Ervilia castanea (Mollusca, Bivalvia) populations adversely affected at CO2 seeps in the North Atlantic

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HIGHLIGHTS

- The bivalve Ervilia castanea was studied at volcanic CO2 seeps and reference sites.
- Abundance, size and net-calciﬁcation were inversely related to CO2 levels.
- Large individuals were scarce or absent at high CO2 sites.
- Recruitment of this bivalve was highest at the CO2 seeps.
- Abundance and size of E. castanea were positively correlated with Chl-a in sediment.

ABSTRACT

Sites with naturally high CO2 conditions provide unique opportunities to forecast the vulnerability of coastal ecosystems to ocean acidification, by studying the biological responses and potential adaptations to this increased environmental variability. In this study, we investigated the bivalve Ervilia castanea in coastal sandy sediments at reference sites and at volcanic CO2 seeps off the Azores, where the pH of bottom waters ranged from average oceanic levels of 8.2, along gradients, down to 6.81, in carbonated seawater at the seeps. The bivalve population structure changed markedly at the seeps. Large individuals became less abundant as seawater CO2 levels rose and were completely absent from the most acidﬁed sites. In contrast, small bivalves were most abundant at the CO2 seeps. We propose that larvae can settle and initially live in high abundances under elevated CO2 levels, but that...
1. Introduction

Increasing atmospheric CO2 concentrations have caused the pH of the surface ocean to fall by 0.1 units since the pre-industrial age (Bindoff et al., 2019; Dupont and Pörtner, 2013; Feely et al., 2009; Keeling et al., 2017) and further decreases, between 0.1 (RCP2.6) or 0.3 (RCP8.5) pH units, are expected by 2100 (Bindoff et al., 2019). This process of ocean acidification affects calcification of marine organisms, leading to permanent damage and reduced viability (Shirayama, 2005) and causing major changes to marine ecosystems (Doney et al., 2012). Shallow-water CO2 seeps provide analogues for the effects of chronic ocean acidification, at spatial and temporal scales that are difficult to mimic in aquaria or mesocosms (Hall-Spencer and Harvey, 2019). Care is needed to assess potential anomalies in temperature, alkalinity, metals and H2S which are frequently found at these volcanic sites and can also influence biological responses, but are not associated with ocean acidification (Vizzini et al., 2013; Mishra et al., 2020).

This type of in situ approach has shown that ocean acidification can lead to ecosystem simplification and reorganization of marine communities (Fabricius et al., 2015; Kroeker et al., 2013a, 2011; Agostini et al., 2018). The structure of benthic marine communities changes greatly when they are exposed to carbonated seawater (Widdicombe et al., 2018). The structure of benthic marine communities changes greatly when they are exposed to carbonated seawater (Widdicombe et al., 2018). Communities tend to change from being dominated by calcareous organisms to non-calcareous ones (Baggini et al., 2014; Christen et al., 2013; Hall-Spencer et al., 2008), granting the latter a perceived advantage in a future acidified ocean (Kroeker et al., 2013b; Vihtakari et al., 2013).

Organisms with calcareous shells and skeletons, such as coralline algae, molluscs and echinodermes, are expected to be adversely affected by increasing CO2 levels (e.g., reductions in abundance or local extinctions; Kroeker et al., 2011; Ricevuto et al., 2012) due to susceptibility of the calcification process to reduced carbonate availability (Bindoff et al., 2019). Garilli et al. (2015) found two gastropod species dwarfed as an adaptation to carbonated seawater, which allowed them to maintain calcification and partially repair shell dissolution. Other studies show that pH (or pCO2) on their own (apart from the carbonate saturation state) can cause physiological effects that result in less energy available for mineralization (Cyrnak et al., 2016; Palmer, 1992; Roleda et al., 2012; Waldbusser et al., 2015).

The early life stages of molluscs are often particularly sensitive to ocean acidification (Clements et al., 2016; Green et al., 2013, 2004; Jansson et al., 2015; Ricevuto et al., 2012). Previous studies have shown stunted growth and shell development (Amaral et al., 2011; Gazeau et al., 2010; Hettinger et al., 2012; Kroeker et al., 2010; Talmage and Gobler, 2010; Waldbusser et al., 2015); reduced sperm motility, fertilization and hatching rates (Barros et al., 2013; Gazeau et al., 2010; Kroeker et al., 2010); slower metamorphosis and delayed settling (Jansson et al., 2015; Talmage and Gobler, 2010); decreased survival (Gazeau et al., 2010; Talmage and Gobler, 2010); changes to biochemical composition (Range et al., 2014a; Talmage and Gobler, 2010); increased frequency of shell abnormalities and dissolution (Barros et al., 2013; Green et al., 2004; Pereira et al., 2015; Talmage and Gobler, 2010); altered metabolic rates (Fernández-Reiriz et al., 2012, 2011; Mellado et al., 2019); altered burrowing behaviour and post-settlement dispersal (Clements et al., 2016) and; decreased settlement success (Gazeau et al., 2010). Furthermore, some species reallocate energy budgets to cope with ocean acidification, with growth and reproduction often compromised to sustain fitness and survival (Clements et al., 2020; Durland et al., 2019; Widdicombe and Spicer, 2008). This type of trade-off has been found to produce negative (Dupont et al., 2012; Hettinger et al., 2012), positive (Parker et al., 2012; Zhao et al., 2018, 2019) and neutral (Clements et al., 2020) carry-over effects from the parental brood stocks to the next generation. These results indicate that the responses of the early life stages of bivalves to increased CO2 go well beyond the detrimental effects on shell calcification.

