FEATURE ARTICLE

Individual and population level effects of ocean acidification on a predator–prey system with inducible defenses: bryozoan–nudibranch interactions in the Salish Sea

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ABSTRACT: Ocean acidification (OA) from increased oceanic CO₂ concentrations imposes significant physiological stresses on many calcifying organisms. OA effects on individual organisms may be synergistically amplified or reduced by inter- and intraspecies interactions as they propagate up to population and community levels, altering predictions by studies of calcifier responses in isolation. The calcifying colonial bryozoan Membranipora membranacea and the predatory nudibranch Corambe steinbergae comprise a trophic system strongly regulated by predator-induced defensive responses and space limitation, presenting a unique system to investigate OA effects on these regulatory mechanisms at individual and population levels. We experimentally quantified OA effects across a range of pH from 7.0 to 7.9 on growth, calcification, senescence and predator-induced spine formation in Membranipora, with or without waterborne predator cue, and on zooid consumption rates in Corambe at Friday Harbor Laboratories, San Juan Island, WA. Membranipora exhibited maximum growth and calcification at moderately low pH (7.6), and continued spine formation in all pH treatments. Spines reduced Corambe zooid consumption rates, with lower pH weakening this effect. Using a spatially explicit model of colony growth, where colony area serves as a proxy for colony fitness, we assessed the population-level impacts of these experimentally determined individual-level effects in the context of space limitation. The area-based fitness costs associated with defense measured at the individual level led to amplified effects predicted for the population level due to competition. Our coupled experimental and modeling results demonstrate the need to consider population-level processes when assessing ecological responses to stresses from changing environments.

KEY WORDS: Predator–prey interactions · Ocean acidification · Inducible defenses · Space competition · Modeling · Membranipora membranacea · Corambe steinbergae

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INTRODUCTION

Ocean acidification (OA) resulting from increased oceanic uptake of atmospheric carbon dioxide presents a variety of environmental stresses to marine organisms and ecosystems (Hofmann et al. 2010, Kroeker et al. 2013). OA causes a decrease in both pH and carbonate ion availability (Feely et al. 2009), often with negative physiological implications for many organisms (Kroeker et al. 2013). Global pH levels have been declining, with continued projections of 0.3−0.4 units within the next century (Feely et al. 2004, 2009). Coastal areas, however, often experience natural pH fluctuations that currently exceed these predicted changes (Doney 2010, Hofmann et al. 2010). In the Salish Sea and San Juan Archipelago, WA, USA, naturally low seawater pH levels result from regular upwelling and river input (Murray et al. 2015). While open ocean pH averages well above 8.0 (Doney 2010), the San Juan Archipelago experiences pH around 7.8 and can reach as low as 7.67 in San Juan Channel during seasonal upwelling (Sullivan 2012, Murray et al. 2015). Consequently, the Salish Sea is a natural laboratory to investigate OA effects where organisms already experience pH values that open ocean organisms will likely not experience until the end of the century.

Organisms that make shells or skeletons from CaCO₃ are common in the Salish Sea, with a third of Puget Sound species identified as calcifiers (Busch & McElhany 2017). Calcifying organisms have been demonstrated to be generally vulnerable to acidification (Kroeker et al. 2013). Decreasing availability of CO₂²⁻ can impede calcification (Feely et al. 2009). In conjunction, the dissolution of calcified structures is favored by the low-saturation states of CaCO₃ polymorphs, calcite and aragonite, that often accompany OA (Orr et al. 2005). Calciﬁer responses to OA have been generally negative, with many organisms exhibiting decreased growth, survival and calcification rates (Kroeker et al. 2013). Some taxa, however, have displayed no effect or even positive effects, demonstrating the importance of quantifying species-specific responses in diverse taxa (Ries et al. 2009). Variable OA responses have been attributed to a number of potential causes, including diversity of calcification mechanisms, presence and thickness of protective organic membranes and solubility of different CaCO₃ polymorphs in addition to differences in capacity for plasticity and prior exposure to low pH (Ries et al. 2009).

While most studies document OA effects on calcifiers in isolation, the true ecological consequences of these stresses occur within interconnected species networks that ultimately structure populations and communities. Skeletal mineralogy alone is an incomplete predictor of OA sensitivity, especially in the context of trophic interactions where direct and indirect effects associated with species interactions may be significant (Busch & McElhany 2017). Physiological and behavioral processes in both predators and prey may be impacted differently by OA. These alterations potentially amplify or reduce organism-level effects of OA on population and community levels, highlighting the importance of considering predator-prey interactions when conducting OA studies (Kroeker et al. 2014). For example, OA may impair predator detection mechanisms, resulting in reduced predator avoidance behavior in fish (Dixson et al. 2010) and marine snails (Manriquez et al. 2013), but can reduce predation by crabs on oysters (Dodd et al. 2015). In these examples, the differential effects of OA on distinct trophic levels indicate a potential for environmental stress to impact populations and communities to different extents and in different directions than suggested by studies of species in isolation. The potential of OA to alter predator-prey interactions, and how these effects may propagate to populations and communities, however, remains poorly understood.

Inducible defenses are an important subset of predator-prey interactions that confer protection to organisms upon the detection of environmental cues of impending predation (Harvell 1990). When defenses incur significant costs, inducibility allows organisms to maximize the benefit of being defended when under attack while avoiding the cost of defense when predation risks are low (Tolrian & Harvell 1999, Ferrari et al. 2010). Predator-induced defenses are common in marine calcifiers, e.g. shell thickening in oysters (Lord & Whitlatch 2012, Scherer et al. 2018), snails (Trussell & Nicklin 2002, Bourdeau 2010), and mussels (Leonard et al. 1999). OA has been found to affect inducible defenses in some species. Both induced shell thickening in the snail Littorina obtusata (Bibby et al. 2007) and predator-induced neck teeth in freshwater Daphnia spp. were reduced by high partial pressure of carbon dioxide (pCO₂) (Weiss et al. 2018). While these studies have investigated the effects of acidification on inducible defenses, few have done so in the context of multiple levels of biological organization.

