The physiological response of the deep-sea coral *Solenosmilia variabilis* to ocean acidification

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ABSTRACT

Several forms of calcifying scleractinian corals provide important habitat complexity in the deep-sea and are consistently associated with a high biodiversity of fish and other invertebrates. How these corals may respond to the future predicted environmental conditions of ocean acidification is poorly understood, but any detrimental effects on these marine calcifiers will have wider impacts on the ecosystem. Colonies of *Solenosmilia variabilis*, a protected deep-sea coral commonly occurring throughout the New Zealand region, were collected during a cruise in March 2014 from the Louisville Seamount Chain. Over a 12-month period, samples were maintained in temperature controlled (∼3.5 °C) continuous flow-through tanks at a seawater pH that reflects the region’s current conditions (7.88) and an end-of-century scenario (7.65). Impacts on coral growth and the intensity of colour saturation (as a proxy for the coenenchyme tissue that covers the coral exoskeleton and links the coral polyps) were measured bimonthly. In addition, respiration rate was measured after a mid-term (six months) and long-term (12 months) exposure period. Growth rates were highly variable, ranging from 0.53 to 3.068 mm year⁻¹ and showed no detectable difference between the treatment and control colonies. Respiration rates also varied independently of pH and ranged from 0.065 to 1.756 µmol O₂ g protein⁻¹ h⁻¹. A significant change in colour was observed in the treatment group over time, indicating a loss of coenenchyme. This loss was greatest after 10 months at 5.28% and could indicate a reallocation of energy with physiological processes (e.g. growth and respiration) being maintained at the expense of coenenchyme production. This research illustrates important first steps to assessing and understanding the sensitivity of deep-sea corals to ocean acidification.

INTRODUCTION

Deep-sea corals (Phylum Cnidaria) are an abundant and diverse group that are found worldwide and, like their shallow water counterparts, several groups are characterised by their ability to form calcium carbonate skeletons. Corals are vulnerable to environmental change resulting from anthropogenic disturbances such as climate change, ocean acidification (OA) (Guinotte et al., 2006; Turley, Roberts & Guinotte, 2007) and fishing.
(Clark & Rowden, 2009; Clark et al., 2015b). To date, no research has been carried out in the New Zealand region on the impacts of climate change, including OA, on this important group of scleractinian stony corals. One study by Thresher et al. (2011) in nearby Australian waters investigated the effects of chronic low carbonate saturation levels on the distribution, growth and skeletal chemistry of several deep-sea corals off southeastern Tasmanian seamounts and found that the distribution of scleractinian corals is constrained by low carbonate saturation levels.

Deep-sea corals are generally found in water temperatures between 4 and 12 °C (Roberts, Wheeler & Freiwald, 2006; Buhl-Mortensen & Mortensen, 2005). This largely corresponds to relatively shallow depths (between 50 and 100 m) at high latitudes, and greater depths (up to 4,000 m) at low latitudes (Roberts, Wheeler & Freiwald, 2006). However, solitary cup corals (e.g., Caryophyllia antarctica, Gardineria antarctica and Flabellum impensum) can be found up to 1,000 m deep in the high latitude waters of Antarctica (Parker & Bowden, 2010). Compared to the large numbers of shallow-water, reef building corals that have been described, only 10 deep-sea scleractinian reef-building species have been described globally (Cairns, 1979; Freiwald et al., 2004).

The South Pacific region, including New Zealand, supports a broad diversity of various deep-sea coral fauna (Williams et al., 2010), the majority of which live between depths of 200 and 1,200 m (Tracey et al., 2011). The scleractinain corals in the region are often found on elevated hard substrate with topographic complexity such as seamounts, knolls, on slope margins, ridges and canyons (Tracey et al., 2011), where they are in an advantageous position to feed in high current areas on particulate organic matter (Wolanski & Hamner, 1998). Branching forms of the scleractinian corals create three-dimensional reef structures in the deep, and these provide key biogenic habitat and refuge for many deep-sea invertebrates, fish and sharks. The community composition of various invertebrates associated with coral-reefs is well described in the literature (e.g., see Henry, Davies & Roberts, 2010). Fish have been seen on, or in close proximity to, stony and other habitat-forming deep-sea corals (Bongiorni et al., 2010; Soffker, Sloman & Hall-Spencer, 2011; Fosså et al., 2012; Purser et al., 2013; Biber et al., 2013; Milligan et al., 2016), and the benefits of deep-sea reef habitats to shark species have also been reported (Henry et al., 2013).