Responses to ocean acidification are species-specific and even closely related species can react differently, depending on various factors, such as life history traits, habitat, sex, nutritional status, adaptation potential and methodological conditions (Calosi et al., 2013; Doney et al., 2012; Dupont and Thorndyke, 2005; Garilli et al., 2015; Kroeker et al., 2013a, 2011; Marçeta et al., 2020; Pörtner, 2008; Vihtakari et al., 2013). Individual responses to ocean acidification are of key importance for any given species, but they also have cascading effects along trophic relations and ecological interactions, making them important drivers in shaping community structures and phase shifts (Kroeker et al., 2011; Range et al., 2014b). Consequently, the direct effects of CO2 on competitors, prey, predators and the quantity and quality of food supply can be translated to density control mechanisms (Metaxas, 2015; Range et al., 2012) and mediated by adaptation or acclimation (Kroeker et al., 2011).

The species targeted in this study, Ervilia castanea (Montagu, 1803), is a semelid bivalve (Morton, 1990) found from south of the British Isles into the Mediterranean (Babio and Bonnin, 1987). It has a brownish, oval and elongated shell, which reaches a maximum length of 15 mm (Macedo et al., 1999; Morton, 1990). In the Azores, where the majority of the bivalve species tend to be ~50% smaller than their continental conspecifics (Morton et al., 2013), the maximum length reported is 6.7 mm (Morton, 1990). This generalized dwarfism of bivalves is commonly found around oceanic islands and has been related to the low availability of food in these oligotrophic waters (Morton et al., 2013). Individuals of this species have a high potential reproductive output, with maturation occurring at a length of 3.5 mm for females and 5.5 mm for males (Morton, 1990). Ervilia castanea occupies subtidal coarse-grained, well-sorted sediments overlain by clean, nutrient poor, oceanic waters, to depths exceeding 40 m (Morton, 1990). In the Azores it is numerically dominant on sandy substrata and mainly predated upon by naticid gastropods (Morton, 1990; Morton et al., 2013).

In this study, we collected in situ evidence about the effects of elevated seawater CO2 on populations of the bivalve Ervilia castanea, along submarine volcanic CO2 gradients in the Azores. We recorded abundance, size-structure, biomass and tissue composition. Three hypotheses were tested: we expected the abundance (Hyp. 1) and net-calcification (i.e., accretion minus dissolution, Hyp. 2) of Ervilia castanea to decrease with rising CO2 levels and for the size structure of the population (Hyp. 3) to be modified along the CO2 gradients.

2. Methods

2.1. Selection of the study sites

The Azores are nine volcanic islands located on the triple junction of the Eurasian, North-American and Nubian tectonic plates. Shallow-water volcanic CO2 seeps occur around several of the Azorean islands.
and seamounts (Couto et al., 2015; Rajasabapathy et al., 2014; Tribollet et al., 2018; Viveiros et al., 2016; Wallenstein et al., 2013, 2009). Two of these coastal sites, located off the islands of São Miguel and Faial, were chosen for this study, based on their shallow depth and accessibility, as well as previous knowledge about the intensity and composition of the gas emissions. Ponta da Lobeira (São Miguel, 37°43′31.8′′N 25°19′01.6′′W, Supplementary Video 1) is a site with strong degassing at 7 m depth, located <2 km from the village of Ribeira Quente, on the south flank of Furnas Volcano. Ponta da Espalamaca (Faial, 38°32′36.4′′N 28°35′48.4′′W, Supplementary Video 2) is a site with more diffuse degassing, at 37 m depth, located 3 km from Horta port (Fig. 1). Supplementary videos one and two show the type of benthic habitat and the degassing fluxes at Ponta da Lobeira and Ponta da Espalamaca, respectively.

2.2. Experimental design

Off each island, three sampling sites, with similar depth, bottom typology exposure to currents and wave action, were selected: a significant degassing site (High CO2); a transitional site (Intermediate CO2), where the seawater carbonate chemistry was still affected, but with no visible CO2 release; and a reference site (Reference), unaffected by the CO2 emissions. Test dives with a pH multiprobe (YSI6600) were used to define these sites. In São Miguel, sampling was done between 3 and 7 July 2014, at depths ranging between 6 and 10 m. All sites in São Miguel were relatively sheltered from the dominant wave action. Sampling in Faial was done between 10 and 15 July 2014, at depths between 28 and 37 m. Given the proximity to the Faial-Pico channel, all the sites in Faial were strongly exposed to tidal currents.

2.3. Sample collection and laboratory procedures

2.3.1. Free gas composition and flux

Free gas samples were collected at the High CO2 sites using Giggenbach bottles (Supplementary Video 3) and gas composition was analysed by titration and gas chromatography (for additional details, Ferreira et al., 2005). Inverted plastic funnels, connected to volumetric flasks, were used to measure gas emission rates. In São Miguel (Ponta da Lobeira), this sampling was stratified into low, medium and high flux areas, defined by visual inspection of bubbling, with three replicate samples collected at each level. In Faial (Ponta da Espalamaca), intensity of bubbling was more homogeneous, so the different levels of emission could not be established and two replicates were collected in each of three haphazardly selected points of emission (A, B and C).

2.3.2. Physicochemical properties of seawater

Seawater temperature, salinity, dissolved oxygen and total pH (pHT) were measured at least once per day at each sampling site, over the course of 3 or 4 days. A YSI6600 multiprobe was used for continuous records (pHT, temperature and salinity). Seawater carbonate chemistry speciation was calculated from temperature, salinity, silicate, phosphate, multiprobe pHt values and TA using CO2SYS (Lewis and Wallace, 1998) with the equilibrium constants determined by Mehrbach et al. (1973) as refitted by Dickson and Millero (1987), by interpolating discrete variables from Niskin sampling (silicate, phosphate and TA) with continuous multiprobe records (pHT, temperature and salinity). Accordingly, the calculations were done using the average values of TA, silicate and phosphate for each vertical profile (i.e., average of bottom and surface samples per sampling site and per day).

