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FEATURE ARTICLE

Experimental impacts of climate warming and ocean carbonation on eelgrass Zostera marina

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ABSTRACT: CO₂ is a critical and potentially limiting substrate for photosynthesis of both terrestrial and aquatic ecosystems. In addition to being a climate-warming greenhouse gas, increasing concentrations of CO₂ will dissolve in the oceans, eliciting both negative and positive responses among organisms in a process commonly known as ocean acidification. The dissolution of CO₂ into ocean surface waters, however, also increases its availability for photosynthesis, to which the highly successful, and ecologically important, seagrasses respond positively. Thus, the process might be more accurately characterized as ocean carbonation. This experiment demonstrated that CO₂ stimulation of primary production enhances the summertime survival, growth, and proliferation of perennial eelgrass Zostera marina from the Chesapeake region, which is regularly impacted by summer heat stress. The experiment also quantified the logarithmic response to CO₂ in terms of shoot proliferation, size, growth and sugar accumulation that was fundamentally consistent with model predictions based on metabolic carbon balance derived from short-term laboratory experiments performed with other eelgrass populations from cool ocean climates and other seagrass species from tropical and temperate environments. Rather than acting in a neutral fashion or as an independent stressor, increased CO₂ availability can serve as a quantitative antagonist to counter the negative impact of climate warming on seagrass growth and survival. These results reinforce the emerging paradigm that seagrasses are likely to benefit significantly from a high-CO₂ world.

KEY WORDS: Climate warming · Ocean acidification · Eelgrass · Carbon balance · Photosynthesis · Temperature · CO₂

INTRODUCTION

The global monthly mean CO₂ concentration of the earth’s atmosphere exceeded 400 ppm in 2016 (Dlugokencky & Tans 2016) and appears to be increasing significantly faster than previously anticipated (IPCC 2013). Present-day concentrations are now higher than at any time in the last 40 million years (Pearson & Palmer 2000). Not coincidentally, the earth’s climate has warmed nearly 1°C since 1880, generating phenological changes in natural ecosystems including the progressively early occurrence of spring activities (budbreak, reproduction, migrant arrival), extended growing seasons (days above freezing), a delay in the onset of autumnal activities including outmigration, leaf coloring and abscission, and shifts in species range and commu-
nity composition (Walther et al. 2002, Cleland et al. 2007).

In addition to being a climate-warming greenhouse gas, CO₂ is a critical and potentially limiting substrate for photosynthesis in both terrestrial and aquatic ecosystems. CO₂ enrichment experiments with terrestrial plants have shown (1) enhanced carbon uptake, even in the presence of photosynthetic acclimation, (2) increased nitrogen and water use efficiencies, and (3) increased dark respiration, although the positive impacts on crop yield have been less than predicted (see review by Leakey et al. 2009 and references cited therein).

Similar to the impacts on terrestrial ecosystems, the increased concentrations of CO₂ dissolved in the oceans of the world will elicit both negative and positive responses among organisms, ultimately potentiating ecological losers and winners. Organisms and ecosystems, including hermatypic corals and marine bivalves (e.g. oysters) and the reef structures they generate, as well as pelagic pteropods, to name but a few, are likely to be negatively impacted as calcification becomes energetically less favorable in an acidified ocean (Kleypas et al. 1999, 2006, Fabry et al. 2008). The dissolution of CO₂ into ocean surface waters, however, also increases its availability for photosynthesis. Although many marine primary producers are predicted to exhibit very little response to ocean carbonation (Mackey et al. 2015), photosynthetic rates of several taxonomic groups, including nitrogen-fixing cyanobacteria (Hutchins et al. 2007), coccolithophores (Rivero-Calle et al. 2015) and the weedy chlorophytes Caulerpa spp. and Ulva spp. (Beer 1994, Hall-Spencer et al. 2008), as well as the highly successful and ecologically important seagrasses (Zimmerman et al. 1997, Invers et al. 2001, Jiang et al. 2010, Campbell & Fourqurean 2013), have been shown to respond positively to increased CO₂ concentrations.

The seagrass Zostera marina L. (eelgrass) is widely distributed on sandy shores and estuaries throughout the temperate Northern Hemisphere, where it plays an important role in sediment stabilization and habitat provision for many ecologically and economically important invertebrates and fish, and represents a significant blue carbon sink (Orth et al. 2006, McLeod et al. 2011). Positive long-term (>1 yr) effects of dissolved aqueous CO₂ concentration ([CO₂(aq)]) on eelgrass performance were previously quantified using a population from the cool and thermally stable eastern Pacific waters of central California, in which the ambient temperature ranged between 12 and 16°C across seasons (Palacios & Zimmerman 2007).

