

“Combined effects of ocean warming and acidification on the larval stages of the European abalone *Haliotis tuberculata*”

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Abstract

This study examined the physiological responses of the larval stages of *Haliotis tuberculata*, an economically important abalone, to combined temperature (17°C and 19°C) and pH (ambient pH and -0.3 units, i.e., +200% increase in seawater acidity) in a full factorial experiment. Tissue organogenesis, shell formation, and shell length significantly declined due to low pH. High temperature significantly increased the proportion of fully shelled larvae at 24 hours post-fertilization (hpf), but increased the proportion of unshelled larvae at 72 hpf. Percentage of swimming larvae at 24 hpf, 72 hpf and 96 hpf significantly declined due to high temperature, but not because of low pH. Larval settlement increased under high temperature, but was not affected by low pH. Despite the fact that no interaction between temperature and pH was observed, the results provide additional evidence on the sensitivity of abalone larvae to both low pH and high temperature. This may have negative consequences for the persistence of abalone populations in natural and aquaculture environments in the near future.

Keywords: ocean acidification, global warming, climate change, marine mollusks, abalone larvae

Introduction

Increased CO₂ concentrations due to anthropogenic activities have led to increased oceanic CO₂ absorption, resulting in a phenomenon called ocean acidification (OA) that is concurrent with ocean warming (OW). Current projections suggest that by the year 2100, the average sea surface temperature will increase by 1°C– 3°C and the water pH will drop by 0.1 to 0.3 units (i.e., +30% to +200% increases in acidity; IPCC, 2014). OA has particularly deleterious impacts on marine calcifiers that produce calcium carbonate (CaCO₃) skeletons (Kroeker et al., 2013; Parker et al., 2013). These minerals are experiencing unprecedented decreases in their saturation states due to ongoing OA. This has a suppressive, narcotic effect and causes acidosis, which impairs many physiological processes (Feely et al., 2004; Portner, 2008; Doney et al., 2009). In contrast, the impacts of increased temperatures due to OW can stimulate physiological processes until they reach a threshold (Pörtner, 2010; Przeslawski et al., 2011; Byrne et al., 2011).

The combined impacts of OA and OW have been addressed with contrasting responses, such that the effects of OA and OW are either exacerbated (e.g., Di Santo 2015; D’Amario et al., 2020; Zittier et al. 2018; Rodolfo-Metalpa et al., 2011) or ameliorated (e.g., Kroeker et al. 2014; Davis et al., 2013; Knights et al., 2020; García et al., 2015; Jiang et al., 2018) in the presence of the other stressor (). Meta-analyses have suggested that the combined impacts of OA and OW on species physiology are especially devastating for the larval stages of many species (Przeslawski et al., 2015; Kroeker et al., 2013), presenting a major bottleneck for population persistence under changing oceanic conditions (Przeslawski et al., 2015). As a result, the responses of these early stages to combined OA and OW are now being paid greater attention (Przeslawski et al., 2015). Temperature and OA-related factors, such as pH, are among the paramount environmental factors that dictate the survival, morphology, physiology, and behavior of marine larvae, and in particular,

calcifying larvae (Przeslawski et al., 2015). Among all of the marine calcifiers studied so far, mollusk larvae are the most vulnerable to the effects of OA (Przeslawski et al., 2015; Kroeker et al., 2013).

The recent marine mollusks include over 43,000 recognized species That includes some of the major CaCO_3 producers (Rosenberg, 2014) fulfilling crucial ecosystem functions, such as creating habitat structures and food sources for benthic species (Parker et al., 2013). Therefore, any negative impacts from environmental drivers can result in high ecological and economic consequences (Narita et al., 2012). Abalones are important mollusks that have high ecological and commercial values, and also provide food for human beings (Cook, 2016; Huchette and Clavier, 2004). However, their natural populations have experienced severe declines due to overexploitation (Micheli et al., 2008; Kashiwada and Taniguchi, 2007) and environmental disturbances such as OW and bacterial diseases (Cook, 2016; Travers et al., 2009; Huchette and Clavier, 2004; Morales-Bojórquez et al., 2008). On the other hand, abalone aquaculture is expanding worldwide, understanding the effects of global change drivers on abalone physiology is an important issue for the management of abalone populations in natural and aquaculture environments (Morash and Alter, 2015).