Atmospheric concentrations of carbon dioxide (CO₂) have increased since pre-industrial times due to anthropogenic emissions. The ocean acts as a carbon sink, absorbing this CO₂, but results in changes to the chemistry of seawater, including a reduction in pH and the availability of free carbonate ions (Hoegh-Guldberg et al., 2007). By the end of this century, OA is expected to cause a decline in oceanic pH by 0.2–0.3 pH units (IPCC, 2013). This is in addition to the pH drop of 0.1 units, which has already been observed since pre-industrial times (Friedrich et al., 2012). OA is enhanced at low temperatures and high pressure, conditions experienced in the deep-sea (Orr et al., 2005; Roberts, Wheeler & Freiwald, 2006; Law et al., 2018). Already impacted by trawling (Clark & Rowden, 2009; Clark et al., 2015a; Clark et al., 2015b), scleractinian corals in deep, cold-water environments are predicted to be affected by global change, such as OA, much sooner than corals in surface waters of more temperate regions (Guinotte et al., 2006; Turley, Roberts & Guinotte, 2007; Sweetman et al., 2017). By 2100, under the high CO₂ RCP8.5 scenario (IPCC, 2013), pH
reductions of >0.2–0.3 pH units from current levels are expected in 23% of deep-sea canyon regions and on 8% of seamounts, the key areas where deep-water corals are typically found (Gehlen et al., 2014). While the response of deep-sea corals to OA, and resulting low carbonate saturation levels, is poorly understood, research such as that by Thresher et al. (2011), and research investigating carbonate saturation horizons in New Zealand waters by Bostock, Mikaloff Fletcher & Williams (2013), indicate that deep-sea corals will be sensitive to such environmental changes. A recent synthesis assessed the potential threat posed by OA to the diversity and productivity of New Zealand marine ecosystems, including corals, and highlighted the knowledge gaps in understanding the impacts (Law et al., 2018).

The skeletons of deep-sea scleractinian corals are most commonly composed of aragonite, the more soluble polymorph of carbonate, which makes them vulnerable to OA-induced dissolution (Anthony et al., 2008). The water’s suitability for carbonate deposition is determined by the carbonate saturation state (Ω). As Ω reduces, the formation of carbonate skeletons becomes increasingly difficult, and the increased energy requirements of calcification can ultimately threaten an organism’s survival. The depth at which seawater is saturated with aragonite is termed the aragonite saturation horizon (ASH). Below this depth, the ocean is under-saturated with respect to aragonite. From studies of distribution (e.g., see Bostock et al., 2015), it is suggested that most deep-sea coral species can probably tolerate some aragonite undersaturation (∆ΩAr ~0.8–0.9). These authors suggested that scleractinian corals should be present in >1% of stations down to 1,800 m water depth, and that some species (e.g., Solenosmilia variabilis) may be tolerant of ∆ΩAr ~0.07, but they concluded it is unclear how deep-sea corals might respond to future OA.

The rapid shoaling of the ASH over the last two decades, measured at 1–2 m yr⁻¹ (Feely et al., 2012), represents a significant threat to deep-sea corals, as it is anticipated it will become challenging for these ecosystem engineers to construct and maintain their skeletons in water under-saturated with respect to aragonite (Guinotte et al., 2006; Tracey et al., 2013; Bostock et al., 2015). Globally, more than 70% of the present deep-sea coral communities will be subject to under-saturated conditions by the end of this century (Guinotte et al., 2006). However, the models used to simulate past and future changes in OA have the largest uncertainties in the Southern Ocean (e.g., Bopp et al., 2013; Orr et al., 2005). Within the New Zealand region, 95% of the habitat-forming scleractinian corals are found above the ASH (Tracey et al., 2013; Bostock et al., 2015). Recent work suggests that, during the present Anthropocene, the ASH has already shoaled by 50 to 100 m over much of the New Zealand Exclusive Economic Zone (Mikaloff-Fletcher et al., 2017). This indicates that the proportion of the region with a carbonate chemistry favourable to aragonitic calcifiers has already shrunk considerably (Mikaloff-Fletcher et al., 2017). Further, Bostock et al. (2015) noted that some scleractinian corals lie below the ASH (i.e., in a zone where conditions seem unfavourable for their growth). These authors hypothesised that previous shifts in the ASH could explain this unexpected result; corals could have established when the ASH was deeper and the waters were supersaturated with aragonite at that depth, and then adapted as the ASH shoaled (Mikaloff-Fletcher et al., 2017). Alternatively, the pattern could indicate that these corals have some capacity to withstand or acclimate to changes in ocean chemistry.
Globally, most experimental work on the effects of OA on corals has been on shallow water species and many studies note significant negative responses to OA. In meta-analyses that included studies of shallow water corals, Kroeker et al. (2010) and Kroeker et al. (2013) highlighted corals as one of the more vulnerable groups to OA. For example, calcification rates may decrease and carbonate dissolution rates may increase in shallow-water Pacific corals when pH is reduced only slightly (pH 7.85–7.95), with substantial impacts when the pH is reduced to 7.60–7.70 (Anthony et al., 2008). This pattern of decreasing calcification rates at lower carbonate concentrations is widely observed in shallow-water corals (Marubini et al., 2008; Herfort, Thake & Taubner, 2008). Cellular level effects on shallow water corals have also been observed, where the photosynthetic activity of the endosymbiont is tightly coupled with the ability of the host cell to recover from cellular acidosis after exposure to OA (Gibbin et al., 2014).