Eelgrass from the Chesapeake region of the western Atlantic, however, are typically exposed to temperatures that range from wintertime lows near 5°C to summertime highs exceeding 25°C. Temperatures can approach 30°C during particularly warm summers, stressing eelgrass meadows throughout the southern Chesapeake Bay (Orth & Moore 1983, Moore & Jarvis 2008, Moore et al. 2012). Eelgrass can acclimate to temperatures below 25°C (Zimmerman et al. 1989), but exposure to simulated heat waves above 25°C causes die-backs (Ehlers et al. 2008) that can be linked to impaired photosynthetic performance (Winters et al. 2011) as well as to the differential effects on respiration and CO₂-limited photosynthesis that make eelgrass vulnerable to negative carbon balance above 25°C (Evans et al. 1986, Zimmerman et al. 1989). CO₂ stimulation of photosynthesis should reduce eelgrass vulnerability to negative carbon balance that is likely to occur with increasing frequency as the climate warms (Zimmerman et al. 1997, 2015, Invers et al. 2001).

The goal of this experiment was to quantify the extent to which CO₂ enrichment can improve the ability of eelgrass to tolerate stressful summer temperatures typical of the Chesapeake/Pamlico region that defines the southern limit of eelgrass distribution in the temperate western North Atlantic and to explore the longer-term impacts on plant size, biomass allocation between shoots and roots/rhizomes, flowering, leaf sugar, and photosynthetic pigment content. This experiment also tested model predictions based on short-term physiological experiments that prolonged CO₂ enrichment could reduce the lethal impacts of high summertime water temperature on eelgrass survival (Zimmerman et al. 2015). The insights into processes controlling carbon balance in seagrasses provided by this experiment offer to improve our predictive understanding of the simultaneous impacts of ocean warming and carbonation/acidification on the photosynthetic metabolism, vegetative growth and reproductive success of seagrasses in the context of a changing ocean climate.

**MATERIALS AND METHODS**

**Experimental facility**

We constructed an experimental facility at the Virginia Aquarium & Marine Science Center, adjacent to Owls Creek, VA, consisting of 20 fiberglass aquaria (3 m³ volume each, Fig. 1). The long axis of each aquarium was oriented in a north–south direction to
minimize the effect of wall shadows on irradiance to the plants. Natural water from Owls Creek was pumped directly into a large holding tank and then flowed by gravity through a bottom diffuser at the inlet of each aquarium. Flow rates were individually adjusted to ensure 10 complete turnovers in each aquarium per day (min. flow = 20 l min⁻¹). Water was discharged through a standpipe at the opposite end that maintained unidirectional (but not laminar) flow and a fixed water depth of 1 m in each aquarium. Water temperature was monitored by a high-precision thermistor (Omega 44005) placed in each aquarium. Thermistors were calibrated every 6 mo at 6 tempera-

tures from 5 to 30°C using a temperature-controlled water bath (Fisher Scientific Iso-temp). Salinity in each aquarium was measured a few times in May and June of 2013 using a hand-held refractometer (data not shown). No aquarium-to-aquarium differences were found among the refractometer measurements. Beginning in July 2013, salinity was continuously monitored within the system using a factory-calibrated SeaBird SBE37-SM CTD installed into one of the aquaria; all values are reported using the Practical Salinity Scale (PSS, Lewis 1980). Incident downwelling plane irradiance (photosynthetically active radiation, in air) was monitored with a factory-calibrated LiCor Li-190 quantum radiometer. A layer of neutral-density window screening (60% transmission) prevented photodamage to upper leaf surfaces in the very shallow water of the aquaria. All instrument data were recorded at 10 min intervals using a National Instruments (NI) data logging and control system and custom software written in LabVIEW®. The aquaria and eelgrass shoots were cleaned weekly to remove biofilms and leaf epiphytes as well as any animals (fish, crabs, littleneck clams) introduced with the water. Mixing within the aquaria was accomplished by forcing air from a high-performance aquarium compressor through a 2 m length of Pentair Bio-Weave® diffuser hose running along the bottom of each aquarium.

Beverage-grade CO₂ was injected into the air stream at each aquarium (except ambient) beginning 1 June 2013 and continued through November 30 2014 (18 mo) using solenoid valves individually controlled by Eutech Alpha pH 190 controller/transmitters equipped with submersible glass electrodes. Controllers were poised at 5 different pH values (4 aquaria at each pH) ranging from ambient (pH ~7.7, no CO₂ addition) to pH 6.0 ([CO₂(aq)] = 2121 µM) at intervals of approximately 0.5 pH unit as indicated on Fig. 1. The 5 pH treatments were distributed evenly between the 2 rows of 10 aquaria (2 replicate pH treatments per row). The position of the pH treatments in each row was determined randomly except that replicate pH treatments were located adjacent to each other in the 2 rows because our initial plan was to heat 1 aquarium of each pH pair 5°C above ambient to achieve stressful temperature conditions during the summer. However, the ambient summertime temperature reached 30°C in both years, and we elected not to heat the aquaria as temperatures
above 30°C are likely to produce acute thermal stress (Zimmerman et al. 1989) that was beyond the scope of this experiment. Since all aquaria were fed from the same water source and our monitoring data revealed no position effects in terms of light, temperature or salinity among the aquaria, we elected not to randomly redistribute the pH pairs in each row after the experiment had begun. Potential effects of aquarium position were subsequently ignored, and each aquarium was treated as an independent replicate in the statistical analysis of the experimental data.