Early life-history stages of abalones are negatively affected by OA and show a high percentage of deformed larvae under low-pH conditions (Byrne et al., 2011; Crim et al., 2011; Guo et al., 2015; Kimura et al., 2011; Zippay and Hofmann, 2010; Wessel et al., 2018; Swezey et al., 2020). Other negative responses that have been reported in abalones in response to temperatures higher than their physiological limit during development include abnormal appearance, reduced growth, and trochophore mortality (e.g., Leighton, 1974; Pedroso, 2017). Although several studies have shown larval abalones to be highly sensitive to OA (Santander-De Leon et al., 2018; Crim et al., 2011;

Tahil and Dy, 2016; Guo et al., 2015; Wessel et al., 2018), the combined effects of OA and OW are still not understood (Gao et al., 2020). Based on the limited data available, OA and OW may have deleterious impacts on abalone populations. Indeed, *Haliotis coccoradiata* embryos were dramatically affected by a combination of high temperatures (+2 to +4°C compared with control 20°C) and acidified conditions (-0.4 to -0.6 pH units), with only a small percentage surviving (Byrne et al., 2011).

Haliotis tubercula is a commercially important species in Europe, for which rearing over the whole life cycle is controlled in aquaculture (Huchette and Clavier, 2004; Courtois de Viçose et al., 2007). As for most marine molluscs, abalone species display a pelago-benthic life cycle with a larval planktonic stage followed by a critical metamorphosis into the benthic juvenile, making the specie highly sensitive to environmental changes (Byrne et al. 2011). The impacts of OA on all stages of the European abalone, *Haliotis tuberculata* (Linnaeus, 1758) have been recently studied (Wessel et al., 2018; Auzoux-Bordenave et al., 2020; Avignon et al., 2020). All the aforementioned studies reported adverse impacts of OA on *H. tuberculata*, especially on the shell growth and calcification. However, *H. tuberculata* larvae were more impaired by OA than was any other life stage, showing significantly adverse effects in survival rate, morphology and development, growth rate and shell calcification (Wessel et al., 2018). While the impacts of OA on abalone larvae are worrying enough, high temperatures present another emerging threat for *H. tuberculata* populations (Travers et al., 2009; Huchette and Clavier, 2004). Furthermore, there is a lack of knowledge on how climate change will modify *H. tuberculata* larval settlement, behaviour and physiology in early life stages.

In this study, we investigated the combined effects of OA and OW on the early life stages of *H. tuberculata* in a full factorial experiment. Abalone larvae were exposed to four OA and OW scenarios: ambient (pH 8.08.0 and 17°C); combined (pH 7.77.7 and 19°C); and individual effect

scenarios (pH 8.08.0 and 19°C; pH 7.77.7 and 17°C) throughout the five days of larval development. We focused on three key larval stages: the trochophore stage, characterized by the set-up of the larval shell at 24 hours post-fertilization (hpf); the mature veliger stage (72 hpf); and the premetamorphic veliger stage (96 hpf), which is the last pelagic life stage before larval settlement (Jardillier et al., 2008; Auzoux-Bordenave et al., 2010). To better understand both the individual and combined effects of OA and OW, we investigated several biological parameters involved in growth, physiology and behavior throughout the larval development cycle of *H. tuberculata*.

Materials and methods

Abalone larvae production

The parental *H. tuberculata* stock was composed of wild broodstock (3 females and 7 males) collected from Saint Quay Portrieux (Brittany, France), and farmed abalones (4 females and 5 males) collected from an offshore sea-cage structure at the France *Haliotis* abalone farm (48°36'50 N, 4°36'3 W; Plouguerneau, Brittany, France). Mixing wild and farmed broodstocks prevent inbreeding and production of enough larvae for the experiment. In France *Haliotis* farm, after a 10-month period in nursery tank, abalone are placed in sea-cages and raised with similar environmental conditions as the wild abalone apart the protection from the predator attacks and high density until the age of 4 years. They are fed with algae collected on the shore, and submitted to the same temperature and pH as the wild broodstock.