In contrast, there are fewer studies on the impacts of OA on deep-sea corals. A synthesis by Maier, Weinbauer & Gattuso (in press) reports that the response of only five deep-sea coral species (Madrepora oculata, Lophelia pertusa, Desmophyllum dianthus, Dendrophyllia cornigera and Caryophyllia smithii) to OA have been investigated. Most of the stony coral studies outside of the Mediterranean region have been confined to one species: L. pertusa (see Table 1 in Maier, Weinbauer & Gattuso, in press). These studies have examined effects of OA (through manipulation of pH or partial pressure of CO$_2$ (pCO$_2$)) after short (∼24 h) and long term (∼10–12 months) exposure, on measures such as calcification, metabolism and skeleton strength (Maier et al., 2009; Hennige et al., 2014; Movilla et al., 2014). Here we took a long-term approach (12 months) to assess the impacts of the projected end-of-century OA scenario on the physiology of an abundant, habitat-forming scleractinian coral species (Solenosmilia variabilis) from New Zealand and the wider southeast Pacific region (Tracey et al., 2011; Thresher et al., 2011). This species is fragile, long-lived, and late to mature (Thresher et al., 2013; Fallon, Thresher & Adkins, 2014; H Neil, DM Tracey, DM Tracey, P Marriott & MC Clark, 2010, unpublished data), and any negative impact of OA on this species could have wider ecosystem consequences.

**METHODS**

**Live sampling of Solenosmilia variabilis**

Field sampling of protected corals was approved by the Department of Conservation (permit number: 35099-CAP) and coral samples were landed under the authority of the Ministry for Primary Industries (permit number: B2014/61361).

Live colonies of *S. variabilis* were sampled during March 2014 from the Louisville Ridge, 700 km east of New Zealand (Fig. 1). Colonies were sampled in depths ranging from 1,220 to 1,370 m from each of two seamount-like guyot features (referred to as seamounts throughout) (Table 1) using an epibenthic sled deployed from the National Institute of Atmospheric Research (NIWA) research vessel RV Tangaroa.

Upon retrieval of the sled, multiple live coral colonies were immediately placed in a chilled bin of seawater and then transferred to an on-board aquarium with a continuous flow-rate (∼50 L h$^{-1}$) of unfiltered seawater, maintained at ∼5 °C. No feeding took place.
throughout the three-week voyage as it was assumed that the corals would obtain sufficient food from unfiltered surface water.

Conductivity, Temperature and Depth (CTD) casts (Seabird 911: Seattle, WA, USA) and water samples were conducted at the sampling sites in order to characterise local seawater and to inform the experimental conditions (Table 1). One CTD cast was taken per site, and the following water samples (one per site): 500 ml for total alkalinity ($A_T$) and 250 ml for dissolved inorganic carbon ($C_T$). Water samples were preserved with mercuric chloride ($\text{HgCl}_2$). The $C_T$ was determined using coulometric analysis of the $\text{CO}_2$ stripped...
Table 2  Experiment seawater conditions. pH, pCO$_2$ and carbonate parameters (average ± SE) calculated from measured pH, alkalinity, temperature and salinity on two separate dates during the experiment. The pH over the entire 12 month experiment averaged 7.88 ± .00004 (control) and 7.65 ± .00007 (treatment).

| Treatment (target) | pH      | $A_T$ (µmol kg$^{-1}$) | pCO$_2$  | $\Omega_{Ar}$ | $\Omega_{Ca}$ |
|-------------------|---------|------------------------|----------|----------------|--------------|
| pH 7.88 (control) | 7.87 ± 0.0004 | 2,257 ± 28.71 | 591.9 ± 7.04 | 1.11 ± 0.02 | 1.76 ± 0.03 |
| pH 7.65 (treatment) | 7.65 ± 0.001 | 2,260 ± 27.51 | 1,017.5 ± 15.71 | 0.69 ± 0.01 | 1.09 ± 0.01 |

from the seawater sample after acid addition (Dickson, Sabine & Christian, 2007). The accuracy of the method is determined by analysis of Certified Reference Material (provided by Andrew Dickson from Scripps Institution of Oceanography) with every sample batch, and is estimated to be ± 1 µmol kg$^{-1}$. $A_T$ was determined using a closed cell potentiometric titration (Dickson, Sabine & Christian, 2007). The accuracy of the method is determined by analysis of Certified Reference Material (provided by Andrew Dickson from Scripps Institution of Oceanography) with every sample batch, and is estimated to be ± 2 µmol kg$^{-1}$.

In situ pH (total scale) was calculated using measured $C_T$, $A_T$, temperature and salinity, and Mehrbach equilibrium constants refit by Dickson & Millero (1987). This calculated pH was used to set the ambient pH conditions for the experiment.

Experimental set up

Once ashore, 12 coral colonies were transferred to NIWA’s Marine Environmental Manipulation Facility (MEMF), Wellington, where they were held in flow through seawater at the temperature measured at the collection site (3.5 °C). After a stabilisation period, the large colonies were carefully broken into small portions to achieve 54 colony fragments comprising live polyps and adjacent branchlets. The number of fragments achieved from each colony ranged from two, up to six. Each colony was kept out of the water for no longer than 1 minute during this process. Each fragment (~4–6 cm in length) was then attached in a fixed orientation to a piece of plastic mesh. Colonies were then randomly assigned to one of 18 identical tanks (4 L) with three coral fragments per tank, whilst also ensuring that fragments originating from the same colony were not included in the same tanks. Corals were maintained in darkness throughout the stabilisation and experimental period.