The pH electrodes were cleaned and the pH controller systems calibrated weekly against NIST SRM 185 & 186 standard buffers (pH 4, 7 and 10). Although $[CO_2]_{aq}$ at pH settings below 7.5 exceeded the range of ocean acidification predicted by the IPCC through the end of this century, estuarine systems are known to experience a much wider and more temporally variable range in pH/CO$_2$ than the open ocean (Duarte et al. 2013, Waldbusser & Salisbury 2014, Ruesink et al. 2015). More importantly, this range provided a useful gradient in CO$_2$ availability required to determine functional responses (slopes and intercepts) necessary for predicting the performance of eelgrass in any CO$_2$ world—past, present or future.

Alkalinity was determined approximately weekly by automated titration using 0.020 N HCl (Gieskes & Rodgers 1973) and showed a strong predictive relationship with salinity: alkalinity (µmol kg$^{-1}$ seawater [SW]) = (49.61 × salinity) + 694.79, $r^2 = 0.86$). Because no other acidifying agents or buffers were employed, alkalinity values derived from the salinity record, along with temperature and pH, were used to determine the total CO$_2$ ($\Sigma$CO$_2$) content in each experimental aquarium as well as the distribution of $\Sigma$CO$_2$ into the constituent fractions of CO$_2$$_{2aq}$, HCO$_3$$^-$ and CO$_3^{2-}$ using CO2SYS (Ver. 01.05, Lewis & Wallace 1998).

**Eelgrass source population and transplants**

Vegetative shoots were collected by SCUBA divers from a recently restored eelgrass meadow in South Bay, a coastal lagoon at the southern end of the Delmarva Peninsula (Orth et al. 2010), and transported to the experimental facility in insulated coolers in May 2013. Approximately 50 individual vegetative shoots with intact rhizomes and root bundles were transplanted into rectangular fiberglass-reinforced plastic containers (0.04 m$^3$ volume, 0.075 m$^2$ surface area) filled with clean intertidal sediment (sand) collected from the Old Dominion University (ODU) sailing center on the Elizabeth River, Norfolk, VA, resulting in an initial density of 700 shoots m$^{-2}$. Five containers were placed into each aquarium at a water depth of 0.85 m, placing the top of the canopy at about 0.5 m beneath the surface of the water at the beginning of the experiment.

**Whole plant performance**

All shoots were counted in every container and their flowering status noted each month from May 2013 to October 2014 (18 mo total). All abscised leaves and floating dead shoots were removed from the aquaria every few days. One shoot from each container was randomly selected each month and marked with a 20 gauge hypodermic needle 1 wk prior to measurement using the hole-punch method (Zieman 1974, Zimmerman et al. 1996). Marked plants were also tagged with a small plastic cable tie to prevent resampling the same shoots in consecutive months. Young unmarked leaves were assumed to be new growth. The length of new leaf material below the punch mark and the total length of all leaves were measured to the nearest millimeter using a flexible meter tape. Leaf width (nearest 0.1 mm) was measured with a digital caliper. Total leaf area (cm$^2$ shoot$^{-1}$) was calculated with a digital caliper. Total leaf area (cm$^2$ shoot$^{-1}$) was calculated by summing the 1-sided area (leaf length × leaf width) of all leaves of the shoot. Absolute growth (cm$^2$ shoot$^{-1}$ d$^{-1}$) was calculated by normalizing the new leaf area by the elapsed time (d) since the shoot was marked. Specific growth (% d$^{-1}$) was calculated by normalizing absolute growth rates by the total leaf area measured at the end of the marking period.

Biomass allocation among shoots, rhizomes, and roots was measured only at the end of the experiment in October 2014 because it required destructive sampling of the shoots. Three shoots were carefully removed from the sediment in each aquarium by hand, to avoid breaking rhizomes and roots, and returned to the laboratory for determination of fresh weight, dry weight, and chemical analysis of leaves, rhizomes, and roots. The average results from each aquarium, without error estimates for within-tank replication, were used in subsequent statistical analyses.

**Chemical composition**

Each month, a segment of leaf No. 2 (No. 1 was the youngest leaf) was collected from each shoot that was marked for growth. Leaf epiphytes were removed by gentle scraping with a razor blade. Fresh
leaf segments were ground in ice-cold 80% acetone, and pigment content (chl $a$, chl $b$ and total carotenoids) was determined spectrophotometrically using the equations of Lichtenthaler & Wellburn (1983). The remaining leaf samples were dried at 60°C for 48 h and ground in liquid nitrogen. Sucrose was extracted from the ground dry tissue 3 times using hot (80°C) ethanol (Zimmerman et al. 1989). The 3 extractions were combined, and an aliquot was evaporated to dryness under a stream of compressed air, redissolved in distilled water, and analyzed spectrophotometrically using a resorcinol assay standardized to sucrose (Huber & Israel 1982).