Inducible defenses can have effects beyond the organism level, with potential propagating effects through food webs and population dynamics of both predators and prey (Miner et al. 2005), which suggests that OA effects on these interactions must
then be assessed at population and community levels. Scaling organism-level interactions to estimate population-level responses can provide quantitative assessments of propagating effects of OA. In this study, we used a well-known inducible defense system to develop a strategy for assessing OA effects in which we empirically measured individual-level responses and used a numerical model to infer population-level effects.

The encrusting bryozoan Membranipora membranacea (hereafter Membranipora) is an abundant colonial epibiont on kelp in the Salish Sea and San Juan archipelago (Fig. 1). Larval recruitment occurs in late spring, with populations colonizing kelp blades as they grow throughout the summer (Seed 1976). Colonies do not typically overgrow each other, and instead divert energy to reproduction when occluded (Harvell & Padilla 1990). Membranipora are ecologically significant invasive biofouling organisms along the east coast of North America, where overgrowth on kelp can cause kelp deforestation (Scheibling & Gagnon 2009, Saunders et al. 2010), especially where specialist nudibranch predators are not prevalent (Lambert et al. 1992) and other predators prefer different bryozoans (Pratt & Grason 2007). Membranipora colonies consist of multiple, genetically identical zooids (Harvell 1984) partially encased in a calcium carbonate skeleton (Smith et al. 2006, Taylor et al. 2015). Larger colonies consist of more zooids, leading to higher reproductive output than colonies with fewer zooids, making space an important limiting resource for Membranipora (Yoshioka 1982a). The cryptic nudibranch Corambe steinbergae (hereafter Corambe) exerts strong episodic predation pressure on Membranipora, feeding by sucking individuals out of zooids, leaving them empty (Yoshioka 1982b). In response, Membranipora exhibits an inducible defense by forming protective chitinous spines upon detecting waterborne chemical cues from Corambe. Spines are produced only on newly formed zooids at the growing margin of the colony, requiring the formation of new calcified zooids (Harvell 1984). Spination begins with formation of corner spines on zooid vertices, followed by additional spines along the walls of the zooids (Harvell 1984). Spines have been demonstrated to be effective in reducing predation (Harvell 1986). However, spines also have associated costs, including reduced colony growth as resources are allocated for defense (Harvell 1986, 1992) and disruption of feeding currents (Grünbaum 1997).

The Membranipora–Corambe system presents a uniquely informative opportunity to understand the consequences of OA effects and predator–prey interactions across multiple scales of biological organization. Membranipora colonies are primarily mineralogically composed of aragonite and low-Mg calcite (Smith et al. 2006), which may affect their ability to grow and calcify in OA conditions. Aragonite is a highly soluble polymorph of CaCO₃, and some heavily calcified bryozoans have been identified as potential bioindicators of OA given their vulnerability (Smith 2009, Fortunato 2015, Taylor et al. 2015). Membranipora is a tractable experimental system, as its modular architecture allows for a single genotype to be divided into multiple experimental treatments. This effective control for genetic variation facilitates quantifying the plastic capabilities of specific genotypes (Harvell 1990). Both the deployment of defense (inducible spine formation) and the degree of predation (consumed zooids) are easily and non-destructively quantifiable. Finally, organizational levels of this system, from zooid to population, are inherently linked by zooid-level processes such as spine formation, which alter colony growth dynamics and ultimately influence space competition and, consequently, fitness.

Fig. 1. Tightly linked multiple levels of biological organization in the bryozoan Membranipora membranacea make it a uniquely informative organismal system in which to investigate OA effects at multiple levels. Individual zooids within a colony: (a1) feeding, undefended zooids (with no spines), and (a2) spines formed along zooid walls of defended zooids. (b) One Membranipora colony comprising many genetically identical zooids with older, developed zooids at the center and younger developing zooids along the perimeter. (c) Multiple colonies competing for space on a kelp blade, with larger colonies occluding smaller colonies. From left to right scale bars are 300 μm, 250 μm, 3 mm and 5 cm.
Membranipora presents a good system for constructing population dynamics models because space competition and reproductive potential, mediated by variations in predation and colony growth rates, can be represented as a spatially explicit process analogous to observable distributions in the field. Using this system to explore connections between individuals and populations, modeled growth patterns which scale up individual responses can provide some insight on OA impacts on predicted area-based fitness and costs of defense under space limitation. Therefore, Membranipora presents an exceptional opportunity to understand population-level implications of an organism-level anti-predator mechanism under OA stress.

The goals of this study were to (1) experimentally quantify the colony-level effects of OA on Membranipora growth, CaCO₃ production and formation of defensive spines, along with rate of predation by Corambe, and (2) use modeling approaches to infer the consequences of observed colony-level impacts at the population level. We addressed the following questions: (1) How does a range of pH reflecting mild to severe OA affect Membranipora colony growth, CaCO₃ production and inducible spine formation and Corambe predation rates on Membranipora? (2) How are predicted organism-level effects of OA and predator cues altered by the inclusion of population-level processes (e.g. space competition among Membranipora colonies)?

We hypothesized that, at the colony level, growth and CaCO₃ production would be reduced by OA in the absence of predator cues, and that reductions in growth would be compounded in the presence of predator cue due to the cost of defense. On the population level, we predicted the associated costs of these growth rate reductions would be further amplified by space competition. We also hypothesized that inducible spines, although chitinous, would be reduced by OA since they form on areas of new growth which require the formation of new calcified skeleton, and that predation rates would be reduced due to direct effects of OA stress on organisms.

MATERIALS AND METHODS

Bryozoan collection and preparation

Membranipora colonies were collected off the floating dock at Friday Harbor Laboratories (FHL) (48.546° N, 123.013° W) in late May and transferred to laboratory sea tables reflecting collection site conditions. Colonies with at least a 5 cm radius of free space were selected as target colonies and cut from the plastic plate using a bandsaw, to produce 10 × 10 cm plates with 1 colony at the center. To minimize shock, colonies were returned to sea tables for 3 d. Other colonies surrounding the target colony were cleared from the plastic plate to prevent interference. This allowed sufficient area for new growth throughout the duration of the experiment, such that in no colony was space limited. Target colonies were dyed with a 200 mg l⁻¹ solution of Alizarin Red S (Sigma Aldrich) in filtered seawater for 18 h and flushed for 6 h, to mark the initial colony area (Saunders & Metaxas 2009). 10 × 10 cm plastic plates with target colonies were scored with a razor blade, broken into separate quarters and returned to sea tables for 24 h prior to the experiment. A total of 24 colonies (distinct genotypes) were included in the experiment. Twelve colonies were exposed to water-borne predator cue of Corambe, and 12 received no predator cue, with each colony genotype fully replicated (roughly equal colony sections) across 4 nominal pH treatments: 7.0, 7.3, 7.6 and 7.9 (Fig. S1 in Supplement 1 at www.int-res.com/articles/supp/m607p001_supp/). We used regression-based approaches to establish Membranipora response curves, which could be easily integrated into a population dynamics model.