2.3.3. Macrofauna

Sediment macrofauna samples were collected by scientific divers, from an area of 0.25 m² excavated to 15 cm depth, using an airlift fitted with a 500 μm mesh bag (Supplementary Video 4). Replicate samples were spaced by 2 to 6 m and haphazardly positioned within each site, except in the High CO2 sites, where the points of gas emission were explicitly avoided. In São Miguel, nine replicates were collected at each site. In Faial, time restrictions and operational limitations while diving at greater depths and in stronger currents, enabled the collection of only six samples of macrofauna in the Reference and Intermediate sites and seven in the degassing site. All organisms retained in the bag were preserved in 5% buffered formaldehyde, hand-sorted and later preserved in 70% ethanol until taxonomic determination and counting. All the Ervilia castanea were individually measured and subsequently pooled per replicate and shell length (1 mm size-classes, 0 to 8 mm). These pools of bivalves were dried for a minimum of 36 h at 60 °C (dry weight - DW) and subsequently burnt in a muffle furnace for 4 h at 550 °C (ash weight - AW). Ash-free dry weight (AFDW) and relative ash weight (%AW) were estimated, per replicate and size-class using Eqs. (1) and (2), respectively:

\[
\text{AFDW} = \frac{\text{Dry Weight} - \text{Ash Weight}}{\text{Dry Weight}} \quad (1)
\]

\[
\%\text{AW} = \frac{\text{Ash Weight}}{\text{Dry Weight}} \times 100 \quad (2)
\]

2.3.4. Sediments

Hand-held PVC cores (3.5 cm diameter per 15 cm length) were used to collect two sediment samples, adjacent to each macrofauna sample. The first core was used for determination of granulometry and organic matter/carbonate content. A set of six sieves (63 μm, 125 μm, 250 μm, 500 μm, 1 mm and 2 mm) was used for sediment pooling per grain size. After sieving, each fraction was weighed and related to the total weight of the sample, as a percentage. The organic matter content and carbonate content in sediments was determined by the loss-of-ignition (LOI) method (Heiri et al., 2001). Sediment was dried in an oven for 72 h at 100 °C (DW100), from which, 10 g of each sample were burnt for 8 h at 550 °C (DW550) and subsequently for 2 h at 900 °C (DW900). Organic matter content (LOI550) and carbonate content (LOI900) were then determined using Eqs. (3) and (4), respectively:

\[
\text{LOI550} = \frac{\text{DW100} - \text{DW550}}{\text{DW100}} \times 100 \quad (3)
\]

\[
\text{LOI900} = \frac{\text{DW550} - \text{DW900}}{\text{DW100}} \times 100 \quad (4)
\]

The second core was used for carbon/nitrogen ratio and photosynthetic pigment analyses. Sediment samples were frozen and kept in the dark for preservation of chloroplastic pigments. Approximately 12 g of frozen sediment were lyophilized in a Telstar LyoAlfa 15 for 48 h, to get the 5 g of material needed. For total organic carbon (TOC) and total nitrogen (TN) samples were homogenized and acidified with...
dilute hydrochloric acid until complete decarbonization. TOC was measured with a Thermo Scientific Flash 2000 elemental analyser. For analyses of the chloroplastic pigments, samples were homogenized and pigments extracted in 90% acetone. Pigment separation was done using reverse phase high performance liquid chromatography (HPLC) and measured with a Gilson fluorescence detector according to Wright and Jeffrey (1997).

2.4. Statistical analysis

Two-way analyses of variance (ANOVA) were used to test for differences in abundance and biomass of Ervilia castanea within each island (São Miguel and Faial). CO₂ level (High, Intermediate and Reference) and size-class were considered as fixed orthogonal factors. The data was untransformed and Cochran’s test was used to test for
hydrogen sulfide (H₂S), methane (CH₄), hydrogen (H₂), helium (He), nitrogen (N₂), oxygen (O₂), and argon (Ar). b.d.l. (below detection limit).

| Site   | Emission rate | CO₂ (mol%) | H₂S (mol%) | CH₄ (mol%) | H₂ (mol%) | He (mol%) | N₂ (mol%) | O₂ + Ar (mol%) | Gas flux (L/min/m²) |
|--------|---------------|------------|------------|------------|-----------|-----------|-----------|----------------|------------------|
| S. Miguel | Low  | 99.49      | b.d.l.     | 1.14E⁻² | 1.39E⁻⁴ | 1.01E⁻¹³ | 0.37      | 0.13           | 0.58             |
| S. Miguel | Medium | 99.86      | 9.66E⁻⁴ | 1.25E⁻² | 4.00E⁻⁵ | 9.95E⁻¹⁴ | 0.11      | 0.02           | 10.26            |
| S. Miguel | High       | 99.58      | 8.97E⁻² | 1.39E⁻² | 3.97E⁻⁶ | 1.06E⁻¹³ | 0.23      | 0.05           | 24.88            |
| Faial   | A       | 98.61      | b.d.l.     | b.d.l.    | b.d.l.    | b.d.l.    | b.d.l.    | b.d.l.         | 2.64             |
| Faial   | B       | 98.91      | b.d.l.     | b.d.l.    | b.d.l.    | b.d.l.    | 3.41E⁻¹³ | 0.15           | 2.64             |
| Faial   | C       | 99.35      | b.d.l.     | b.d.l.    | b.d.l.    | 2.24E⁻¹³ | 0.56      | 0.09           | 2.64             |