**Statistical analysis**

Within-aquarium replicate measures of each performance property were combined each month to generate statistically independent means for each aquarium (without error), resulting in 4 statistically independent replicate measurements for each CO$_2(aq)$ treatment each month. Although CO$_2(aq)$ concentrations in the aquaria were manipulated using the pH controllers, they were also affected by salinity/alkalinity and especially water temperature such that [CO$_2(aq)$] was not absolutely fixed across time (see Fig. 2e). Consequently, statistical significance of treatment effects were determined using a repeated-measures ANCOVA implemented in the mixed model analysis of the generalized linear models component of IBM SPSS Statistics 22 using Month (i.e. time) as the fixed factor (within subjects) and pH as the covariate (between subjects). The aquaria were treated as repeated subjects. Statistical differences among CO$_2(aq)$-dependent slopes across time were determined by post hoc analysis of the 95% confidence intervals derived from the mixed model analysis. ANCOVA results for significance and linearity of the individual regression slopes for each monthly measurement are available from the Biological and Chemical Oceanography Data Management Office data repository (www.bcodmo.org/project/2141). Effects of pH on plant size and biomass allocation among leaves, rhizomes, and roots at the end of the experiment were quantified using Model I linear regression. All error terms are expressed as standard errors unless otherwise noted.

**Heat maps**

Measures of whole plant performance and chemical composition are presented as colored contour plots across the experimental domain defined by CO$_2(aq)$/pH (vertical axes) and time (horizontal axes) using the mean value for each treatment at each time point. The 5 CO$_2(aq)$/pH treatments are indicated by the tick marks on the left vertical axis of each heat map. Tick marks on the temporal (horizontal) axes indicate the middle of each month when performance measures were taken. Regression slopes and their standard errors relating whole plant performance and chemical composition to log[CO$_2(aq)$] were computed using all the aquarium replicates (4 per CO$_2(aq)$/pH treatment) in a Model I regression. White fields on the heat maps indicate missing data values.

**RESULTS**

**Experimental conditions**

The outdoor aquarium system used for this experiment exposed all treatments to daily and seasonal variations in total solar radiation, ambient temperature and salinity. Irradiance varied seasonally in response to solar elevation and day length, producing 3-fold more total daily irradiance in summer than winter (Fig. 2a). Day-to-day variations in irradiance resulting from weather and clouds often exceeded the seasonal range in total daily irradiance, particularly in summer. The seasonal cycle in water temperature lagged the pattern in total daily irradiance by 6 to 8 wk (Fig. 2b). Temperature varied among the aquaria by <1°C and cycled daily by ~5°C around the mean values in summer and 2°C in winter. Summer high temperatures reached 30°C each summer and exceeded the 25°C threshold for eelgrass stress (Evans et al. 1986, Zimmerman et al. 1989) for at least 1 h d$^{-1}$ on 97 and 124 d in 2013 and 2014, respectively. Water temperature exceeded the 25°C threshold all day long for 48 and 73 d in 2013 and 2014, respectively. Winter low temperatures averaged about 5°C from January through March 2014 and approached 0°C on a few days in February 2014.

Hydrography of the Owls Creek source water is dominated by tidal exchange with the mid-Atlantic Ocean at Rudee Inlet, just south of the Chesapeake Bay mouth (Sisson et al. 2010), resulting in a mean salinity of 24 ± 3 (PSS) punctuated by brief spikes of low salinity resulting from local rainfall events and runoff into the watershed (Fig. 2c). Unlike irradiance and temperature, there was no obvious seasonal pattern in the salinity signal. Aragonite saturation of the source water, calculated from measured
Fig. 2. Environmental conditions during the experiment. (a) Total daily irradiance. Open symbols represent unscreened values measured by the PAR sensor. Filled symbols represent the irradiance incident on the plants below the 40% reduction window screening. (b) Ambient temperature scaled according to the left vertical axis. Solid black line indicates the mean for all aquaria. Gray shading around the mean line indicates daily maximum and minimum values. Open symbols indicate the number of hours each day that temperature exceeded the 25°C threshold for temperature stress (right vertical axis). (c) Daily mean salinity starting in July 2013. Solid black line indicates the mean for all aquaria. Gray shading around the mean line indicates daily maximum and minimum values. (d) Daily mean aragonite saturation calculated from salinity, alkalinity, pH, and temperature using CO2SYS. Solid black line indicates the mean for all aquaria. Gray shading around the mean line indicates daily maximum and minimum values. (e) Daily mean CO2(aq) calculated for each pH treatment from salinity, alkalinity, pH and temperature using CO2SYS. Solid lines indicate the means for each treatment. Pastel shading around each line indicates daily maximum and minimum values. Red line: pH 6.1, orange: pH 6.5, green: pH 6.9, blue: pH 7.4, black: pH 7.7 (ambient). (f) Hourly values of pH (open symbols) and CO2(aq) (filled symbols) of the ambient treatment (no CO2 addition) from 2 to 8 August 2014. Major ticks on the time axis indicate local midnight (Eastern Standard Time). PSS: Practical Salinity Scale; SW: seawater
values of salinity, alkalinity and pH, was consistently <1 (median $\Omega = 0.7$) but showed a high degree of daily and seasonal variability (min. $\Omega = 0.04$, max. $\Omega = 4.2$). Aragonite saturation was higher during the day ($\Omega = 0.96 \pm 0.01$) than at night ($\Omega = 0.60 \pm 0.01$), and more variable in summer ($\Omega$ daily range = 0.21 to 2.48) than in winter ($\Omega$ daily range = 0.15 to 1.40, Fig. 2d).