Experimental setup

In total, 4 tanks were used in the FHL Ocean Acidification Environmental Laboratory (OAEL) to establish each of these pH conditions by bubbling CO₂ as described by O’Donnell et al. (2013). Temperature in tanks was maintained at 13°C, reflecting typical ambient temperature at the collection site during the summer months (Murray et al. 2015). Weekly water samples were taken for carbonate chemistry measurements including spectrophotometric pH (total scale) and total alkalinity in accordance with best practices (Reibesell et al. 2010). CO₂SYS v.1.05 (Pierrot et al. 2006) was used to calculate additional carbonate system parameters from measured pH, total alkalinity, salinity and temperature (Table 1). Temperature and pH were monitored continuously throughout the experiment using Durafet electrodes, which were calibrated weekly. Tanks included a total of 8 isolated chambers, 6 of which were used to contain 4 Mem-
**Table 1.** Average (±1 SD) carbonate chemistry data for experimental seawater tank treatments at the Ocean Acidification Environmental Laboratory (OAEL). Total alkalinity (TA) and salinity were measured weekly and pH (total scale) in tank chambers was monitored daily throughout the study. Additional carbonate system parameters (dissolved inorganic carbon [DIC], partial pressure of carbon dioxide [pCO$_2$], aragonite saturation state [Ω$_{Ar}$]) were calculated using CO2Sys (Pierrot et al. 2006) with dissociation constants (K1, K2) from Mehrbach et al. (1973) refit by Dickson & Millero (1987), HSO$_4$ constant from Dickson (1990) and Boron constant from Lee et al. (2010).

<table>
<thead>
<tr>
<th>pH</th>
<th>Temp (°C)</th>
<th>Salinity (psu)</th>
<th>pH (total scale)</th>
<th>TA (μmol kg$^{-1}$)</th>
<th>DIC (μmol kg$^{-1}$)</th>
<th>pCO$_2$ (μatm)</th>
<th>Ω$_{Ar}$</th>
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</thead>
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<td>7.0</td>
<td>12.8 ± 0.1</td>
<td>28.6 ± 0.1</td>
<td>7.13 ± 0.06</td>
<td>2022 ± 19</td>
<td>2141 ± 32</td>
<td>3462 ± 504</td>
<td>0.23 ± 0.03</td>
</tr>
<tr>
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<td>29.2 ± 0.1</td>
<td>7.38 ± 0.02</td>
<td>2024 ± 25</td>
<td>2060 ± 27</td>
<td>1937 ± 94</td>
<td>0.44 ± 0.02</td>
</tr>
<tr>
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<td>28.9 ± 0.1</td>
<td>7.66 ± 0.03</td>
<td>2022 ± 19</td>
<td>1984 ± 21</td>
<td>986 ± 74</td>
<td>0.81 ± 0.05</td>
</tr>
<tr>
<td>7.9</td>
<td>13.2 ± 0.1</td>
<td>29.0 ± 0.3</td>
<td>7.88 ± 0.02</td>
<td>2023 ± 20</td>
<td>1920 ± 21</td>
<td>589 ± 28</td>
<td>1.30 ± 0.05</td>
</tr>
</tbody>
</table>

*branipora* colony sections each. Within each tank, 3 chambers received water pre-treated with predator cue from a separate chamber which housed *Corambe*, to induce defensive spines without actual predation. The predator chambers contained 12 nudibranchs for a final concentration of 4 nudibranchs l$^{-1}$, exceeding the threshold concentration required to induce a defensive response (Harvell 1986). Surface area of each nudibranch was measured prior to the experiment, and distributed such that each predator chamber had 12 similarly sized nudibranchs with a total surface area of approximately 130 mm$^2$. Another 3 chambers served as controls, receiving water not treated with predator cue. All chambers received source water from the main tank with independent inflow and outflow lines such that chambers did not share water, which allowed us to restrict predator cue to only predator cue treatments. However, we potentially experienced minor contamination of predator cue in control treatments due to occasional overflows of predator cue chambers into the main water source, which may have created diffuse predator cue water for control treatments. Four colony sections were randomly assigned to each tank chamber, and rotated among chambers within a treatment every 2 d to control for chamber effects. One predator-exposed colony section was damaged during the experiment and all corresponding sections of that colony were removed. The flow-through system was plumbed with unfiltered seawater and each chamber was fed 50 000 cells ml$^{-1}$ of *Isochrysis galbana* and 5000 cells ml$^{-1}$ of *Dunaliella* sp. 3 times daily to encourage bryozoan growth.

**Part 1: Bryozoan growth, senescence and inducible defense formation**

A paddle system constructed from PVC pipe and a motor (Strathmann 2014) above each OAEL tank generated mixing in each chamber sufficient to keep algal cells suspended to ensure effective *Membranipora* feeding. Colony sections on plastic squares were attached to swinging paddles using rubber bands, and exposed to experimental treatments for 15 d. Growth and zooid senescence were monitored for each colony section using photographs taken every 5 d with a Nikon D3400 camera with macro lens. Colony section surface area was measured in ImageJ. Colony section growth rates were calculated as the exponential rate of increase in surface area, because colony area increases roughly in proportion to the number of feeding zooids. Growth rates were calculated by solving for $r$ using the equation below, where $A_{t_{15}}$ is the colony surface area after 15 d, $A_{t_0}$ is the initial colony surface area and $t$ is the duration of the experiment (15 d):

$$A_{t_{15}} = A_{t_0} e^{rt} \tag{1}$$

Zooids in photos were classified as feeding (fully developed), developing, or degenerated (brown body), and counted using the ImageJ cell counter plugin. Fully developed zooids had a full gut (see center zooids in Fig. 1b) while developing zooids could not yet feed and did not yet have a developed gut (see peripheral zooids in Fig. 1b). Degenerated zooids referred to senesced zooids, not those consumed via predation. To assess spine length and quantity, a Lumenera Infinity microscope camera mounted on a dissecting microscope was used to take photo transects for each colony section. Colonies were held at a 45° angle for photographing using a 3D printed mount and a micromanipulator. For each colony section, 3 uniform transect lines, 1 cm apart, were photographed through the new growth area and stitched together. The length of the closest corner spine to each of 5 randomly selected locations along each transect line was measured and corrected for the photographing angle.
Part 2: Nudibranch feeding rates