The wild and farmed abalones were transferred from the sea and sea cages to the farm, respectively.

There were given time to acclimatize to the farm's conditions over three months with natural

running seawater maintained at 15°C and pumped twice a day from the sea, with *ad libitum* feeding to assure optimal reproduction maturity. Spawning was induced at the France *Haliotis* farm following usual procedures. Briefly, abalones were detached from the rearing aquarium and placed individually into 5-L buckets. Spawning was stimulated with ultraviolet light while gradually heating the filtered seawater from 18°C to 21°C over the course of 1 h. Abalones were allowed to spawn for a maximum of 5.5 h from the start of the experiment. Once spawned, the eggs were pooled and then divided into 12 batches. Spermatozoa from each of the 12 males (one male per batch of eggs) were added separately to avoid spermatric competition (Harney et al., 2018) at an optimal sperm concentration of approximately 100,000 spermatozoa per egg (Huchette et al., 2004). After 1 h, the fertilized eggs were pooled again and their density was estimated under a binocular microscope. The pooled fertilized eggs were then divided into 12 samples of 900,000 embryos per batch and transferred to the 12 hatching boxes.

Experimental design

To study the individual and combined impacts of OA and OW, we applied a full factorial design of two temperatures (17°C low and 19°C elevated) and two pH values (8.0 high and 7.7 low; with three replicate tanks per treatment (12 tanks in total; Fig. 1). We considered 17°C to be the ambient (“low”) temperature, as it was the temperature experienced by abalones during the summer reproduction period in Northern Brittany. According to Gac et al. (2020), temperature in the Northern Brittany reach maximal summer value of 17°C and present short-term variability of 0.1 to 0.4 °C mainly related to the tidal cycle and the day-time. Similarly, the high pH corresponded to the naturally occurring pH during autumn in Northern Brittany (Qui-Minet et al., 2018), that is, the high pH in this experiment was the ambient pH of the seawater pumped into the France *Haliotis*

site. The high temperature and low pH conditions were chosen based on the global projections for the coming decades (IPCC, 2014). We chose the worst-case pH and temperature scenarios; however, the high temperature treatment in this study is about 1°C lower than the worst scenarios predicted for 2100. Abalone larvae were exposed to the experimental conditions from 1 h post-fertilization until they reached the premetamorphic veliger stage. The temperature was controlled using two central heating systems. For the 17°C treatment and pre-heating of the 19°C treatment, we used an Aquahort Ltd Heat Pump (26 kW; THP26-3). To reach 19°C, we supplemented this pump with a Charot Heat Pump (6 kW; 4911 type). Each replicate was composed of three parts: I) a food-safe plastic head tank (60 L), in which water of the appropriate temperature and CO₂ level was premixed and homogenized with a bubbling system; II) a food-safe 22.5-L hatching box (30 cm x 50 cm x 15 cm), which was supplied with reduced-flow running water from the head tank to avoid flushing the eggs out while maintaining the desired pH and temperature. This box harbored the fertilized eggs to allow them to hatch within the first 18 h. An evacuation pipe was connected to the rearing tank to allow automatic larval transfer once the larvae could swim; this transfer was completed over four hours. III) A 350-L epoxy-coated larval tank, which was supplied by the head tank, to rear hatched larvae that had left the hatching box. Natural 3-µm filtered seawater was allowed to run through the tanks at all times (open circuit). To prevent larvae from leaving the tanks, a 40-µm net was placed at the outflow. The flow rate from the head tanks to the rearing tanks was between 200–400 mL·min⁻¹. All tanks were exposed to an artificial photoperiod of 12 h light:12 h dark, which was provided at a light intensity of 900–1000 lux at the water surface using 68-W natural spectrum light-emitting diode (LED) lamps (Solar Natur, JBL). As pre-settlement abalone larvae are lecithotrophic (non-feeding), they were not fed during the experiment.