Ambient $\text{CO}_2(aq)$ concentrations in source water averaged $55 \pm 19 \mu\text{mol kg}^{-1} \text{SW}$, depending on season and time of day (Fig. 2e, black line). Prior to the onset of $\text{CO}_2$ enrichment on 1 June 2013, all aquaria experienced nearly identical variations in $[\text{CO}_2(aq)]$ as well as temperature, salinity, alkalinity and pH. The source-water pH was more variable and ~0.5 units lower during summer than winter, resulting in pre-dawn $[\text{CO}_2(aq)]$ as much as 10 times above air saturation (~15 µmol kg$^{-1}$ SW). $[\text{CO}_2(aq)]$ approached (Fig. 2f) and occasionally dropped below (data not shown) air saturation during the daytime, thereby increasing the effect of $\text{CO}_2$ limitation on photosynthesis of the plants in the unmanipulated aquaria. Despite natural fluctuations in ambient salinity, pH and $\text{CO}_2(aq)$ of the source water, the experimental $\text{CO}_2$ manipulation produced consistent $[\text{CO}_2(aq)]$ and pH values across the treatments throughout the duration of the experiment (Fig. 2e, compare colored lines). The 3 wk gap in $[\text{CO}_2(aq)]$ and aragonite saturation data during July and August 2014 resulted from a lightning strike that temporarily disabled the NI data logger. Although pH data were not recorded during this period, the pH controllers for all aquaria operated normally. The irradiance and CTD sensors were equipped with backup data loggers that prevented loss of those data during this period. Because temperature across the aquaria varied by <0.2°C, temperature data from the CTD was used to fill the July to August 2014 gap in recorded aquarium temperature.

Concentrations of macronutrients in Owls Creek were monitored weekly by the Virginia Aquarium staff as part of their water quality monitoring program. Although there is considerable variability in ambient macronutrient concentrations in the creek waters, the mean concentration of dissolved inorganic nitrogen ($\text{NO}_3^- + \text{NO}_2^- + \text{NH}_4^+$) was $11.3 \pm 0.6 \mu\text{M}$ and orthophosphate was $1.4 \pm 0.1 \mu\text{M}$ (W. M. Swingle unpubl. data), well above the levels required to saturate eelgrass growth (Zimmerman et al. 1987) but not high enough to cause eelgrass decline (Burkholder et al. 1992, Brun et al. 2002, van der Heide et al. 2008).

### Plant performance features

Shoot numbers per container, a measure of plant survival, remained stable across all treatments for 2 mo after the initial transplanting in May 2013, indicating no significant effects of aquarium location or systemwide transplant shock (Fig. 3a, heat map). Although the $\text{CO}_2(aq)$ treatments were initiated on 1 June 2013, the slopes of shoot number vs. $\log[\text{CO}_2(aq)]$ were not significantly different from 0 in June and July, indicating no detectable effect on shoot numbers during the first 2 mo of $\text{CO}_2(aq)$ enrichment (white symbols, Fig. 3a, Table 1). In August 2013, however, shoot numbers increased with $[\text{CO}_2(aq)]$, as indicated by the statistically significant positive slopes (repeated-measures ANCOVA p << 0.05, Table 1). By the end of summer 2013, shoot numbers of the 823 µM $\text{CO}_2(aq)$ (pH 6.5) treatment nearly doubled through vegetative propagation, while less than half of the originally transplanted shoots survived in the ambient $\text{CO}_2(aq)$ treatment. Shoot losses in the ambient $\text{CO}_2(aq)$ treatment began in July 2013 as water temperature peaked and continued throughout the winter of 2013 to 2014. In contrast, shoot numbers increased slowly via vegetative propagation through the fall and winter of 2013 in the 3 highest $\text{CO}_2(aq)$ treatments and then increased dramatically at the beginning of 2014 (winter). This dramatic increase in vegetative propagation coincided with the increase in daily solar irradiance following the winter solstice (Figs. 2a & 3a). The slope of shoot number vs. $\log[\text{CO}_2(aq)]$ was highest from January to May of 2014 (Fig. 3a, white symbols and lines). Senescence of the flowering shoots after seed ripening in June and July 2014 (Fig. 3b) caused a transient loss of shoot numbers, most evident at high $[\text{CO}_2(aq)]$ because of the high flowering rate, that subsequently recovered due to a new round of vegetative shoot propagation in the late summer and fall of 2014 (Fig. 3a). In contrast, vegetative shoot losses continued in the ambient $\text{CO}_2(aq)$ treatment throughout the summer and fall of 2014. Differentiation of vegetative shoots into flowering shoots began as early as January 2014 in the other $\text{CO}_2(aq)$ treatments and showed a strong, positive response to $[\text{CO}_2(aq)]$ throughout the flowering period that ended in June 2014, reaching 50% of the shoot population grown under the 2 highest $\text{CO}_2(aq)$ treatments during April and May 2014 (Fig. 3b, Table 1). In contrast, <10% of the surviving shoots flowered in the ambient $\text{CO}_2(aq)$ treatment, and the first flowering shoots did not appear until May 2014. Vegetative shoot size (cm$^2$ total leaf area) increased with $[\text{CO}_2(aq)]$ throughout the summer and
early fall 2013 period of warm water temperature (>20°C), decreased through the cold winter of 2014 and increased with the return of warm water in the summer of 2014 (Fig. 3c, Table 1).