After colonies received a 15 d predator cue or control treatment and growth rates were measured in the absence of predation, 1 nudibranch was released on each colony section in each pH condition to determine feeding rates. Nudibranchs were starved and acclimated for 24 h in their respective feeding pH treatments before being placed on *Membranipora* colony sections. We measured nudibranch length (diameter in mm from photographs), sorted nudibranchs from shortest to longest and paired them with colony sections such that the longest nudibranch was placed with the colony section of the largest surface area. Nudibranch lengths ranged 2.52−8.73 mm with a mean length of 5.30 mm. Initial photographs of colonies were taken to establish baseline conditions of zooids. All nudibranchs were initially placed on the center of the target colony’s growing margin and confined by adding 300 μm mesh cages with a 3D printed frame cable-tied to the swinging plastic paddles above each colony to prevent predation on colonies other than the target colony. Nudibranchs were allowed to consume zooids for a 24 h period, after which final photographs were taken and the number of empty zooids, from predation, were quantified.

Part 3: Quantification of bulk calcium content in bryozoan skeletons using mass spectrometry

One skeleton sample was taken from the new area grown during the experiment from each colony section. Skeleton samples were roughly 100 mm² but were variable depending on available area of new growth in each colony section. Samples were transferred to a 35 mm Petri dish, and sequentially rinsed with milliQ water, a 10% bleach solution, a final milliQ rinse and 95% ethanol to dry out the sample. Areas of skeleton samples were measured in ImageJ. Each rinsed and dried skeleton sample was prepared in 2 parts (A and B) for elemental analysis to quantify Ca content as a proxy for bulk skeletal CaCO₃ (similar to methods in Wood et al. (2008) and Findlay et al. (2011) that quantified total Ca). For Part A, skeletal samples were removed from their respective Petri dishes, weighed on a microbalance, dissolved in 2.5 ml of 5% HNO₃ and diluted to a final 5 ml with milliQ water in a 15 ml falcon tube. For Part B, residual skeletal material was rinsed into a 2.5 ml 5% HNO₃ solution and diluted to a final 5 ml solution with milliQ water. A 10 ml solution composed of 1 ml Part A or B, 100 μl of 100 ppb Sc, Rh, and Y (used as an internal standard spike), and 8.9 ml 2% HNO₃ was analyzed using an inductively coupled plasma mass spectrometer (ICP-MS; Thermo model iCAP RQ) for bulk Ca determination. Samples were analyzed for ⁴³Ca and ⁴⁴Ca. Raw elemental intensities in counts s⁻¹ were blank-corrected, further corrected for ⁸⁷Sr²⁺ interference and instrumental drift with internal standardization before conversion to concentrations (ppm). ⁴³Ca intensities were used to calculate final Ca concentrations and were converted to total Ca (mg) using exact dilution factors for each sample, and normalized to either weight of skeleton (Part A), as an assessment of skeletal composition, or to area of the sample (Parts A + B), as a measure of skeletal density.

Statistical analyses (Parts 1−3)

For each experimental response variable, a series of linear or generalized linear mixed effects models were used to test pH and predation cue as fixed effects and included colony as a random effect to account for variation across genotypes. Quadratic relationships, using pH² as a fixed effect, were also tested for the response variables growth rate, zooid senescence and skeletal density, which displayed non-linear trends with respect to pH. Akaike’s information criterion (AIC) was used to determine the best fitting model for all analyses (see Supplement 1). For growth rate analysis, linear mixed effects models with identity link functions and Gaussian response distributions tested main effects of pH, pH² and predation cue. Zooid senescence was presented as a proportion of degenerated zooids relative to total colony zooids, and generalized linear mixed effects models with logit link functions and binomial response distributions tested main effects of pH, pH² and predator cue. For spine length analysis, linear mixed effects models with identity link functions and Gaussian response distributions with an additional nested random effect of colony section within colony tested for fixed effects of pH and predator cue. For Corambe zooid consumption rate analysis, generalized linear mixed effects models with log link functions and a Poisson distribution tested the main effects of pH and previous colony predator exposure (defended vs. undefended colonies) with nudibranch length as a covariate and colony, nudibranch and colony size included as random effects. For skeletal density analysis, linear mixed effects models with an identity link function and a Gaussian distribution
tested the fixed effects of pH, pH$^2$ and predation cue. Statistical analyses were conducted in R v.3.4.1 (R Core Team 2017) using the ‘lme4’ package (Bates et al. 2014) to construct mixed effects models.

**Part 4: Integrating experimental data into bryozoan population dynamics model**

A spatially explicit population dynamics model was constructed to investigate the implications of empirically determined colony growth rates for *Membranipora* population structure, distributions and space competition. The model approximates *Membranipora* colonies as aggregations of repeating hexagons representing zooids. These hexagons, with side length $p$ and acute angle $\theta$, fit together in an approximately circular form if $\pi/\theta$ is an integer (Fig. 2) (Grünbaum & Shephard 1987). Modeled zooids are not exact replicates of zooid geometry, but are the most similar polygons that can fully tile the plane without gaps or alterations in shape.