3.3. Population structure of Ervilia castanea

A total of 3134 Ervilia castanea individuals were collected during this study, representing 58% of the overall abundance of macrofauna. E. castanea was the dominant organism in all sampling sites, except for the Intermediate site in Faial, where Nematodes of the genus Metoncholaimus were numerically dominant. Densities of E. castanea up to 1140 ind·m⁻² were recorded in São Miguel, which generally had larger abundances, relative to Faial (Fig. 3). In Faial, the size-frequency distribution was clearly skewed towards smaller individuals at the High CO₂ site, while in São Miguel the sites tended to have a more similar distribution. Average abundances (across all size classes) consistently peaked at reference sites and this effect was significant in both Islands (Table 3). In contrast, the smallest size class (<1 mm), composed of recently settled postlarvae (i.e., plantigrades, sensu Carriker, 1961) and juveniles (spat), showed consistently larger abundances in sites affected by CO₂ than in reference sites, which explains the significant interaction term for São Miguel. The dominant size class in Faial (1–2 mm) also showed a similar pattern, reaching peak abundance in the high CO₂ site (Fig. 3).

3.4. Biomass of E. castanea

The smallest (<2 mm) and largest (>5 mm) size classes were excluded from the biomass analyses, due to lack of precision in weighing and absence in the high CO₂ sites, respectively. The average biomass per individual (DW, AW and AFDW) increased with shell length for the three intermediate size classes (between 2 and 5 mm shell length, Fig. 4). This pattern was visible in all sampling sites, except for the Intermediate site in Faial, where individuals larger than 2 mm were absent. Relative ash weight differed significantly among sites (%AW, Table 3), with the dominant size classes showing consistently larger values in reference sites (92% in both islands), than in sites affected by CO₂ emissions (83% in São Miguel and 75% in Faial). Although no quantification of shell dissolution or strength was done in this study, alterations were clearly apparent on the few larger individuals found at Intermediate and High CO₂ areas, particularly around the umbo, where it was usually associated with some degree of erosion to the periostracum (Fig. 5B and D).

3.5. Relationship between sediment characteristics and population structure of E. castanea

DistLM marginal tests indicated that 13 out of the 15 sediment variables considered were significantly related to the distribution and size structure of E. castanea (Table 4). The only exceptions were silt (sediment finer than 63 μm) and total nitrogen (TN). These two variables were, therefore, excluded from the model selection. The best overall DistLM solution included 4 sediment variables: 2 mm grain size, carbonate content, chlorophyll a and other pigments (diadinoxanthin, diatoxanthin, zeaxanthin and lutein). The coarser sediments were strongly associated with the intermediate and high CO₂ sites in Faial (Fig. 6), which also had the smallest abundances of E. castanea above 2 mm shell length (Fig. 3). Chlorophyll a was the dominant photosynthetic pigment in all sampling sites,
followed by fucoxanthin. All pigments showed higher concentrations in the reference sites, relative to the ones affected by CO₂, and this pattern was strongly associated with the intermediate size classes (2–5 mm shell length) of E. castanea. The reference site in Faial was also characterized by elevated values of carbonates (CaCO₃) and organic matter (OM), relative to all the other sampling sites (Table 4).

4. Discussion

Gas released from both seep systems was mainly CO₂ (>98.6%), with N₂, O₂, Ar and He also present in the dry phase. Residual fractions of other typical hydrothermal gases (H₂S, H₂ and CH₄) were also detected at Ponta da Lobeira (Table 1). The sulfuric component was, however, only present when the gas flux was medium to high (≥10 L/min/m²) and in relatively small concentrations (<10 ppm), when compared to the sub-aerial fumaroles of the Furnas Volcano (Caliro et al., 2015) or to other submarine vents (Tarasov et al., 2005). Boatta et al. (2013) recorded much higher concentrations of H₂S (400 ppm) in the vent bubbles at Vulcano Island, Italy, concluding that only a small proportion of sulfide enters or remains in the aquatic phase and that the dissolved levels dropped below the detection limit at 5 m distance from the emission point. Accordingly, although we cannot dismiss the possibility of

![Fig. 2. Violin plots for pH (total scale), CO₂ partial pressure (pCO₂), saturation of calcite (ΩCa) and aragonite (ΩAr); the frequency distribution (grey area), median (white dot) and quartiles (black bars) are presented for each site in each island, across all depths and dates of sampling.](image-url)
toxic gases, other than CO2, affecting the bivalves at Ponta da Lobeira, this potential confounding factor was clearly less important here than in most previously studied shallow vent sites. Other types of contaminants usually present in vent emissions, such as trace metals, were not assessed in our study. Their adverse biological effects are, however, usually localized to a few hundred meters from the vents (Vizzini et al., 2013), so it is unlikely that they could have affected the Intermediate CO2 sites.