Absolute growth rates responded positively within 6 wk of initiating the CO₂ treatment in June 2013, before differences in shoot survival or shoot size were statistically significant (Fig. 4a, Table 1). Absolute growth rates continued to respond in a positive, logarithmic fashion to the [CO₂(aq)] gradient throughout the summer and early fall, during which time water temperature consistently exceeded the previously established 25°C threshold for stress. Absolute growth rates declined in all treatments, and the CO₂(aq) effect (slope) decreased to near 0 during the winter period of low light availability and cold temperature. The CO₂(aq) effect, however, resumed as temperature increased during spring 2014 and continued throughout the warm summer of 2014. When normalized for plant size, the effect of [CO₂(aq)] on relative growth rate disappeared (slopes = 0), but the seasonal pattern of high summer growth and low winter growth remained strong (Fig. 4b, Table 1). Statistically significant dips in relative shoot growth (but not absolute growth) observed between March and June 2014 coincided with the maturation of flowering shoots that may place higher demand on carbon resources than vegetative shoots.

At the end of the 18 mo experiment, plants exposed to the highest CO₂(aq) treatments were about 3 times larger than the plants grown at ambient [CO₂(aq)] (Fig. 5a, Table 2). Further, the fraction of biomass allocated to aboveground leaves increased across the [CO₂(aq)] gradient such that the ratio of Shoot/(Root + Rhizome) was 50% higher in plants grown at the
highest \([\text{CO}_2(aq)]\) (Fig. 5b, Table 2). However, the distribution of biomass between roots and rhizomes remained constant across the \([\text{CO}_2(aq)]\) gradient (Fig. 5c, Table 2). Visual examples of the differences in surviving shoot numbers, plant size and biomass allocation across treatments are illustrated in Fig. 6.

Leaf sugar content also responded positively to higher \([\text{CO}_2(aq)]\) (Fig. 7a, Table 1). Differences across the \([\text{CO}_2(aq)]\) gradient were most pronounced during the summer of 2013, when leaf sugar concentration in the highest \([\text{CO}_2(aq)]\) treatment was 5-fold higher than the ambient treatment, and the slope across all treatments approached 200 \(\mu\text{mol sucrose (g}^{-1}\text{ fresh weight)(log[CO}_2(aq)]^{-1}}\). Sugar concentrations peaked in all \([\text{CO}_2(aq)]\) treatments during January and February 2014, when ambient temperature (Fig. 2b), rates of vegetative shoot proliferation (Fig. 3a) and leaf growth (Fig. 4a) were at a minimum, and flowering shoot differentiation was just beginning (Fig. 2b). Although the relationship between leaf sugar and \([\text{CO}_2(aq)]\) remained positive throughout the duration of the experiment, the slope declined during the summer of 2014 as vegetative shoot proliferation increased following the senescence of flowering shoots (Fig. 3a).

As with all other performance measures, leaf total chlorophyll (\(\text{chl} a + \text{chl} b\)) concentrations were identical across all treatments at the beginning of the experiment in May 2013 (Fig. 7b, Table 1). Unlike the other performance measures, however, total chlorophyll concentrations decreased in response to increasing \([\text{CO}_2(aq)]\) (Fig. 7b, Table 1). Although total chlorophyll increased in all treatments during winter in a manner reminiscent of photoacclimation, the slope remained significantly negative throughout the course of the experiment (Table 1). Total carotenoid content also decreased in response to increasing \([\text{CO}_2(aq)]\), and the slope remained significantly negative throughout the experimental period (Fig. 7c, Table 1).

### DISCUSSION

This experiment demonstrated that \(\text{CO}_2\) stimulation of primary production can enhance the summertime survival, growth, and proliferation of perennial eelgrass from the Chesapeake region that is regularly impacted by summer heat stress (Moore & Jarvis 2008, Jarvis et al. 2012, Moore et al. 2012). These results reinforce the emerging paradigm that eelgrass may benefit significantly from a high-\(\text{CO}_2\) world. Furthermore, the experiment demonstrated a logarithmic response to \([\text{CO}_2(aq)]\) in terms of shoot proliferation, size, growth, and sugar accumulation that was fundamentally consistent with model predictions based on meta-