The model accounts for the key processes of energy acquisition by mature feeding zooids, translocation of that energy to the growing edge to support formation and development of new zooids and occlusion by neighboring colonies preventing new zooid formation. Colony growth is mediated through zooid-level algorithms for energy translocation. A system of ordinary differential equations is used to describe changes in zooid energy content, mass and developmental state over time. The rate of change of energy content of the $j^{th}$ zooid is defined as:

$$\frac{dE_{t_{oj}}}{dt} = q_{in_{j}} + q_{f_{j}} - q_{out_{j}} - q_{g_{j}} - q_{m_{j}}$$  \hspace{1cm} (2)

where $E_{t_{oj}}$ is the total available energy content of zooid $z_{j}$; $q_{in_{j}}$ is the rate of energy translocated into zooid $z_{j}$ from upstream neighboring zooids; $q_{out_{j}}$ is the rate of energy translocated out of zooid $z_{j}$ to downstream neighbors; $q_{f_{j}}$ is the rate of energy intake from feeding; $q_{g_{j}}$ is the rate of energy allocated for the production of zooid tissue; and $q_{m_{j}}$ is the rate of energy allocation for basal metabolism. All rates have units of J d$^{-1}$. Additional equations and parameters specifying how each term is calculated are presented in Text S1 & Table S1.

**Model parameters**

The model, implemented in Python v.2.7, establishes a 2D domain which simulates a kelp blade randomly seeded with *Membranipora* colonies that grow and compete for space. Colony growth rate is not an explicitly designated parameter in the model, but is an emergent property of energy and growth dynamics within colonies. Model parameters were adjusted to achieve the correct range of growth rates determined from experiments (see Table 2). Model colony growth rates were calculated using the slope of log-transformed colony area versus time between Days 20 and 35 of the simulation, corresponding to the method by which growth rates were experimentally determined. In the model, colony growth rates were driven by the combined parameter $\Delta q = q_{f} - q_{m}$ i.e. the difference between energy input from feed-
ing and basal metabolic rate. Therefore, we adjusted Δq to generate simulated growth rates for several single, unoccluded colonies (Fig. 3). We then linearly interpolated across calculated Δq values to find values resulting in growth rates that matched experimentally determined values for each pH and predator condition (see Table 2).

We used model simulations to test consequences of space limitation and competition on the cost of defense and OA effects. These simulations contained 16 simultaneously settled colonies. We used 2 types of simulations, one set with all 16 colonies growing with the defended colony growth rate or undefended colony growth rate at each pH (10 replicate runs), and a second set of simulations containing 8 colonies with pH-specific undefended colony growth rates and another 8 colonies with the defended colony growth rates (25 replicate runs). Simulations were 75 d long with 0.5 d time step intervals, approximating the length of *Membranipora* growing season. Model domain dimensions were 50 × 100 mm, a smaller scaled estimate of typical kelp blade area, with typical settlement densities (Seed 1976, Yoshikoa 1982a). Smaller domain sizes allowed us to acquire similar space competition data without the computational expense required for larger domain sizes representative of actual kelp blade sizes. Colony areas at each time step were averaged to account for edge effects, and plotted as growth trajectories over 75 d. Growth trajectories from both types of simulations were compared to establish the cost of defense in the context of competition with colonies of identical growth rates, versus competition between colonies of different growth rates.

Using average colony growth trajectories over time, we calculated 2 metrics to compare the cost of defense in simulations with colonies of identical growth rates versus simulations with undefended and defended colony competition. We defined the intra-population cost of defense as the difference in mean colony area from populations of uniform growth rate, corresponding to either all defended or undefended colonies, at the time when undefended colonies have filled the domain. We defined a full domain at the time at which the increase in mean colony area decreased below 0.5 mm². We defined the inter-population cost of defense as the difference in mean colony area between undefended and defended colonies at the end of a 75 d duration, from model simulations where equal numbers of both colony types competed against each other for space. These cost metrics were used as area-based fitness proxies for each pH.

**RESULTS**

**Part 1a: Bryozoan growth rates**

*Membranipora* colony sections grew in all treatments (mean ± SD growth rate = 0.076 ± 0.019 d⁻¹, n = 92; Table 2, Fig. 4a). Predator-exposed colonies exhibited slower growth rates compared with control colonies (predator cue: p = 0.0003), representing the cost of defense; however, this effect depended on pH (pH × predator cue: p = 0.0003). The reduction of

<table>
<thead>
<tr>
<th>pH</th>
<th>Predator treatment</th>
<th>Mean Δq (J d⁻¹)</th>
<th>SD</th>
<th>n</th>
<th>Model Δq (J d⁻¹) used for</th>
<th>Mean</th>
<th>SD</th>
<th>n</th>
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<tr>
<td>7.9</td>
<td>Control</td>
<td>0.087</td>
<td>0.013</td>
<td>12</td>
<td>0.852</td>
<td>0.871</td>
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<tr>
<td></td>
<td>Predator-exposed</td>
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<td>0.017</td>
<td>11</td>
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<td>0.071</td>
<td>0.017</td>
<td>11</td>
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<tr>
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<td>0.014</td>
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<td>0.014</td>
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<td></td>
<td>Predator-exposed</td>
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<td>0.015</td>
<td>12</td>
</tr>
<tr>
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<td>0.012</td>
<td>11</td>
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<td>0.012</td>
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<td>12</td>
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<td>0.016</td>
<td>11</td>
<td>0.402</td>
<td>0.065</td>
<td>0.016</td>
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Table 2. Average *Membranipora* growth rates for each treatment determined by experiments which were later replicated in a spatially explicit bryozoan population model. Corresponding Δq values (i.e. the difference between energy input from feeding and energy output to basal metabolic rate) used to replicate specific treatment growth rates in the model are listed (see Fig. 3).

![Fig. 3. Relationship between Δq (difference between energy input from feeding and energy output to basal metabolic rate) and *Membranipora* colony growth rate in modeled colonies. This plot shows simulated growth rates in single, unoccluded (non-space limited) colonies across a range of Δq values. Other parameters are fixed as described in Table S1](image-url)
growth rates in predator-exposed colonies was observed in all pH treatments except 7.0 (Fig. 4a). Growth rates responded to pH treatments parabolically (pH²: p < 0.0001, pH: p < 0.0001), with the shapes of the curves dependent on predator cue (pH² × predator cue: p = 0.0004). Both inferred growth peaks occurred at lower than ambient pH: 7.63 (control colonies) and 7.59 (predator-exposed colonies). Control colonies exhibited faster growth rates than predator-exposed colonies at pH 7.9, 7.6 and 7.3 (0.016, 0.022 and 0.013 d⁻¹ faster respectively), with a peak at pH 7.6. At pH 7.0, model trends showed growth rates were substantially reduced in both cue and no cue treatments (Fig. 4a). Average growth rates presented in Table 2 were used to inform parameter inputs for model simulations. Complete mixed effects model analysis and best fit model output are presented in Tables S2.1 & S2.2 in Supplement 1.