Seawater temperature, salinity and total alkalinity varied within narrow ranges among the different sampling sites and these variations were independent of the CO2 emissions. Seawater pH, on the other hand, varied substantially among sites. This was mainly due to the increased frequency of extremely low pH values (down to 6.8) at the sites affected by CO2 emissions, rather than a consistent decrease of the average pH at those sites. Despite this, the gradient of CO2 partial pressure in São Miguel was clearly within the range predicted by the IPCC for the end of this century, which varies between 421 (RCP2.6) and 936 (RCP8.5) ppm (Hoegh-Guldberg et al., 2014; Pörtner et al., 2014). The gradients in seawater carbonate chemistry observed in São Miguel can, therefore, be unambiguously applied to infer on the response of benthic organisms to present conditions (Reference site) and future scenarios of ocean acidification (Intermediate and High CO2 sites). The gradients in carbonate chemistry were less clear for Faial, due to the greater depth, stronger currents and the fact that pH data was collected along the entire water column (37 m). These conditions complicate the interpretation, as variations exceed the average scenarios for ocean acidification in the foreseeable future. Nevertheless, the regulation of carbonate chemistry in coastal waters involves a multitude of drivers (i.e., watershed processes, nutrient inputs, changes in ecosystem structure and metabolism), apart from anthropogenic CO2 emissions, so variations of this magnitude are not unusual for this type of habitat (Duarte et al., 2013). In fact, the potential adaptations of coastal ecosystems to this increased environmental variability, induced by interactions between global and local drivers, are critically important to

Table 2
Seawater carbonate chemistry variables (mean ± SE) at each sampling site (R - Reference; I - Intermediate CO2; H - High CO2) in São Miguel (SM) and Faial (FA). The variables represented are: temperature (Temp), salinity (Sal), pH (total scale), total alkalinity (TA), CO2 partial pressure, bicarbonate (HCO3−), carbonate (CO32−), ion hydroxide (OH−), saturation of calcite (ΩCa) and aragonite (ΩAr). Each entry represents the average of bottom and surface measurements per sampling site and per day (n = 3).

| Location | Temp (°C) | Salinity (ppt) | pH | TA (μmol/kg) | TCO2 (μmol/kg) | pCO2 (μatm) | HCO3− (μmol/kg) | CO32− (μmol/kg) | OH− (μmol/kg) | ΩCa | ΩAr |
|----------|-----------|----------------|-----|--------------|----------------|-------------|-----------------|----------------|--------------|-----|-----|
| SM-R     | 20.13     | 36.93          | 8.10| 2371.47      | 2056.80        | 347.80      | 1827.08 ± 0.43  | 218.65         | 5.04          | 5.15 | 3.36 |
| ± 0.01   | ± 0.00    | ± 0.00         | ± 0.07| ± 0.27      | ± 0.45         | ± 1.02      | ± 195.25        | ± 0.63          | ± 0.01        | ± 0.00 | ± 0.00 |
| SM-I     | 20.06     | 36.98          | 8.03| 2350.92      | 2104.04        | 456.63      | 1995.06         | 4.34           | 4.60          | 3.00 |     |
| ± 0.01   | ± 0.00    | ± 0.00         | ± 0.00| ± 1.02      | ± 2.69         | ± 195.25    | ± 0.63          | ± 0.01         | ± 0.01        | ± 0.01 |     |
| SM-H     | 20.54     | 36.94          | 7.86| 2397.89      | 2199.16        | 818.95      | 2021.84 ± 1.92  | 151.51         | 3.31          | 3.57 | 2.33 |
| ± 0.00   | ± 0.00    | ± 0.00         | ± 0.06| ± 1.42      | ± 10.07        | ± 0.77      | ± 0.02          | ± 0.02         | ± 0.01        | ± 0.01 |     |
| FA-R     | 20.14     | 37.65          | 8.10| 2371.70      | 2650–46        | 345.94      | 1817.50 ± 1.33  | 222.00         | 5.12          | 5.20 | 3.39 |
| ± 0.04   | ± 0.01    | ± 0.00         | ± 0.00| ± 0.85      | ± 1.59         | ± 0.53      | ± 0.02          | ± 0.01         | ± 0.01        | ± 0.01 |     |
| FA-I     | 19.91     | 37.62          | 8.04| 2345.20      | 2654.58        | 594.01      | 1811.93 ± 8.37  | 213.05         | 5.11          | 4.99 | 3.26 |
| ± 0.05   | ± 0.01    | ± 0.01         | ± 0.00| ± 6.31      | ± 38.00        | ± 3.34      | ± 0.01          | ± 0.08         | ± 0.05        |      |     |
| FA-H     | 19.58     | 37.56          | 8.02| 2353.48      | 2609.69        | 744.14      | 1839.47 ± 5.16  | 205.37         | 4.75          | 4.81 | 3.14 |
| ± 0.05   | ± 0.01    | ± 0.01         | ± 0.08| ± 4.52      | ± 44.22        | ± 2.06      | ± 0.05          | ± 0.05         | ± 0.03        |      |     |

Fig. 3. Abundance (mean ± SE) of the bivalve Ervilia castanea per size-class and CO2 level in each of the islands; bars with different letters denote significant differences among CO2 levels within each size class (SNK tests, p < 0.05); values in parenthesis represent mean abundances, across all size classes, at each CO2 level in each of the islands.

Table 3
Analyses of variance for the abundance and relative ash weight (%AW) of Ervilia castanea for each island; CO2 level and Size Class are fixed orthogonal factors; df: degrees of freedom; MS: mean squares; # denotes groups of samples with heterogeneous variances (Cochran’s tests, p < 0.05).

| Source of variation | Abundance | Relative ash weight (%AW) |
|---------------------|-----------|--------------------------|
|                     | São Miguel | Faial |                     | São Miguel | Faial |
|                     | df | MS | F-ratio | p | df | MS | F-ratio | p |
| CO2 level           | 2  | 14,529 | 20.166 | <0.001 | 2 | 1042 | 3.342 | 0.038 |
| Size Class          | 6  | 6945 | 9.64 | <0.001 | 7 | 302 | 0.97 | 0.456 |
| CO2 × Size          | 12 | 3408 | 4.73 | <0.001 | 14 | 356 | 1.143 | 0.328 |
| Residual            | 168 | 720 | | | | | |
|                      | 128 | 312 | | | | | |
| CO2 level           | 2  | 0.0697 | 5.009 | 0.01 | 1 | 0.0807 | 7.728 | 0.016 |
| Size Class          | 2  | 0.0195 | 1.4 | 0.256 | 2 | 0.0294 | 2.814 | 0.097 |
| CO2 × Size          | 4  | 0.0224 | 1.763 | 0.151 | 13 | 0.0314 | 0.003 | 0.958 |
| Residual            | 51 | 0.0119 | | | 16 | 0.0214 | | |
forecast their sensitivity and vulnerability to future ocean acidification and cannot be ignored.