The tanks were fed seawater via a continuous flow-through system (~130 mL m$^{-1}$). Seawater pH was 7.88, to mimic conditions measured at the collection sites (Table 1). The corals were fed twice weekly with a 3 mL mixture consisting of 10% commercial coral food (JBL Koralfuid, Neuhofen, Germany) and 10% commercial shellfish diet (larval shellfish diet 1800, Reed Mariculture, Campbell, CA, USA), that was diluted to the required volume with 1 µm filtered seawater (FSW). Corals were maintained in this tank set-up, with regular feeding, for three months before the experiment began, increasing the likelihood that each colony had a similar nutritional status at the beginning of the experiment.

After three months, the experiment was initiated, with nine control tanks and nine treatment tanks established. Corals in the control group were exposed to ambient pH 7.88 (pCO$_2$ 519 ppm). In comparison, treatment corals were exposed to low pH 7.65
(pCO₂: 920 ppm) (Table 2). The reduced pH level was based on projected changes to seawater pH through to the year 2100 (Bopp et al., 2013; IPCC, 2013; Orr et al., 2005). The pH in the treatment tanks was reduced gradually over three days until it reached the treatment value. Temperature was held at 3.5 °C in all tanks.

**Seawater manipulation and measurement**

FSW from Wellington Harbour, adjacent to the facility, was chilled to 3.5 °C and fed to separate header tanks before being delivered to the experimental tanks at 130 mL min⁻¹ in a flow-through system. The pH was adjusted through the diffusion of food grade CO₂, which was controlled using Sensorex S150C pH probes (Garden Grove, CA, USA). The pH probes in each header tank were calibrated regularly with TRIS and AMP buffers. Water samples were taken from each header tank on two occasions during the 12-month experiment, preserved with HgCl₂, and analysed for determination of A⁻ as described above. These measurements of pH (on each day the water samples were taken) and A⁻, along with temperature and salinity, were used to calculate pCO₂ and ΩAr of each experimental treatment using the refitted (Mehrbach et al., 1973) equilibrium constants (Dickson & Millero, 1987).

**Evaluating Solenosmilia variabilis responses**

Responses were assessed using a variety of measures at regular intervals over the 12-month experiment. At the beginning of the experiment, all coral fragments were photographed and buoyant weighed. Subsequently, at bimonthly intervals over a 10-month period, measurements of polyp mortality (via live polyp counts), linear skeletal extension and/or three-dimensional step-wise growth, (referred to as linear growth throughout), and loss of coenenchyme tissue were made. The coenenchyme is the outer tissue covering the coral skeleton, that links the coral polyps and provides protection for the developing exoskeleton; loss of this tissue was evaluated via changes in colour saturation (detailed below). Respiration rate (O₂ consumption) was measured on two occasions, at six and 12 months.

**Polyp mortality**

Polyp mortality was measured every two months by making a visual count of the number of live polyps on each fragment. Each tank had three fragments and polyp mortality was averaged for each tank to get a single average per tank (n = 9). The total percentage remaining of the initial polyp count at each time point was then calculated using the following equation:

\[
100 - \left[ \left( \frac{P_1 - P_J}{P_1} \right) \right] \times 100
\]

Where \( P_1 \) is the polyp count taken at the first time point and \( P_J \) is the polyp count at each of the subsequent \( J \)th time points.

**Linear growth**

Each coral fragment was photographed at bimonthly intervals to obtain a measure of linear growth. Because fragments were cable-tied in a fixed position, they remained in the same...
orientation throughout the experiment, and it was possible to locate and measure the same branch through time. From the digital images, linear growth was determined by selecting an easily identifiable feature on the colony fragment, such as a branching point or a specific linear growth feature. Measurements were then taken from this distinctive point, along the axis of linear growth, to the area just below a live polyp where the calcification process occurs. Measurements were made using the software ImageJ © (Schneider, Rasband & Eliceiri, 2012). Where possible a maximum of four such measurements were taken for each fragment. Where multiple measurements were taken, these were then averaged to achieve a single linear growth rate for each fragment. Each tank had three fragments and the single fragment linear growth rates were averaged for each tank to get a single average per tank ($n = 9$). Only branchlets that were $\sim 2–5$ cm long at the beginning of the experiment were selected for measurement.

**Tissue loss**

Images taken to measure linear growth rates were also analysed to determine colour saturation, which was used as a proxy for the coenenchyme covering the branch and polyp areas of the coral skeleton. Our method used to assess colour change was based on that of Winters et al. (2009). Images taken during the experiment were cropped to remove the background and then colour-profiled using the colour histogram plugin on ImageJ. The entire 2D image of each coral fragment was profiled at each time point. This profile provides a mean value of intensity for each of the red, green and blue colour channels. A pilot study was used to confirm that a loss in intensity of the red colour channel corresponded to a loss of coenenchyme (see Supplementary Information).

The relative intensity for the red colour channel was calculated using the following equations:

$$T = R + G + B$$
$$R_r = \frac{R}{T}$$
$$G_r = \frac{G}{T}$$
$$B_r = \frac{B}{T}$$

where $T$, the total intensity of an image; $R$, mean intensity of the red channel; $G$, mean intensity of the green channel; $B$, mean intensity of the blue channel; and $R_r$, $G_r$ and $B_r$, relative intensity of the red, green and blue channels, respectively (Winters et al., 2009).