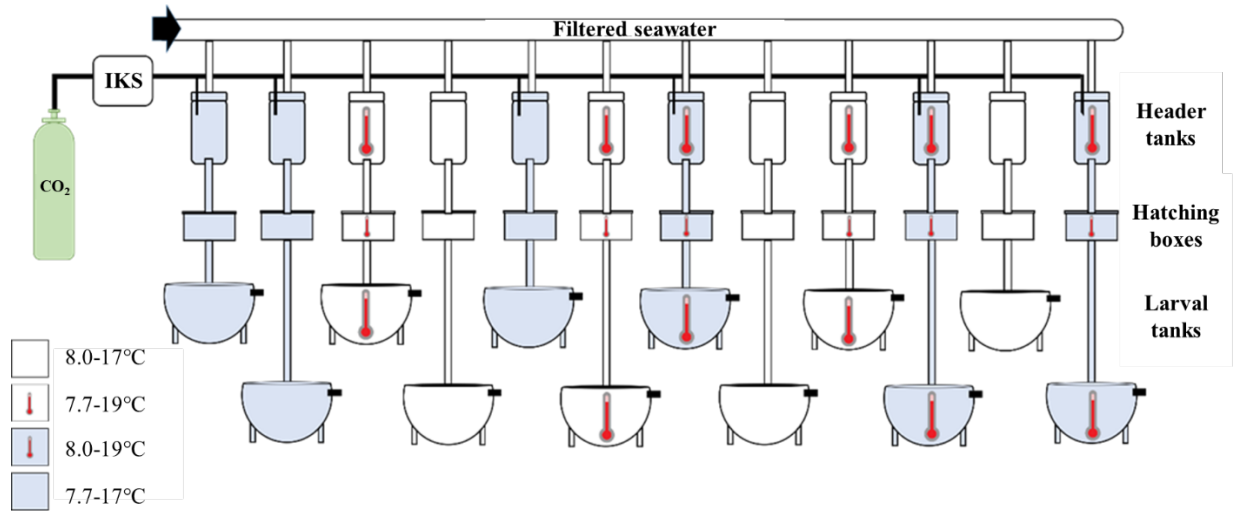


Fig. 1 Simplified design of the experimental system showing the flow of filtered seawater from the header tanks, the styrene hatching boxes, and the 350-L larval tanks (12 in total). Seawater exited the system via the black outflow at the top of larval tanks. IKS is a pH-regulated electrovalve system. The four treatments are indicated using two colors (white and blue) and a red thermometer for high temperatures.

Carbonate chemistry and pH control

Carbonate chemistry and pH control were monitored according to the methods of Avignon et al. (2020). Low-pH seawater was provided by bubbling CO₂ (Air Liquide, Paris, France) into the tanks through electrovalves regulated by a pH-stat system (Aquastar, IKS Computer System, Karlsbad, Germany) (one electrovalve and bubbling CO₂ system per header tank). The desired low-pH value was adjusted to be 0.3 units lower than the high pH, corresponding to natural pH fluctuations along the Northern Brittany coast (the total scale (pH_T) range of 7.9–8.2; Qui-Minet et al., 2018). The values of the pH-stat system were adjusted from daily measurements of the

electromotive force in the header tanks using a pH meter (Metrohm 826 pH mobile, Metrohm AG, Herisau, Switzerland) with a glass electrode (Metrohm Ecotrode Plus, Metrohm AG, Herisau, Switzerland). The electromotive force values were converted to pH units on the pH_T after calibration with Tris-HCl and 2-aminopyridine-HCl (AMP) buffers (Dickson et al., 2007).

Temperature and salinity were measured daily using a portable conductivity meter (ProfiLine Cond 3110, WTW, Oberbayern, Germany). Total alkalinity (A_T) was measured twice ($n = 12$ and $n = 9$) from 50-mL samples taken from each experimental tank. Seawater samples were filtered through 0.7- μm Whatman GF/F membranes, immediately poisoned with mercury chloride, and stored in a dark place at room temperature for later analysis.