Hyp. 1 and Hyp. 2 were corroborated: *Ervilia castanea* significantly decreased in abundance and net-calcification, measured as relative ash weight, with increasing CO2 levels. Individuals of all sizes were represented in reference sites, while larger individuals (above 3 mm shell length) were scarce or absent in sites affected by CO2 emissions. This steep decline in the abundance of larger individuals was probably caused by high levels of post-settlement mortality, as a consequence of delayed or unsuccessful metamorphosis of recently settled pediveligers into postlarvae and juveniles (Bayne, 1965; Talmage and Gobler, 2009, 2010), reduced shell calcification (Clements and Hunt, 2017; Green et al., 2013, 2009, 2004), growth impairment (Parker et al., 2010) or a combination of these factors (Waldbusser et al., 2010).

The average biomass per individual (DW, AW and AFDW) varied among size-classes, but not among sampling sites. Given the relatively small contribution of the soft tissues (AFDW) to the overall biomass (DW), the pattern observed for relative ash weight (%AW) seems to have been mainly driven by variations in the mineral fraction of the shells (i.e., calcium carbonate content). Negative net-calcification rates have frequently been observed for infaunal and epifaunal marine bivalves, in response to elevated CO2 (see reviews by Gazeau et al., 2013 and Clements and Hunt, 2017). Although the relative contributions of accretion and dissolution to this net-effect cannot usually be disentangled, several authors suggest increased post-deposition dissolution as the main cause for the observed losses of shell mass (Findlay et al., 2011; Nienhuis et al., 2010; Range et al., 2012; Rodolfo-Metalpa et al., 2011; Tunnicliffe et al., 2009). In fact, accretion depends on both

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**Fig. 4.** Individual body weight of the dominant size classes of *Ervilia castanea* per CO2 level in each island; dry weight (outline, means ± SE) and ash weight (filled area, mean); percentage values on top of the bars denote relative ash weight (mean).

**Fig. 5.** Microphotographs of specimens of *Ervilia castanea* from different size classes and CO2 levels: A (1–2 mm, reference), B (1–2 mm, high CO2), C (2–3 mm, reference), D (2–3 mm, high CO2). Scale bar is 1 mm.
Table 4
Sediment characteristics (means ± SE) by island and location. Granulometry (<63 μm, 63 μm, 125 μm, 250 μm, 500 μm, 1 mm, 2 mm and 4 mm), organic matter (OM), carbonate content (CaCO₃), total organic carbon (TOC) and total nitrogen (TN) are expressed as percentages. Other pigments include diadinoxanthin, diatoxanthin, zeaxanthin and lutein. C:N ratio was not independently determined, but calculated, so it was not considered in the DistLM models.

| Island       | Reference (n = 9) | Intermediate (n = 9) | High (n = 9) | Reference (n = 6) | Intermediate (n = 6) | High (n = 10) |
|--------------|-------------------|----------------------|-------------|-------------------|----------------------|-------------|
| São Miguel   |                   |                      |             |                   |                      |             |
| <63 μm       | 0.127 ± 0.032     | 0.130 ± 0.072        | 0.102 ± 0.007 | 0.327 ± 0.080     | 0.071 ± 0.012        | 0.061 ± 0.010 |
| 63 μm        | 2.970 ± 0.857     | 2.625 ± 0.053        | 1.980 ± 0.375 | 7.420 ± 0.598     | 0.783 ± 0.613        | 0.245 ± 0.043 |
| 125 μm       | 37.00 ± 7.220     | 15.60 ± 2.080        | 38.10 ± 2.200 | 45.00 ± 1.570     | 19.40 ± 3.450        | 24.50 ± 1.480 |
| 250 μm       | 40.40 ± 4.550     | 59.90 ± 2.390        | 48.80 ± 0.827 | 22.00 ± 1.600     | 42.20 ± 6.320        | 45.70 ± 4.010 |
| 500 μm       | 15.2 ± 4.61       | 20.4 ± 3.61          | 9.3 ± 1.38   | 8.04 ± 1.17       | 3.26 ± 0.59          | 1.3 ± 0.063  |
| 1 mm         | 2.8 ± 1.09        | 3.29 ± 0.818         | 1.35 ± 0.268 | 7.02 ± 0.983      | 6.44 ± 0.869         | 8.78 ± 1.26  |
| 2 mm         | 0.595 ± 0.237     | 0.381 ± 0.117        | 0.137 ± 0.036 | 3.82 ± 0.458      | 13 ± 3.23            | 13.1 ± 2.17  |
| 4 mm         | 0.438 ± 0.347     | 0.039 ± 0.031        | 0.006 ± 0.006 | 5.91 ± 2.82       | 15.6 ± 4.93          | 6.34 ± 2.11  |
| OM (LOI550)  | 1.149 ± 0.053     | 0.769 ± 0.030        | 1.046 ± 0.334 | 3.124 ± 0.168     | 1.073 ± 0.038        | 1.18 ± 0.071  |
| CaCO₃ (LOI900) | 0.477 ± 0.043   | 0.219 ± 0.035        | 0.574 ± 0.042 | 6.516 ± 0.393     | 0.085 ± 0.018        | 0.209 ± 0.048 |
| Fucoxanthin (μg/g) | 0.426 ± 0.051 | 0.095 ± 0.013        | 0.063 ± 0.014 | 0.211 ± 0.023     | 0.04 ± 0.007         | 0.054 ± 0.027 |
| Chl a (μg/g) | 1.097 ± 0.13      | 0.231 ± 0.029        | 0.169 ± 0.047 | 0.717 ± 0.116     | 0.072 ± 0.019        | 0.12 ± 0.064  |
| Other pigments (μg/g) | 0.008 ± 0.008 | 0.007 ± 0.012        | 0.06 ± 0.011 | 0.061 ± 0.013     | 0.076 ± 0.012        | 0.079 ± 0.010 |
| TN           | 0.077 ± 0.016     | 0.007 ± 0.012        | 0.06 ± 0.011 | 0.061 ± 0.013     | 0.076 ± 0.012        | 0.079 ± 0.010 |
| TOC          | 0.161 ± 0.014     | 0.142 ± 0.010        | 0.166 ± 0.029 | 0.256 ± 0.032     | 0.159 ± 0.018        | 0.138 ± 0.006 |
| C:N ratio    | 2.44 ± 0.338      | 2.100 ± 0.201        | 2.685 ± 0.499 | 5.016 ± 0.972     | 2.348 ± 0.402        | 1.947 ± 0.266 |