Calculating the percentage of relative brightness for the red colour channel, rather than using the mean brightness, suppresses the influence that any changes in illumination, exposure or internal camera processing, may have on the brightness of each channel (Richardson et al., 2009).

The percentage change in relative intensity of the red colour channel was then calculated using the following equation:

$$R_R = 100 \times \frac{(S_{R1} - S_{R2})}{(S_{I1} \times \frac{T_1}{T_2})}$$

Gammon et al. (2018), PeerJ, DOI 10.7717/peerj.5236
where $R_R$, the relative intensity of the red colour channel; $S_{R1}$, the mean intensity of the red colour channel at time point one; $S_{R2}$, the mean intensity of the red colour intensity at time point two; $T_1$, time point one and $T_2$ time point two.

### Respiration rate

At six and 12 months, one fragment per tank was randomly selected ($n = 9$ for each treatment and time point), and respiration rates measured. Respiratory oxygen consumption was measured in a 500 mL chamber, sealed by an o-ring (Fig. 2).

Each chamber was equipped with a magnetic stirrer to ensure homogeneity of oxygen ($O_2$) around the coral fragments. A glass vial with a Presens Pst 3 $O_2$ sensor (Regensburg, Germany) glued to its end was inserted through a hole in the chamber lid, so that it made contact with seawater in the chamber. The $O_2$ sensor was two-point calibrated before each run, using 0% and 100% saturated seawater; 0% saturated seawater was obtained by dissolving 1 g of sodium sulphite (Sigma-Aldrich, St. Louis, MO, USA) in seawater, and 100% saturation was achieved by bubbling air through seawater for 30 min. The chambers were placed in a 3.5 °C water bath and kept in darkness. Each coral fragment was left to settle in its chamber for a minimum of 20 min before the chamber was sealed. The chambers remained in the water bath for the duration of the measurement and were kept in darkness to prevent any photosynthetic activity in the seawater. Total $O_2$ used by each coral fragment was calculated as the difference between the initial and final oxygen concentrations measured within each chamber.

Each run consisted of five incubation chambers, each housing a different coral fragment. The duration that each fragment was kept in a chamber varied depending on the coral’s respiration rate, a period ranging from 5 to 7 h. Measurements in the chambers were...
terminated if the O₂ saturation dropped below a pre-determined 90%. The water volume within each chamber was measured at the end of each experiment.

The O₂ concentration in each chamber at the start and end of the experiment was standardised to µg L⁻¹ and an hourly rate of O₂ consumption calculated for each individual. The protein content per individual was used to normalise the respiration rate (µg O₂ mg protein⁻¹ h⁻¹). Samples were initially frozen and the frozen tissue removed from the skeleton matrix with an airbrush and transferred to a snap-lock bag containing 5 mL of distilled water. The protein slurry produced was then poured into a 250 mL beaker. The snap-lock bag was rinsed into the beaker twice with 5 mL of distilled water to remove any residual protein. The protein slurry was homogenized further using an electric homogenizer (Proxxon micropower driver, Föhren, Germany), and the total quantity of homogenized material noted. A 5 µL subsample of homogenized protein slurry was transferred to a 96-well plate and analysed with the Coomassie Brilliant Blue protein assay (Bradford, 1976), and a spectrophotometer (EnSpire 2300 Multilabel Plate Reader: PerkinElmer, Waltham, MA, USA). The protein concentration of each 5 µL sub-sample was then adjusted for the total volume of each sample and the total protein content of each individual coral fragment calculated.

Statistical analyses
Statistical analyses were carried out using the software package SPSS (Coakes & Steed, 2009). Data were initially tested for normality, and transformed if they did not meet assumptions. A Friedman test was used to analyse data for both polyp mortality and the loss of coenenchyme (data were not normally distributed, and the assumption of normality could not be met using log transformations). Data were categorized into 12 groups which represented each of the monthly time points (zero, two, four, six, eight and 10 months of exposure) for the treatment and control pH samples.

A rm-ANCOVA was used to compare the average linear growth rate of individuals between the control and treatment groups. The difference between the linear growth lengths of each branchlet, for each sample, was compared between each time point and linear growth presented as mm linear extension per year. Seamount and colony of origin were included as covariates to ensure that they had no confounding effects on the response variable.

Respiration data were log transformed to meet the assumption of normality. The significant effect of treatment, and interactive effect of time since exposure with treatment, were tested using a two-way ANCOVA. Respiration chamber, seamount of origin and colony of origin were included as covariates to ensure that they had no confounding effects on the response variable.

RESULTS
At the end of the 12-month experiment, all corals, in both the treatment and control groups, had live polyps, indicating that the experimental conditions were appropriate to maintain viable corals.
pH was maintained at target concentrations for the duration of the experiment. The average pH for the treatment group was 7.650 ± 0.00007 (range: 7.604–7.699), and the average pH for the control group was 7.876 ± 0.00004 (range: 7.823–7.920). These averages are calculated from >2,800 pH probe measurements, taken throughout the 12-month experiment.