A_T was determined from approximately 50 g of weighed samples using a potentiometric titration at 25°C with 0.1 M HCl and using a Titrino 847 plus Metrohm. The balance point was determined by the Gran method (Gran, 1952) according to Haraldsson et al. (1997). The accuracy of this method was $\pm 2 \mu\text{mol}\cdot\text{kg}^{-1}$ and was verified by Certified Reference Material 182, provided by A. Dickson (Scripps Institute of Oceanography, University of South California, San Diego, United States). The seawater carbonate chemistry analysis included dissolved carbonate (CO_3^{2-}), bicarbonate (HCO_3^-), dissolved inorganic carbon (DIC), pCO_2 , aragonite saturation state (Ω_{ar}), and calcite saturation state (Ω_{calc}); these were determined by entering the values of pH_T , A_T , temperature, and salinity into CO2SYS software (Lewis and Wallace, 1998) using constants from Mehrbach et al. (1973) as refitted by Dickson and Millero (1987).

Hatching success

Under pH 8.0 and 17°C, trochophore larvae typically hatch at 18–20 hpf (Jardillier et al., 2008). To estimate the percentage of larvae that hatched in the 12 experimental hatching boxes, we collected all of the unhatched larvae that were deposited at the bottom of the hatching boxes at 24 hpf, after the hatched larvae could swim and had evacuated to the 350-L rearing tank. Nine 1-mL replicates were sampled from a 5-L bucket and fixed with 90% ethanol. Then, the total number of unhatched larvae per tank was counted under a binocular microscope. To estimate the percent hatching success of each tank, we used formula 1.

$$(1) = \frac{\text{number of swimming larvae at 24 hpf}}{(\text{number of swimming larvae at 24 hpf} + \text{number of unhatched larvae})} \times 100$$

Morphometric assessment and percentage of swimming larvae

The post-embryonic developmental stages of *H. tuberculata* were explained in detail by Jardillier et al. (2008) and Auzoux-Bordenave et al. (2010). Our morphometric measurements were focused at 24 and 72 hpf, which corresponded to the free-swimming trochophore and mature veliger stages, respectively. Therefore, only the swimming larvae in the tank water column that corresponded to these larval stages were sampled for further analyses.

The development timing of the larvae was investigated under a binocular microscope at 24, 72, and 96 hpf to verify the larval stages before sampling according to Jardillier et al. (2008) and Auzoux-Bordenave et al. (2010). At 24 and 72 hpf, the total number of larvae swimming in each tank was estimated from six 10-mL replicates per tank ($n = 18$ per treatment). For morphometric, birefringence, and scanning electron microscopy (SEM) analysis, 10–15 L of seawater containing larvae was collected from each tank and filtered through a 40- μm sieve, then aliquoted into 15-mL

tubes. Larvae were concentrated at the bottom of each sample by adding few drops of 70% ethanol; then, the samples were fixed and stored in 70% ethanol for polarized light microscopy and SEM analysis. At 96 hpf, all of the swimming larvae were filtered out of the 350-L tank using a 40- μ m sieve and placed into buckets containing 5 L of seawater adjusted to the same temperature and pH as in their respective treatments. The percentage of swimming larvae at 96 hpf was calculated from eight 1-mL samples ($n = 24$ per treatment). The percentage of swimming larvae was calculated at 24 hpf, 72 hpf and 96 hpf as the total number of swimming larvae in the 350-L tank at each time point divided by the initial number of larvae that hatched.