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<td>&lt;0.001</td>
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<tr>
<td></td>
<td>(\text{pH})</td>
<td>1</td>
<td>48</td>
<td>21.29</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>Month × (\text{pH})</td>
<td>18</td>
<td>227</td>
<td>10.85</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Shoot size (cm(^2))</td>
<td>Month</td>
<td>18</td>
<td>200</td>
<td>2.97</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>(\text{pH})</td>
<td>1</td>
<td>90</td>
<td>97.39</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>Month × (\text{pH})</td>
<td>18</td>
<td>210</td>
<td>2.52</td>
<td>0.001</td>
</tr>
<tr>
<td>Absolute growth rate (cm(^2) d(^{-1}))</td>
<td>Month</td>
<td>18</td>
<td>203</td>
<td>5.93</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>(\text{pH})</td>
<td>1</td>
<td>80</td>
<td>72.22</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>Month × (\text{pH})</td>
<td>18</td>
<td>201</td>
<td>4.74</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Specific growth rate (d(^{-1}))</td>
<td>Month</td>
<td>18</td>
<td>178</td>
<td>1.63</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>(\text{pH})</td>
<td>1</td>
<td>104</td>
<td>0.05</td>
<td>0.827</td>
</tr>
<tr>
<td></td>
<td>Month × (\text{pH})</td>
<td>18</td>
<td>178</td>
<td>1.18</td>
<td>0.278</td>
</tr>
<tr>
<td>Sucrose ((\mu\text{mol g}^{-1}\text{ FW}))</td>
<td>Month</td>
<td>17</td>
<td>291</td>
<td>4.68</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>(\text{pH})</td>
<td>1</td>
<td>19</td>
<td>261.87</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>Month × (\text{pH})</td>
<td>17</td>
<td>291</td>
<td>3.83</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>(\Sigma\text{chl (a + b)}) (mg g(^{-1}) FW)</td>
<td>Month</td>
<td>14</td>
<td>252</td>
<td>2.98</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>(\text{pH})</td>
<td>1</td>
<td>18</td>
<td>144.44</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>Month × (\text{pH})</td>
<td>14</td>
<td>252</td>
<td>2.50</td>
<td>0.002</td>
</tr>
<tr>
<td>(\Sigma\text{Carotenoids}) (mg g(^{-1}) FW)</td>
<td>Month</td>
<td>13</td>
<td>234</td>
<td>1.58</td>
<td>0.093</td>
</tr>
<tr>
<td></td>
<td>(\text{pH})</td>
<td>1</td>
<td>18</td>
<td>138.69</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td></td>
<td>Month × (\text{pH})</td>
<td>13</td>
<td>234</td>
<td>1.61</td>
<td>0.083</td>
</tr>
</tbody>
</table>

Table 2. Regression statistics for the effect of \(\text{pH}\) on \(\Sigma\text{Plant biomass, Shoot/(Root + Rhizome) and Root/Rhizome at the end of the experiment in October 2014 shown in Fig. 5. FW: fresh weight.\)

<table>
<thead>
<tr>
<th>Dependent variable</th>
<th>Slope</th>
<th>SE</th>
<th>Intercept</th>
<th>SE</th>
<th>(r^2)</th>
<th>df</th>
<th>(F)</th>
<th>(p)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\Sigma\text{Plant biomass (g FW)})</td>
<td>-1.46</td>
<td>0.47</td>
<td>10.10</td>
<td>3.33</td>
<td>0.390</td>
<td>14</td>
<td>9.0</td>
<td>0.01</td>
</tr>
<tr>
<td>Shoot/(Root + Rhizome)</td>
<td>-1.86</td>
<td>0.64</td>
<td>17.23</td>
<td>4.54</td>
<td>0.388</td>
<td>14</td>
<td>8.9</td>
<td>0.01</td>
</tr>
<tr>
<td>Root/Rhizome</td>
<td>-0.12</td>
<td>1.05</td>
<td>3.19</td>
<td>7.48</td>
<td>0.001</td>
<td>14</td>
<td>0.016</td>
<td>0.90</td>
</tr>
</tbody>
</table>
The results of the experiment in the Chesapeake Bay showed that the increase in CO₂ availability eliminated the lethal effects of temperature on whole plant sugar accumulation, growth, and vegetative proliferation. At present, it is unclear whether growth under elevated [CO₂(aq)] prevented thermal stress directly or provided the shoots with sufficient carbon reserves to support growth.

Heat stress experiments performed on eelgrass populations from the Adriatic and Baltic Seas without a CO₂ subsidy at a maximum temperature of 25°C revealed significant up-regulation of genes coding for a variety of stress proteins in addition to significant shoot losses (Bergmann et al. 2010, Winters et al. 2011). Although this experiment produced similar shoot losses in the absence of a CO₂ subsidy, increasing CO₂ availability eliminated the lethal effects of temperature on whole plant sugar accumulation, growth, and vegetative proliferation. At present, it is unclear whether growth under elevated [CO₂(aq)] prevented thermal stress directly or provided the shoots with sufficient carbon reserves to support growth.
metabolic repair without negatively impacting growth and other performance features. The increased plant size and shoot number, however, suggests that the CO$_2$ effect was far greater than the simple suppression of stress, and for the eelgrass population examined here, the proximate effect of temperature was on metabolic carbon balance. There may be significant population-level differences in the ability of CO$_2$ to rescue eelgrass from thermal stress, as significant ge-