**Polyp mortality**
A visible increase in polyp mortality was noted throughout the experiment. No polyp mortality occurred in the first two months in either the control or treatment groups, although it subsequently increased over time. While there was a treatment effect (Friedman test, $\chi^2(11) = 107.769$, $p = 0.001$), post hoc tests (Wilcoxon-signed rank test) showed that these differences were between different time points of the same treatment and that there was no change in polyp mortality within the treatment group, relative to the control. However, from six months onward, polyp mortality was consistently higher in the low pH group. The greatest loss in polyp mortality occurred in the low pH group from four (where colonies still had 92.04% ± 7.45 of their polyps remaining) to six months of exposure (where colonies only had 61.58% ± 7.19 of their polyps remaining). The difference between these two time points represents a loss of 30.46% of initial polyp counts over just a four-month period.

**Linear growth**
The average linear extension rate at the control pH was 1.558 ± 0.226 mm year$^{-1}$ and at the reduced pH was 1.603 ± 0.260 mm year$^{-1}$. Linear growth rates were highly variable between individual coral fragments, ranging from 0.583 to 3.068 mm year$^{-1}$.

Linear growth rate was also independent of time of exposure for both the reduced pH and control groups (rm-ANCOVA, $F_{4,40} = 0.481$, $p = 0.749$ and $F_{4,52} = 0.274$, $p = 0.893$, respectively). The seamount of origin also had no effect on the linear extension rate of the treatment colonies (rm-ANCOVA, $F_{4,40} = 0.769$, $p = 0.552$) or the control colonies ($F_{4,52} = 0.577$, $p = 0.681$). For these reasons, both time of exposure and seamount were excluded from the final analyses, which then found no effect of reduced pH on the linear extension rate of *S. variabilis* (rm-ANCOVA, $F_{1,25} = 0.017$, $p = 0.899$).

**Coenenchyme loss**
While both the control and treatment groups lost colour throughout the experiment, colour loss was significantly greater in fragments held at reduced pH, a finding that was apparent at all time points (i.e., 2, 4, 6, 8 and 10 months; Wilcoxon-signed rank analysis post hoc analysis, Friedman test, $\chi^2(11) = 130.617$, $p = 0.001$; Fig. 3). After two months, the colour intensity of the control group was 97.61% ± 1.933 of that measured at the start of the experiment, while the low pH group retained 94.396% ± 0.738 of its colour intensity. By comparison, at 10 months there was, on average, a difference of 5.28% between the percentage of initial colour remaining between the treatment and control groups.

**Respiration rate**
Coral respiration rates were higher in the control pH than in the low pH at both the six and 12-month time points (Fig. 1) and, for all fragments, were higher at the 12-month time point.
Figure 3  The effect of seawater pH on the loss of coenenchyme tissue of the coral *Solenosmilia variabilis*. Images A–B are of the same colony at control pH (pH 7.88). Images C–D are of the same colony at the treatment pH (pH 7.65). Images A and C were taken prior to the start of the experiment, and images B and D after three months into the experiment. (continued on next page...)

Full-size DOI: 10.7717/peerj.5236/fig-3
Figure 3 (…continued)
Note the marked reduction in the intensity of the red/pink colouration to a pale colour once the coral had been exposed to low pH for several months (D). Photos of *S. variabilis* were analysed for the relative percentage of intensity in the red colour channel (*n* = 17 per time-point per treatment). The mean percentage remaining (± 1 SE) of the initial relative intensity is presented (E). The solid line represents the treatment group and the broken line represents the control group. Significant differences from the control are shown by ** *p* < 0.01 and *** *p* < 0.001 (Wilcoxon-signed rank analysis).

Figure 4  The effects of seawater pH on the respiration rate of *Solenosmilia variabilis*. Respiration rate (µmol O₂ mg protein⁻¹ h⁻¹) of colonies after exposure to reduced pH (pH 7.65; dark grey) or control pH (pH 7.88; light grey) for six- and 12-month exposure. (*n* = 9 for each treatment and time-point; values are means ± standard error (SE)).

For fragments in the control group (pH 7.88), respiration was 179% and 31% higher than for coral colonies exposed to low pH (pH 7.65) after six and 12 months, respectively (Fig. 1). Also of note is that the respiration rates at reduced pH increased by 225% between the six and 12-month time points.

While the statistical analyses indicated that respiration rate was not influenced by pH (two-way ANOVA, *F*₁,₂₄ = 3.200, *p* = 0.086; Fig. 4), there was a significant effect of time, where respiration rates were higher, for both the control and treatment groups, at the 12-month time point (two-way ANOVA, *F*₁,₂₄ = 0.977, *p* = 0.007). There was, however, no interactive effect between pH treatment and time (two-way ANOVA: *F*₁,₂₄ = 0.101, *p* = 0.350).
DISCUSSION

This study investigated physiological responses to reduced pH in *S. variabilis*, a habitat-forming scleractinian coral species common around New Zealand and the wider southeast Pacific region.

*S. variabilis* colonies were maintained for 12 months under reduced pH conditions (pH 7.65, $\Omega_{\text{Ar}} = 0.69 \pm 0.01$), and various aspects of their physiological response were investigated over that time. While the colonies were generally robust to OA conditions (there was no mortality), there was significant loss of coenenchyme tissue cover at low pH (Fig. 3), and indications of effects on respiration rates (Fig. 4). Respiration rate was relatively low in the reduced pH treatment, particularly at the six-month time point when it was 179% higher in control conditions (Fig. 4), although this effect was not statistically significant. There was no treatment effect on mortality of polyps or linear growth rates.