Slide preparation for morphometric and birefringence analysis

To study morphometric characteristics and birefringence, 12 slides per larval stage (1 replicate per experimental tank) were prepared with the ethanol-fixed larvae (Wessel et al., 2018). Approximately 100 larvae were whole-mounted in about 500 μ L of glycerol. All ethanol was removed before the samples were transferred into the glycerol. The slides were kept at room temperature for 5 to 10 min to allow any remaining ethanol to evaporate and let the larvae settle. Six spots of vacuum gel were deposited at the corners and middle edges of a square coverslip to prevent the larvae from being crushed. After placing the coverslip over the glycerol, the slides were gently sealed with clear nail polish. Each slide contained 150 larvae per treatment per larval stage. The first 50 larvae that were observed per slide, regardless of their orientation, shape, and development, were photographed for morphological analysis with an Olympus binocular microscope (Olympus, Hamburg, Germany) under phase contrast and under polarized light. The same microscope was equipped with polarizing filters for the birefringence analysis. To avoid bias, a coding system was used to prevent the person taking pictures and analyzing the photos from learning which slides corresponded to the particular treatments. All images were taken with a

digital camera (DS-Ri1, Nikon) at 20X magnification and a 40-ms exposure. Images were analyzed in ImageJ software (1.52a).

Morphometric analysis

Larval development, shell formation, and shell size were analyzed for larvae lying on the lateral side (n = 125–133 larvae per treatment). We used semi-quantitative categorizations to analyze larval development, whereby each larva was scored on a scale from 0 to 2 as one of three morphological categories (Table 1; Fig. 2).

Table 1 Semi-quantitative morphological categories of abalone larvae at 24 and 72 hpf. Categories were based on shell formation and soft tissue morphogenesis according to the developmental cycle of *Haliotis tuberculata* at $17 \pm 0.5^{\circ}\text{C}$ (Jardillier et al. 2008).

Score	Shell formation	Soft tissue morphogenesis
2	Normal shell	Tissues normally developed
1	Shell partially and/or abnormally developed	Abnormal or partially developed
0	No shell	Undeveloped

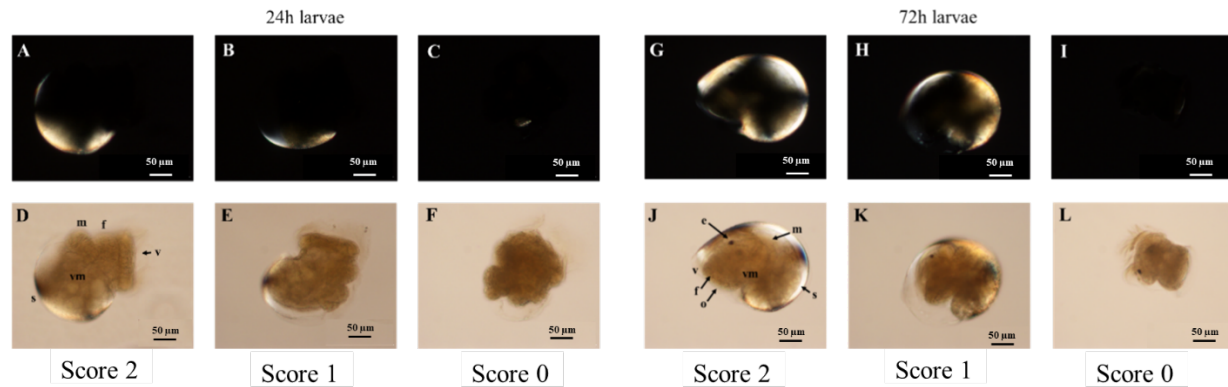


Fig. 2 Polarized microscope (A–C and G–I) and light microscope (D–F and J–L) images showing the morphological variables used for larval scoring at 24 hours post-fertilization (hpf; left) and 72 hpf (right). **A** and **G** Larvae with normal shell development (score = 2; the shell field covers the posterior area of the larval body, including the light half-circled area). **B** and **H** Partially and/or abnormally developed shell (score = 1). **C** and **I** Larvae with no shell development (score = 0). **D** Larvae at 24 hpf with normal tissue development (score = 2). **J** Larvae at 72 hpf with normal tissue development (score = 2). **E** and **K** Partially and/or abnormally developed larvae (score = 1). **F** and **L** Underdeveloped larvae (score = 0). velum (v), mantle (m), shell (s), foot (f), visceral mass (vm), eyes (e), and operculum (o)

The maximum larval length (24 hpf) or total shell length (72 hpf) were measured using ImageJ software and used as indicators of larval size, with all individuals lying on the lateral side (Fig. 3). The mean length of larvae from each tank was calculated, and the average value for all tank replicates was presented as the mean larva size for each treatment (n = 3 tanks per treatment).