Fig. 6. Distance-based redundancy analysis (dbRDA) for the best solution of distance-based linear model (DistLM). Vector overlays represent significant variables included in the model, selected with the BEST procedure and Akaike’s information criterion; top panel represents explanatory (sediment) variables and bottom panel the response variables (abundances of E. castanea per size class); the length of the vectors represents the relative influence of each variable to the ordination; São Miguel – SM, F – Faial, Reference – Intermediate, H – High CO₂.

Carbonate saturation of the surrounding water and the physiological status of the organism, which is largely determined by food availability, whereas dissolution relates solely to the external saturation state (Marshall et al., 2019).

Dissolution of the shell is recognized as an important source of post-settlement mortality in bivalves, having been associated with the exponential loss of individuals after the transition from the larval to the benthic stage (Green et al., 2004). The type of shell damage observed at sites affected by CO₂ emissions has been previously reported in several laboratory (Bressan et al., 2014; Gazeau et al., 2014; Range et al., 2012) and in situ studies (Rodolfo-Metalpa et al., 2011; Thomsen et al., 2010). Other studies have also found that increased CO₂ levels are often associated with decreases in the mechanical properties of the shell, such as, thickness (Bressan et al., 2014; Gaylord et al., 2011; Talmage and Gobler, 2010), density (Klok et al., 2014; Rühl et al., 2017), hardness and fracture resistance (Beniash et al., 2010; Dickinson et al., 2013; Gazeau et al., 2013; Welladsen et al., 2010). Thinner or weaker shells have also been shown to increase bivalve mortality through several pathways, mediated by interactions with other environmental stressors (Dickinson et al., 2012), predators (Grey et al., 2007; Sanford et al., 2014), competitors (Coleman et al., 2014) or parasi-tes (MacLeod and Poulin, 2015).

A number of recent studies have shown that the detrimental effects of elevated CO₂ on the calcification and growth of bivalves can be mitigated or avoided when the food supply is not limiting (Melzner et al., 2011; Pansch et al., 2014; Ramajo et al., 2016b, 2016a; Thomsen et al., 2013). We did not assess if food limitation occurred during our study, but sediments at reference sites had greater amounts of chlorophyll-a and inorganic carbon, relative to sites affected by CO₂ emissions (Fig. 6, Table 4). This implies that reference sites were better feeding grounds for intrafaunal bivalves. Concentrations of chlorophyll-a in seawater showed a similar pattern, with the smallest values consistently being found at the High CO₂ sites (unpublished data). Chlorophyll-a was, therefore, negatively correlated with CO₂ level and positively correlated with abundance and size of E. castanea. This strongly suggests that the indirect effects of CO₂, mediated by trophic interactions (i.e., food availability), may be equally or even more important than its direct effects on growth and survival of these clams.

Exposure to elevated CO₂ during development can also lead to reduced fitness and increased post-settlement mortality, due only to reductions in growth rates and energetic reserves (Hettinger et al., 2012; Waldbusser et al., 2015). Some of these mechanisms probably contributed to the observed effects on the population densities of E. castanea. Inferring on the relative importance of direct and indirect effects would, however, require manipulative experiments, which was clearly beyond the scope of the present study.
500 μm mesh used for sampling (Bayne, 1965; Chicharo and Chicharo, 2001). Although larval settlement was not directly measured, this suggests that settlement rates were higher in sites affected by CO2 emissions than in reference sites. The fact that newly settled postlarvae were found only in high CO2 sites (Fig. 5B) also seems to corroborate this hypothesis.

Other populations of the same genus are known to have extreme temporal variations in abundance (i.e., boom-and-bust dynamics - Albano et al., 2016), which implies highly variable settlement rates. Potential spatial or temporal differences in settlement among experimental sites could, therefore, also have influenced the abundance and size-frequency distribution of the populations. The preponderance of small bivalves in naturally acidified environments has previously been observed and interpreted as evidence of continuous recruitment (Metaxas, 2015). As far as we are aware, however, there is no experimental evidence for positive effects of CO2 on larval settlement of calcifying benthic marine invertebrates (Gazeau et al., 2013; Parker et al., 2013), while neutral (Bechmann et al., 2011; Crim et al., 2011) or negative effects (Cigliano et al., 2010; Ricevuto et al., 2012) have been observed. Accordingly, a direct enhancement of bivalve settlement by elevated levels of CO2 seems extremely unlikely. Any positive effect was probably indirect, regulated by density-dependent control mechanisms.