**Part 1b: Zooid senescence**

Zooid senescence was quantified as the number of degenerated zooids formed during the experiment as a proportion of the total zooids in the colony section (mean ± SD proportion degenerated = 0.14 ± 0.18, n = 92; Fig. 4b). pH affected zooid senescence trends (pH: p < 0.0001), but was dependent on predator cue (predator cue × pH: p < 0.0001). Both control and predator-exposed colonies exhibited parabolic senescence patterns (pH²: p < 0.0001), demonstrating dramatic increases in senescence at pH 7.0 with curves specific to predator-exposure treatment (pH² × predation cue: p < 0.0001). Control colonies expressed a 2.16x greater proportion of degenerated zooids than predator-exposed colonies only at this lowest pH level; senescence rates were similar across higher pH values (Fig. 4b). Senescence trends reflected the complement of growth rate trends observed in both
Part 1c: Inducible defense formation

Spines were formed in all treatments (mean ± SD length = 76.33 ± 71.94 μm, n = 1350), but were 117.43 μm longer on average in predator cue treatments than in control treatments (Fig. 4c). Spines were produced in some control colonies (mean length = 19.79 μm) but were significantly shorter than those produced by predator-exposed colonies (predator cue: p < 0.0001). A weak but significant effect of pH was detected for both control and predator-exposed colonies, indicating that spine length at lower pH was slightly decreased (pH: p = 0.01). Complete mixed effects model analysis and best fit model output are presented in Tables S4.1 & S4.2. Abundance of spines per treatment is presented in Fig. S2 with corresponding analyses in Tables S5.1 & S5.2.

Part 2: Nudibranch consumption rates

Consumption rates of Membranipora zooids by Corambe were measured as the number of zooids consumed over a 24 h period (mean ± SD consumption = 74.29 ± 49.69 zooids, n = 92; Fig. 5). Spines deterred feeding in nudibranchs (spine presence: p = 0.022) but this effect was dependent on pH, with reduced zooid consumption rates in higher pH treatments but not in lower pH treatments (spine presence × pH: p = 0.016). As pH decreased, zooid consumption rates on control colonies decreased, while consumption rates on predator-exposed colonies increased (Fig. 5), essentially removing the benefit of defense. Nudibranch length also significantly influenced zooid consumption rate, with larger nudibranchs consuming more zooids (nudibranch length: p = 0.001). Complete mixed effects model analysis and best fit model output are presented in Tables S6.1 & S6.2.

Part 3: Bulk calcium content quantification of bryozoan skeletons using ICP-MS

Total mg Ca sample⁻¹ was calculated and normalized to skeletal area as a metric of skeletal density (Fig. 4d). Skeletal density (mean ± SD density = 0.0023 ± 0.0007 mg Ca mm⁻², n = 90) was significantly affected by pH (pH: p < 0.0001) and displayed a parabolic trend (pH²: p < 0.0001) with a weak peak at target pH 7.6. This peak pH was similar to pH values associated with maximum growth (control: 7.63; predator-exposed: 7.59). There was no significant difference between skeletal density of control and predator-exposed colony sections. Complete mixed effects model analysis and best fit model output presented in Tables S7.1 & S7.2. Bulk calcium content normalized to skeletal weight is presented in Fig. S3 with corresponding analyses in Tables S8.1 & S8.2.

Part 4: Membranipora population dynamics model results

Model output (Fig. 6) enabled visualization and quantification of predicted population-level costs of defense at different OA conditions. The cost of defense was dependent on space competition between colonies and the variations in their growth rates. In simulations where all colonies grew with identical growth rates (either defended or undefended), all colonies eventually approached the same average colony size of 250 mm², regardless of growth rate (Fig. 7). As expected, slower-growing colonies required a longer time to reach this average size. In this case, the intra-population cost of defense demonstrated the area deficit of a slower-growing colony at the time at which the faster-growing colonies have
filled the domain. However, this trend was dramatically altered when colonies from mixed populations containing both undefended and defended colonies competed for space. The inter-population cost of defense demonstrated the area deficit of defended colonies in comparison with undefended colonies at the end of the simulation duration. Defended, slower-growing colonies were prevented from reaching the average size by competing undefended, faster-growing colonies which exceeded the average size. Mixed population competition increased the cost of defense with inter-population costs of defense exceeding intra-population costs of defense between pH 7.9 and 7.3. At pH 7.0, all colonies grew too slowly to fill the domain over a 75 d simulation, thus costs of slower growth using these 2 metrics could not be calculated. In addition, undefended colonies at pH 7.0 grew slower than defended colonies and did not exhibit a cost of defense.

Inter-population costs of defense were higher than intra-population costs of defense at pH 7.9−7.3. The intra-population costs of defense were calculated as 37.9, 58.4, and 17.2 mm² for pH values 7.9, 7.6 and 7.3 respectively (label a in Fig. 7). In comparison, the inter-population costs of defense were much higher, at 183.2, 207.5, and 121.9 mm² for pH values 7.9, 7.6 and 7.3 respectively (label b in Fig. 7). Costs of defense initially increased at mild OA conditions (pH 7.6) but were reduced in severe OA conditions (pH 7.3). Compounding costs of slow growth in the context of competition resulted from a reduced growth rate and the inability to grow into space already occupied by faster-growing colonies.

In model simulations, all colonies settled simultaneously, to control for the effect of settlement time which likely influences colony competitiveness. Faster-growing colonies quickly gained an early competitive advantage over slower-growing colonies, which was maintained throughout the simulations. Overall, smaller colony growth rates yielded smaller population size structures. Population size structures are represented as density plots displaying the frequency of different sized colonies at the beginning, middle and end of model simulations (Fig. 8).