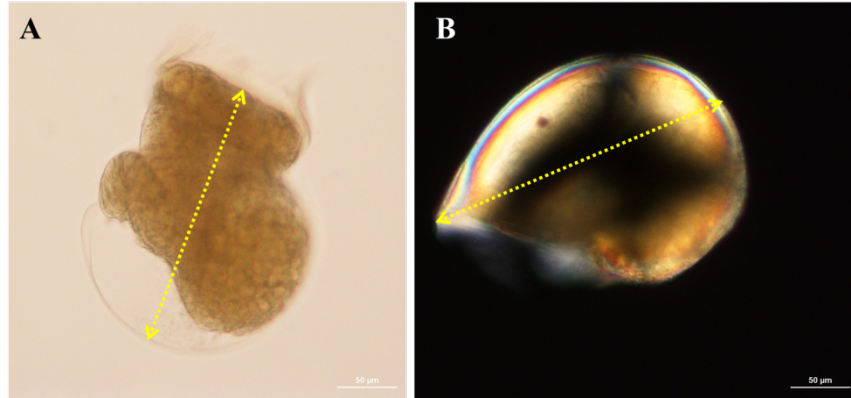


Fig. 3 Larval length measured at **A** 24 hours post-fertilization (hpf) and **B** 72 hpf. Arrow heads show the two ends of the measured lengths. Scales are 50 µm.

Birefringence analysis

We measured birefringence on a scale of 0–255 under cross-polarized light using an Olympus microscope, according to the method described by Wessel et al. (2018). Cross-polarized light that passes through CaCO_3 (an anisotropic material) is double-refracted, and shells with higher CaCO_3 content double-refract more light. Therefore, birefringence intensity can be used as a proxy for evaluating shell mineralization (Noisette et al., 2014). Birefringence was measured for at least 120 larvae per treatment at 24 and 72 hpf using ImageJ. The mean grayscale level (in pixels) was determined for each area of the larval shell showing birefringence, i.e., two zones for 24 hpf larvae, and three zones for 72 hpf larvae.

Scanning electron microscopy

Larvae stored in 70% ethanol were used for SEM analysis (four larvae per treatment). Samples were dehydrated in a series of increasingly concentrated ethanol solutions (90%, 95%, and 100%) and were critical point dried with liquid CO_2 . Finally, the samples attached to the SEM stubs were

gold-coated and observed at 5 kV with a SIGMA 300 FE-SEM scanning electron microscope (Plateau Technique de Microscopie Electronique, MNHN, Concarneau, France).

Larval swimming behavior

At 72 hpf, a 1-L sample was taken from each 350-L tank by carefully dipping a plastic bucket into the middle of the tank and filling it to its maximum capacity. The bucket was covered with a lid to minimize CO₂ exchange and ensure pH stability. The samples were kept in insulated boxes to maintain their respective temperature treatments while they were transported (< 45 min) to the Institut Français de Recherche pour l'Exploitation de la Mer (IFREMER) experiment station (Argenton, France). Upon arrival, the buckets were placed in water baths adjusted to their respective treatment temperatures to recover from their transport. After two hours, the samples were examined for the behavioral analysis.

