Differences between hourly calcification rates under light and dark treatments are most likely driven by light, enhanced by the effect of

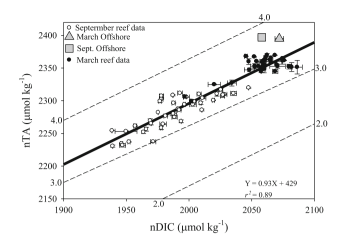


Fig. 10 Salinity normalized (S = 36.6) total alkalinity versus dissolved inorganic carbon from data collected at 50 sampling stations distributed across the Bermuda reef northern platform in March and September, 2012 (*open circles*), with offshore end members from samples collected in the Sargasso sea during corresponding reef sampling expeditions (*filled triangle and square*). Hashed lines are aragonite saturation state contours at 25 °C. The slope of the type II linear regression of the reef data (*black line*) equals 0.93 ± 0.04 (*n* = 79, *r*² = 0.89)

respiration and photosynthesis on Ω_{arag} under dark and light conditions, respectively.

Though the sensitivity of normalized calcification to a unit change in Ω_{arag} was found to be similar and high at hourly and seasonal time scales, i.e., 63 % per unit change in $\Omega_{\rm arag}$, we believe this sensitivity is misleading. In the chamber study, this high sensitivity can be explained by the covariant nature of light and $\Omega_{\rm arag}$, which both have a positive effect on calcification, with the former having a much stronger effect. In the seasonal study, this high sensitivity may be a result of the covariant nature of temperature, light, and $\Omega_{\rm arag}$. On the other hand, it is important not to underestimate the role that ocean acidification may play in controlling demographics in coming decades. Not assessed in field studies such as this one is the possibility that a unit change in saturation state, pH, or pCO₂ can have a much larger impact on growth or net photosynthesis when the temperature approaches the upper thermal limit of the coral, i.e., a synergistic interaction between pH and temperature (Anthony et al. 2008; Crawley et al. 2010; Kaniewska et al. 2012).

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