**DISCUSSION**

We sought to understand the impacts of OA-related stress at both the organism and population level using the bryozoan *Membranipora* and its specialist nudibranch predator *Corambe* as a case study. Using this organismal system, we developed a spatially explicit model to infer population-level consequences of experimentally determined individual-level responses to OA, which would otherwise be difficult to assess. Key characteristics of bryozoan life history, such as the cost of defense and the impor-
tance of space limitation, were maintained within the range of near-future seawater pH projections (7.6, 7.3). At pH 7.0, likely not representative of conditions that are or will soon be experienced, we observed apparent physiological limits in *Membranipora*; colonies at this low pH suffered low growth rates and high zooid senescence. Here, we focus on responses and ecological interactions for pH 7.9−7.3, above this apparent toxicity limit. When scaled to population levels in the context of space competition, individual-level reductions in growth rate associated with the cost of defense were amplified, resulting in multiplicative decreases in colony area, and likely reduced fitness since reproduction is generally proportional to area. The population-level cost of defense increased at moderate OA conditions but decreased at more severe OA conditions likely due to interacting effects of increased growth of undefended colonies at moderately low pH and space competition. Our results demonstrate the need to consider population-level mechanisms such as space competition when making assessments of ecological consequences of environmental stress.

*Membranipora* growth rates and calcification under OA

Contrary to our hypotheses, we observed maximum growth rates near target pH 7.6, corresponding to a moderately high pCO$_2$ of 985 μatm. This is lower than current average ambient pH conditions in the Salish Sea, but not outside the range normally experienced by *Membranipora* in the field (Murray et al. 2015). Previous studies have also documented increased growth in bryozoans at moderately low pH.
conditions. The calcifying bryozoans *Electra pilosa* (Saderne & Wahl 2013), *Celleporella cornuta* (Swezey et al. 2017a) and *Jellyella tuberculata* (Swezey et al. 2017b) exhibited increased growth in response to moderately high pCO₂, corresponding with moderately low pH conditions, relative to ambient (*E. pilosa*, 1200 μatm; *C. cornuta*, 1150 μatm; *J. tuberculata*, 1050 μatm). *E. pilosa* displayed a parabolic growth response curve exhibiting maximum growth at 1200 μatm (Saderne & Wahl 2013), similar to our results, suggesting these types of response curves may not be unusual in bryozoans.

Maximum skeletal density also occurred near target pH 7.6, reflecting observed trends in growth rate and implying a positive relationship between growth rate and skeletal density with a possible physiological link. Observed trends in growth and calcium production could reflect the calcification mechanism used by *Membranipora*, which is currently poorly understood. Many calcifiers secrete CaCO₃ from a calcifying fluid space of elevated pH, making it energetically favorable to convert HCO₃⁻ to CO₃²⁻ rather than extracting CO₃²⁻ directly from seawater. Micro-electrode studies have shown calcifying fluids to be upwards of 2 units above ambient seawater pH (Al-Horani et al. 2003). Ries et al. (2009) documented increases in calcification with respect to acidification in multiple taxa of marine calcifiers, and suggested that these increases could be due to strong control over calcifying fluid pH and H⁺ regulation at the calcification site used to convert HCO₃⁻ to CO₃²⁻. In *C. cornuta*, OA also increased the expression of protective organic coverings in areas of new growth, which can prevent dissolution while allowing calcification to continue (Swezey et al. 2017a). A combination of strong chemical control over a calcifying space and protective membranes might be responsible for *Membranipora’s* robust calcification response to OA.

In the bryozoan *C. cornuta*, increased growth was associated with costs including reduced investment in reproduction and lighter skeletons (Swezey et al. 2017a). We did not find a cost associated with calcifi-
cation in Membranipora. However, it is possible that our observed increases in growth and skeletal density in Membranipora are associated with costs that were not measured in this study, such as reduced per-zoid reproduction or altered metabolic costs.

**Spine formation in OA conditions**

Inducible spine formation was sustained across the entire range of pH conditions we investigated, with low pH correlated with slightly shorter spines. Membranipora spines are chitinous (Harvell 1984), rather than calcified, so their chemical composition may be less vulnerable to pH changes than zooids’ CaCO₃ skeletons. Predator cues in all pH conditions induced spines in consistent patterns, and were deposited concurrently with new skeleton formation. Induction of spines suggests that Membranipora colonies were able to detect predator cues in seawater with pH as low as 7.0. Shorter spines in lower pH may imply slight impairment of predator detection mechanisms in Membranipora due to OA. If so, this would be consistent with diminished chemosensitivity observed in other taxa (Dixson et al. 2010, Manríquez et al. 2013, Dodd et al. 2015, Weiss et al. 2018). Our observations are perhaps most analogous to Weiss et al. (2018), who documented decreases in the length of induced neck teeth in Daphnia spp. due to decreased predator-detection ability at higher pCO₂. While our data do not suggest strongly impaired predator detection in Membranipora, predator cue concentrations in our treatments may have been high enough such that cues were detected despite decreased sensitivity. By using high levels of cue, we effectively tested for physiological limits to spine formation, but not cue detection thresholds. Spine formation occurred in some control colonies due to likely diffuse cross-contamination of predator cue that made it effectively a ‘low predation cue’ treatment. In these low-level induction conditions, abundance of corner spines decreased with decreasing pH. This is consistent with the hypothesis that OA may affect defense formation, but only at very diffuse predator cue concentrations.