Intraspecific interactions, like competition for resources (i.e., food or space) or predation (i.e., ophryan and larviphy), have previously been recognized as important regulating mechanisms for population size in filter-feeding bivalves, by imposing a negative relationship between recruitment success and densities of adult conspecifics (Andre et al., 1993; Andre and Rosenberg, 1991; Comtet and Desbruyères, 1998; Vania et al., 2014). It seems likely, therefore, that the high densities of adults in reference sites could lead to reduced rates of recruitment.

Chemical cues released by adult conspecifics are known to induce larval settlement and metamorphosis in several species of bivalves (Gosling, 2015; Porri et al., 2007). While the terms settlement and metamorphosis are often used interchangeably (Hadfield and Paul, 2001), they actually designate separate processes: metamorphosis is a definitive morphogenetic event; settlement, on the other hand, is reversible, as pediveligers and plantigrades of some species can swim up from the benthos and settle again in a new location (Bayne, 1964; Hadfield and Paul, 2001). Given the absence of E. castanea adults, chemical cues may be lacking from sites with elevated CO2, which could have contributed to delayed metamorphosis and/or increased post-settlement dispersal, while having no effect on larval settlement itself. Although these potential mechanisms were not investigated in our study, the simultaneous occurrence of high levels of intraspecific predation (in reference sites) and the lack of chemical cues from adult conspecifics (in sites affected by CO2) could explain the observed trends in the abundance of postlarvae and juveniles.

According to Morton (1990), E. castanea in the Azores reach sexual maturity at a shell length of 3.5 mm for females and 5.5 mm for males. The survival threshold in sites with elevated levels of CO2 was clearly under that size range, suggesting that individuals at those sites are not producing offspring. Being an open system, however, larvae may originate from non-acidified sites, and only after settlement become subjected to the effects of elevated CO2 (Dupont and Pörtner, 2013; Kroeker et al., 2013b). Overall, and even though reproduction was not investigated in our study, these results suggest that populations of E. castanea in high CO2 sites are probably sink populations.

5. Conclusions

One of the advantages of conducting ocean acidification research using natural CO2 gradients is that the responses observed are not restricted to the direct effect of carbonate chemistry, but also include the indirect consequences of trophic interactions and changes in habitat complexity (Fabricius et al., 2015; Hale et al., 2011; Range et al., 2012). In the future, transplant experiments including measurements of mineralization and dissolution of the shell, physiological, metabolic and genetic endpoints could enlighten and complement the responses observed in this study. There is also a need to consider other stressors (e.g., temperature, nutritional conditions, pollution), as future ecosystem changes will be driven by the synergistic effects of multiple stressors, instead of their isolated action (Bagnini et al., 2014; Hale et al., 2011; Kroeker et al., 2013b). Future work should also include environmental benthic variables, such as sediment pH and temperature, as these can significantly vary from water column values (Clements et al., 2016; Dashfield et al., 2008).

We have shown that shallow–water volcanic CO2 emissions in the Azores can be used to assess the effects of ocean acidification in central Atlantic benthic ecosystems, complementing similar studies in the region (Viotti et al., 2019) and in other parts of the world (Fabricius et al., 2015; Hall-Spencer et al., 2008; Harvey et al., 2018). Our results also emphasized the importance of food availability (i.e., chlorophyll a) in mediating the effects of CO2 on filter feeders. Although E. castanea is not commercially exploited, these findings raise concerns about the effects of ocean acidification on coastal bivalve populations with similar life history traits and on coastal ecosystems that depend on them for food provision or commercial exploitation.

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CRediT authorship contribution statement

Marta Martins: Investigation, Formal analysis, Visualization, Writing – original draft. Marina Carreiro-Silva: Funding acquisition, Conceptualization, Methodology, Resources, Formal analysis, Writing – review & editing, Project administration. Gustavo M. Martins: Funding acquisition, Conceptualization, Methodology, Resources, Investigation, Formal analysis, Writing – review & editing, Project administration. Joana Barcelos e Ramos: Funding acquisition, Conceptualization, Methodology, Investigation, Resources, Formal analysis, Writing – review & editing, Project administration. Éfrica Vieiros: Funding acquisition, Conceptualization, Methodology, Resources, Formal analysis, Writing – review & editing, Project administration. Ruben P. Couto: Funding acquisition, Conceptualization, Methodology, Investigation, Formal analysis, Resources, Writing – review & editing, Project administration. Hélio Parra: Investigation, Formal analysis. João Monteiro: Investigation, Formal analysis, Writing – review & editing. Francesca Gallo: Methodology, Investigation, Formal analysis, Writing – review & editing. Catarina Silva: Investigation, Formal analysis, Writing – review & editing, Alexandra Teodósio: Funding acquisition, Resources, Investigation, Writing – review & editing. Katja Guillon: Methodology, Investigation, Resources, Writing – review & editing. Jason M. Hall-Spencer: Funding acquisition, Conceptualization, Methodology, Investigation, Writing – review & editing, Supervision. Luís Chicharo: Funding acquisition, Methodology, Investigation, Formal analysis, Writing – review & editing, Supervision. Pedro Range: Funding acquisition, Conceptualization, Methodology, Investigation, Formal analysis, Visualization, Writing – review & editing, Project administration, Supervision.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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