To date, published studies on how OA might influence deep-sea corals have varied results, even within different populations of the same species. This is demonstrated by Georgian et al. (2016) who tested the physiological response to OA of *L. pertusa* colonies from two geographically different populations (Gulf of Mexico, USA and Tisler Reef, Norway). The Gulf of Mexico corals exhibited reductions in net calcification and respiration, while Tisler Reef corals showed only slight reductions in net calcification and elevated respiration. The authors concluded that these differences were likely the result of environmental differences (e.g., depth, pH, food supply) between the two regions. In another experiment on *L. pertusa*, Maier et al. (2009) found that incubating *L. pertusa* for 24 h in seawater with pH lowered by 0.15 and 0.3 units, relative to the ambient level, resulted in calcification being reduced by 30 and 56%, respectively. In another short term study, Hennige et al. (2014) investigated the response of *L. pertusa* to increased CO$_2$ conditions (750 ppm) over 21 days. *L. pertusa* corals exposed to increased CO$_2$ had significantly lower respiration rates than corals in control conditions, but found no corresponding change in calcification rates. In a longer-term study, Movilla et al. (2014) found a decline in the calcification of *D. dianthus* after 314 days of exposure to elevated pCO$_2$ (800 µatm). Interestingly, in another long-term experiment, over 12 months, Hennige et al. (2015) observed a decrease in the structural integrity of dead exposed *L. pertusa* skeleton, when exposed to increased CO$_2$ conditions. Such studies provided a platform for our current study.

The $\Omega_{\text{Ar}}$ was <1 in both the control and treatment waters in our experiment (Table 2). From a broad survey of New Zealand coral species and carbonate saturation, Tracey et al. (2013) and Bostock et al. (2015) identified a strong dependency of coral distribution on $\Omega_{\text{Ar}}$ and $\Omega_{\text{Ca}}$. However, many deep-sea stony corals can cope with some degree of aragonite undersaturation ($\Omega_{\text{Ar}} \sim 0.8–0.9$) with some species tolerant of $\Omega_{\text{Ar}} \sim 0.7$ (Bostock et al., 2015), a value lower than the $\Omega_{\text{Ar}}$ of 0.69 ± 0.01 in our low pH treatment. While it has been noted that some stony corals lie below the ASH (e.g., Bostock et al., 2015; Baco et al., 2017), including in the New Zealand region, such as those found along the Louisville Seamount Chain (Bostock et al., 2015), we did note reduced coenenchyme tissue cover, and indications of elevated respiration rates at these levels.
**Linear growth rates**

This study found no treatment effect on the linear growth rates of corals which were seen to be highly variable, ranging from 0.583 to 3.068 mm y\(^{-1}\). The measured linear growth rate is comparable to results from radiocarbon dating studies of *S. variabilis* by Fallon, Thresher & Adkins (2014), who reported linear growth-rates ranging between 0.84–1.25 mm y\(^{-1}\), and by H Neil, DM Tracey, DM Tracey, P Marriott & MC Clark (2010, unpublished data), who again showed similar linear growth of 0.25–1.3 mm y\(^{-1}\). Linear growth rates that are independent of pH have been found in other deep-sea corals from various *in aquaria* studies, including *M. oculata* (Maier et al., 2013b) and *L. pertusa* (Form & Riebesell, 2012; Maier et al., 2013b). While it was found that *M. oculata* was not affected when pH was manipulated to end-of-century projections, when the partial pressure of CO\(_2\) was reduced to pre-industrial levels, calcification rates in this species increased (Maier et al., 2012). This provides important information about the historical effect of OA on the calcification of deep-sea corals and indicates that the present-day calcification rates may have already declined due to an anthropogenic increase in the concentration of atmospheric CO\(_2\). Although no net effect of OA on linear extension was observed in this study, it is important to note that measurements were only taken after several months and undetected shorter-term changes may have occurred.

**Respiration rate**

Respiration rate was highly variable, ranging from 0.065 \(\mu\)g O\(_2\) g protein\(^{-1}\) h\(^{-1}\) to 1.178 \(\mu\)g O\(_2\) g\(^{-1}\) protein\(^{-1}\) h\(^{-1}\). These results are low compared to respiration rates found by Dodds et al. (2007) for the branching scleractinian *L. pertusa*, who found a respiration rate of about 0.5 \(\mu\)mol g\(^{-1}\) h\(^{-1}\). Here, the respiration rate of *S. variabilis* was relatively low in the reduced pH treatment, particularly at the six-month time point. Interestingly, this difference decreased at the 12-month time point, and this could be indicative of acclimation. A similar response was found by Maier et al. (2013a) in *M. oculata* and *L. pertusa*. The authors attributed the observed increase in respiration rate to an increase in energy supply as a result of regular feeding, thus sustaining an elevated level of coral metabolism. Regular feeding, and its impact on coral metabolism, can mask the effects of OA in experimental work. This was found by Büscher, Form & Riebesell (2017) who concluded that, while the deep-sea coral, *L. pertusa*, is capable of calcifying under elevated CO\(_2\) and temperature, its condition (fitness) is more strongly influenced by food availability rather than changes in seawater chemistry. With the natural habitat of *S. variabilis* being so inaccessible, it is difficult to predict the amount of food, including particulate organic matter and sources of plankton, reaching colonies in situ.