**Nudibranch consumption rate responses**

Zooid consumption rates by nudibranchs were slightly affected by pH, as shown by a significant interaction of spine presence and pH. With decreasing pH, nudibranch consumption increased in the presence of spines. Lower pH conditions apparently reduced the benefit conferred by induced spines in higher pH conditions, with nudibranchs consuming nearly equal numbers of zooids from defended and undefended colonies at the lowest pH. This trend suggests that OA stress fundamentally compromised the benefit of defenses in Membranipora, and represents potential differential effects of pH on predators and prey. This could have resulted from a variety of mechanisms acting on nudibranchs, bryozoans or both. OA could have effects on feeding physiology and energetic requirements in Corambe. OA stress has been shown to decrease the energetic content of some prey taxa, potentially increasing feeding rates in predators to meet their energetic requirements (Kroeker et al. 2014). Future experiments are needed to establish effects of OA on physiological processes of both Corambe and Membranipora, as we did not measure them in our study. Effectiveness of protective spines in Membranipora could also have been altered by OA. Consistent with the slight decrease in spine length at lower pH, shorter spines may not confer the same protective advantage as the longer spines induced at higher pH. Alternatively, OA stress may have compromised the structural integrity of spines, enabling higher consumption rates of defended zooids. OA has been shown to negatively affect defensive polymorphs in the bryozoan Schizoporella errata (Lombardi et al. 2011). While prey detection in Corambe may be sensitive to OA, like in other molluscs that have exhibited decreased sensory ability in OA conditions (Manríquez et al. 2013), our experiments did not test for impaired prey detection as Corambe was confined to a small area on top of colonies requiring only short-range detection of prey. Therefore, differences in consumption rates across pH are not likely due to impaired prey detection, but this mechanism may be relevant in the field and merits further investigation.

**Population-level insights from experimental data: inferences from a spatially explicit model**

Our experiments quantified the roughly exponential growth patterns exhibited by isolated colonies in the absence of geometric constraints on growth. The model provides insight into the implications of these different growth rates in a more realistic context where space is frequently limiting and where colony growth is constricted by dynamically interacting competitive neighbors. Compared to experimentally determined growth rates in the absence
of space competition, simulated space limitation in our model reduced area-based colony fitness. Our results reaffirm and add quantitative detail to the importance of density-dependence in assessing *Membranipora* area-based colony fitness. Modeling provided new insights into effects of environmental stress, allowing us to consider population-level processes in evaluating the relative effects of pH and predation pressure on area-based colony fitness and competitiveness. Within the pH range 7.9−7.3, our model results show potential amplification of the cost of defense due to space competition pressure, demonstrating how growth patterns of competing colonies shape individual area-based fitness. Specifically, the cost of defense increased in mixed populations containing defended and undefended colonies, representing both the costs incurred by reduced growth and space occupied by faster-growing colonies. In addition to emphasizing the need to assess fitness in the context of populations, our results also highlight the selective advantages of inducibility, where these high costs of reduced growth are minimized and only incurred when predation pressure is sufficiently high. Our model only considered intraspecific competition; however, if *Membranipora* is less vulnerable to OA, this may confer competitive advantage over other species with which it shares a common substrate such as other bryozoans, tunicates and epibionts experiencing space limitation.

Our model simulations represented competition between colonies that are undefended or defended throughout the entirety of their growing period, to evaluate the extremes of fitness tradeoffs associated with defenses. In the field, predation cue concentrations would need to be regularly high to maintain constant production of defenses, and positions of defended colonies would likely be dictated by predator placement in the field and not random, as used in our model. Therefore, our model simulations most accurately represent competition between constitutively spined colonies (those that are spined even in the absence of predators, which constitute 6.2% of the population; Harvell 1986) and unspined colonies. The high cost of defense in the context of space competition emphasizes the benefit of inducibility, where costs are only incurred when necessary. Predation was not included in our model, so we did not quantify the benefits of defense across OA conditions which may offset some of the predicted costs. Beyond demonstrating that both predator–prey relationships and intraspecies competition can influence OA responses in *Membranipora* and *Corambe*, our modeling results also suggest that organismal-level effects are oftentimes insufficient to estimate population-level effects of environmental stress in space-limited species.

**Membranipora is relatively robust to OA**

Given the metrics measured in our experiments and modeled population-level effects, *Membranipora* in the Salish Sea appear to be less vulnerable to OA than many other calcifiers, particularly within the range of near-future predictions, and may be well-suited to tolerate changes in ocean chemistry. This relative robustness may be in part attributable to the frequency of naturally low and fluctuating pH in the Salish Sea, and to conditions in their immediate microhabitats on kelp. pH values in the Salish Sea fluctuate seasonally between 7.9 and 7.6 (Murray et al. 2015), and have been recorded as low as 7.3 in Washington state coastal environments (Wootton & Pfister 2012). Diffusional boundary layers on kelp can create distinctly different microenvironments for *Membranipora*, modulated by photosynthesis and respiration, which may provide a partial refuge from chemical changes occurring in ambient seawater (Cornwall et al. 2014). Fluctuations of oxygen concentrations and pH within this micro-environment may also exceed ambient conditions serving to precondition bryozoans to low pH environments, especially in slow-flow environments where boundary layers are thicker (Noisette & Hurd 2018). While we considered conditions of pH 7.6 to be suboptimal, this may not in fact be very stressful for *Membranipora* if it is within the pH range normally experienced within the boundary layer.

**Implications for Membranipora population dynamics**

On the east coast of North America, *Membranipora* is a significant invasive biofouling organism in kelp forests that contributes to kelp forest defoliation (Scheibling & Gagnon 2009, Saunders et al. 2010). Population-level size structures in our simulations may have implications for kelp forest health, which are culturally and ecologically important ecosystems. From our study, *Membranipora* does not appear particularly vulnerable to OA, having maintained growth, CaCO₃ production and spine formation at least to pH 7.3, with maximum growth and calcification near pH 7.6. Predicted population structures of
undefended colonies were larger under pH 7.6 conditions, suggesting Membranipora may have an invasive edge at lower pH conditions. Our observations and simulations of undefended colonies are most relevant for predictions of invasive potential on the east coast, since Corambe is not prevalent there (Lambert et al. 1992). Membranipora may represent an example of a relatively resilient species to ocean change as studies of increased temperature have demonstrated increased growth as well (Saunders et al. 2010), although potential interactions of OA and warming have not been investigated.

CONCLUSIONS

Our study demonstrates the need to consider both interspecific interactions and intraspecific interactions on both organism and population levels in assessments of effects of ocean acidification. Overall, Membranipora colonies were less vulnerable to moderate OA conditions than hypothesized but responses were heavily influenced by predator exposure and space competition among conspecifics. By modeling population-level processes, we were able to understand effects of individual-level experimental responses to OA under the pressures of space limitation. More broadly, these environmentally induced geometric tradeoffs associated with space limitation can be applicable to other modular encrusting organisms affected by changing environments.

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