Fig. 5. (a) Total plant biomass (leaves + roots + rhizomes) at the end of the experiment in October 2014 plotted as a function of pH/CO$_2$(aq) treatment revealed a significant linear relationship (Table 2). (b) Fraction of total biomass allocated to aboveground leaves at the end of the experiment in October 2014 plotted as a function of CO$_2$ treatment revealed a significant linear relationship (Table 2). (c) Fraction of belowground biomass allocated between roots and shoots was not significantly affected by CO$_2$ treatment (Table 2). FW: fresh weight; SW: seawater

Fig. 6. Eelgrass growing at (a) 823 µM CO$_2$(aq), pH 6.5 and (b) 55 µM CO$_2$(aq), pH 7.7 (ambient) after 6 mo growth in the experimental chambers in October 2013. White bars at the top and bottom of the pictures are 20 cm long. (c) Differences in eelgrass shoot size across the CO$_2$ gradient in October 2014
netic and ecotypic diversity exists throughout eelgrass populations distributed throughout the Northern Hemisphere (Backman 1991, van Lent & Verschuure 1994, Olsen et al. 2004). In addition, the high organic carbon content of some sedimentary environments may subject seagrasses to increased vulnerability to sulfide anoxia, particularly at elevated temperature (Pedersen et al. 2004, Borum et al. 2005, Govers et al. 2014). However, CO2 stimulation of seagrass photosynthesis generates a proportional increase in belowground release of photosynthetic oxygen from the roots and rhizomes of seagrasses that may also mitigate vulnerability to sulfide intrusion (Bodensteiner 2006).

Sugar signaling has significant impacts on metabolism, growth, stress responses, and development from embryogenesis to senescence, including flowering (Rolland et al. 2006, Springer & Ward 2007), and provides a mechanism whereby [CO2(aq)] can affect the dynamics of flowering shoot production observed here. The onset of flowering in natural eelgrass populations has been reported to occur earlier in the year as latitude decreases and ambient water temperature increases (Phillips et al. 1983). In the California experiment (Palacios & Zimmerman 2007), flowering shoots first appeared in April in all CO2 treatments, and the flowering process continued throughout the summer. Even though winter water temperature was cooler by 6°C in the experiment presented here, flowering shoot initiation began here in January under high [CO2(aq)] and was completed by early July in all treatments, nearly 3 mo earlier than observed in the California experiment. Further, the flowering rate observed here (50% of the shoot population) was nearly double that observed in the prior experiment at high [CO2(aq)]. Although both experiments were conducted at the same latitude (36.8° N) and the pattern of incident irradiance was nearly identical at the 2 locations, the light intensity experienced by eelgrass in the California experiment (33% of surface irradiance) was half that used here. This difference in the daily pattern of light intensity may also be responsible for the higher flowering rates (50% of the shoot population at high [CO2(aq)]) observed here relative to the California experiment (25% at high [CO2(aq)]).
Nonetheless, the rates of flowering observed under elevated [CO₂(aq)] in both experiments were 3 to 5 times higher than the 10% levels typically observed in natural populations (Hemminga & Duarte 2000). Taken together, these results suggest that sugar accumulation may be critical in regulating the flowering process in eelgrass, while the direct effects of temperature are of secondary importance.

Sucrose accumulation resulting from elevated CO₂ availability can also result in the down-regulation of light-saturated photosynthetic capacity via expression of the Rubisco small subunit (Moore et al. 1999, Walters 2005). However, there was no significant down-regulation of pigment-specific photosynthetic capacity during the course of this experiment (Celebi 2016). Instead, the down-regulation of leaf pigment content under high [CO₂(aq)] observed here suggests that elevated CO₂ availability may be triggering photoaclimation mechanisms sensitive to the redox state of the electron transport chain and/or the production of reactive oxygen species resulting from CO₂-stimulated photosynthetic activity (Pfannschmidt et al. 2009).

Taken together, the experimental results presented here demonstrate that CO₂ availability can affect the temperature tolerance, vegetative growth, and flowering effort of Chesapeake region eelgrass via its impact on photosynthetic carbon fixation, sugar accumulation, and metabolic carbon balance. These results support previous model simulations that increased CO₂ availability associated with climate change may permit the survival of eelgrass in the Chesapeake region in spite of a warming climate (Zimmerman et al. 2015). Although it can be difficult to extrapolate the results of carefully controlled aquarium studies to the performance of organisms in the field, the experimental system employed here captured the full suite of natural estuarine variability in ambient temperature, solar irradiation, salinity, alkalinity and pH, strengthening our confidence in the reality of the responses observed here. However, even these aquarium experiments fall short of the ecosystem-level realism afforded by free-air CO₂ experiments (Ainsworth & Long 2005, Leakey et al. 2009). The technology to perform similar free-ocean CO₂ experiments is improving, but many challenges remain to its successful implementation, particularly for long-term, multifactor experiments (Gattuso et al. 2014). In the meantime, progress in understanding the impacts of climate change on marine ecosystems is likely to depend heavily on a combination of controlled experiments, numerical modeling, and field observations capable of integrating natural variability and scaling the results across time and space (Andersson et al. 2015).

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