**Tissue loss**

A visible loss of coenenchyme was noted from both the control and treatment colonies in the first two months of the experiment, although this loss was significantly greater from the treatment colonies. Tissue loss in the control group is consistent with stress and colony deterioration, which is not surprising given that deep-sea corals are difficult to maintain in a healthy state *in aquaria*; indeed, to our knowledge, *S. variabilis* has never previously
been maintained for more than a few weeks in this state. The significantly greater effect of reduced pH on the rate of tissue loss highlights that other physiological mechanisms were also playing a part. A loss of tissue when exposed to OA conditions has also been observed in tropical corals (e.g., *Pocillopora damicornis* and *Oculina patagonica*), but the tissues of these two species regenerated when the corals were returned to ambient pH (*Kvitt et al., 2015*). For these shallow, warm-water corals, reduced pH induced tissue-specific apoptosis, a breakdown of coenenchyme and a subsequent loss of the colonial form. The tissue loss of *S. variabilis* seen here could represent the early stages of a similar response and warrants a longer-term study. Interestingly, the same rate of polyp mortality over time was observed in both the treatment and control colonies, while coenenchyme loss was greater in the treatment group. The coenenchyme has a function in connecting each neighbouring polyp and protecting the growing skeleton. A loss of the coenenchyme could mean a shift away from the coral’s ability to produce a colonial three-dimensional matrix (*Hennige et al., 2015*). Reverting to solitary and non-calcifying polyps has been proposed as an evolutionary mechanism which has allowed corals to survive through geological periods of unfavourable calcification conditions (*Kvitt et al., 2015*), and could explain several “reef gaps” in the geological records (*Wood, 1999*).

Alternatively, the observed loss of coenenchyme could represent a reallocation of energy. That is, corals in the treatment group may have been diverting energy away from the maintenance of tissues, allowing them to maintain other metabolic requirements (e.g., linear growth, respiration and reproduction). For this reason, tissue loss in corals is considered a better indicator of physiological stress than skeletal linear growth (*Anthony, Connolly & Willis, 2002*). *Maier et al. (2016)* show that the energy required for calcification in *M. oculata* is a small fraction (~1-3%) of overall metabolic requirements. Assuming that the energy requirements for calcification in *S. variabilis* are similar, this substantiates our comment that tissue loss may be a better indicator of physiological stress than linear growth and partly explains why this study found no treatment effect on the linear growth rates of corals.

## CONCLUSION

Deep-sea corals are typically difficult to study due to their poor survival rate in laboratory conditions. For this reason physiological studies of their responses to environmental change have been limited to date. The data presented here for *S. variabilis* represent an important first-step towards understanding the biology of this ecologically important species, and to our understanding of the sensitivity of deep-sea corals to OA. In New Zealand specifically, the lack of knowledge of organism responses is well recognised: the potential threat posed by OA to the diversity and productivity of marine ecosystems (including to corals) is classed as medium for vulnerability, low to medium for knowledge of established response, and low for understanding mechanistic response, ecosystem interaction, and interaction of other stressors (*Law et al., 2018*). This study found that *S. variabilis* lost tissue in response to OA and we hypothesize that this could represent a reallocation of energy, with corals diverting energy away from the maintenance of non-essential tissue. It is assumed however,
that an organism would not continue to break down tissues to help support skeletal
three-dimensional linear and/or step-wise growth, as a threshold will ultimately be reached
where the animal becomes seriously compromised. If this were to happen, then there would
be major changes to the structure and function of this species as an important ecosystem
engineer in the deep-sea.

This study has signposted the need to better understand the long-term implications and
mechanisms of OA on colony tissue loss, the most notable effect of decreased pH observed.
To our knowledge, this study is the first to apply a technique of measuring tissue loss to a
deep sea coral in an experiment designed to measure the corals’ response to OA. Studies,
such as this, which find a limited response in those physiological variables which are
typically measured (e.g., respiration and linear growth), should consider what the potential
cost of maintaining those parameters may be. Here we demonstrate an additional measure,
of tissue loss, which could be routinely included in future studies to gain a more holistic
understanding of the organisms’ response. We also recommend that future studies assess
the impact of OA on skeletal morphology and density, which were not assessed here. Such
impacts have the potential to change colony integrity and survival. Combined with ongoing
and more refined modelling work to inform future projections of the ASH and CSH in the
South Pacific, this study nevertheless improves our knowledge on the impacts of OA on
this important and ecologically vulnerable coral group in the New Zealand region.

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Author Contributions
• Malindi J. Gammon conceived and designed the experiments, performed the experiments, analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
• Dianne M. Tracey conceived and designed the experiments, performed the experiments, contributed reagents/materials/analysis tools, authored or reviewed drafts of the paper, approved the final draft.
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• Vonda J. Cummings and Simon K. Davy conceived and designed the experiments, contributed reagents/materials/analysis tools, authored or reviewed drafts of the paper.

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Data Availability
The following information was supplied regarding data availability:
The raw data are provided in a Supplemental File.

Supplemental Information
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