The pH treatments were randomly selected for behavioral analysis, alternating the elevated and low temperatures. Before filtration, each bucket's temperature and pH were measured to ensure that the treatment conditions had been maintained. Larvae were concentrated in a 40- μ m sieve and collected with a pipette. A total of 10–20 larvae per sample were distributed into 24-well plates. The wells (16 mm in diameter) were filled with 500 μ L of natural seawater with the same temperature and pH as in their original treatment (n = 3 or 4 replicates per tank). All larvae were allowed to habituate for 5 min prior to starting the test. Videos were recorded in the DanioVision Observation Chamber[®] (Noldus Information Technology, the Netherlands) using a camera (Basler, GigE) fitted with a 55-mm lens that was positioned 15 cm away from the microplate. Each video was recorded for 30 seconds at 30 frames per second and 1920 \times 1080 resolution. All processed

videos were analyzed under completely treatment-blind conditions using the software EthoVision XT 13.0 (Noldus Information Technology, Wageningen, the Netherlands). This software tracked active larval swimming for a maximum of 16 larvae simultaneously (not exceeded in our case) within a selected area. To reduce uncertainty, we pre-set a few criteria that were followed for each video. For example, if the larvae hit the walls of the well or if they collided, the behavioral variables after the collision were excluded from further analysis. Therefore, only larvae that were detected for at least 10 consecutive seconds without interruption were used in our analyses. If a larva did not move for 30 consecutive seconds, then no signal was detected by the program and a value of zero was indicated for the different behavioral variables of that larva.

The 2D_{distance} moved was measured as the distance (cm) that each larva traveled from the center point of the well over 30 sec. The mean velocity ($\text{cm} \cdot \text{ms}^{-1}$) was calculated by dividing the distance moved by the unit of time (ms). The mean meander ($\text{deg} \cdot \text{cm}^{-1}$) was calculated as the mean change in a larva's direction of movement relative to its distance moved; this provided an indication of how convoluted the larva's trajectory was.

Larval metabolism

At 72 hpf, larval respiration rates were determined using closed incubations in acrylic respirometry chambers (Engineering and Design Plastics Ltd, Cambridge, UK). We carried out six incubations for each experimental tank: three incubations with larvae in 120-mL respirometry chambers, and three incubations with larvae-excluded seawater (control) in 80-mL respirometry chambers. We applied a correction to compensate for the differences in chamber volume. About 100 L of live larvae were siphoned from the experimental tanks and resuspended in 1 L of filtered seawater from

the experimental tanks to reach a density of 100 larvae·mL⁻¹. The incubation chambers that contained larvae were filled with this same larval resuspension, whereas the control chambers were filled with filtered seawater from the experimental tanks. The chamber temperatures were maintained at the larval culture temperature during the incubations, either at room temperature (17°C) or at an high temperature (19°C) using a circulating water bath. The respiration rates were calculated from the measured differences in oxygen concentration during each incubation using a noninvasive fiber-optical system (FIBOX 3, PreSens, Regensburg, Germany) made up of an optical fiber and a reactive oxygen spot attached to the inner wall of each chamber. The reactive oxygen spots were calibrated using a 0% oxygen buffer that was prepared by dissolving 10 g of Na₂SO₃ into 1 L of seawater, and a 100% oxygen buffer that was prepared by bubbling air into 1 L of seawater for 20 min to achieve oxygen saturation. Oxygen consumption was measured over 1.5 to 2 hours and checked for linearity. Respiration rates were corrected for oxygen consumption in the controls and normalized to the number of larvae ($\mu\text{mol O}_2\cdot\text{larva}^{-1}\cdot\text{h}^{-1}$).

After each incubation (n = 9 per treatment), the exact number of larvae per chamber was determined from six 100- μL sample replicates per chamber that were distributed into a six-well plate.

For two of the three replicate incubations per experimental tank (n = 6 per treatment), the larvae were removed from the chambers, rinsed with distilled water to remove salt, and filtered through a pre-weighed Whatman GF/F 0.7- μm membrane. Filters containing larvae were dried in an oven at 60°C for 48 h to determine their DW. The relative CaCO₃ content (%) was calculated from the ash weight after burning at 550°C for 5 h and the DW of each sample ($\text{g CaCO}_3\cdot\text{g}^{-1}\text{ DW larvae}$).

Larval settlement

Larval settlement, which involves attachment to the substrate to achieve metamorphosis, occurs about 4 days after fertilization in *H. tuberculata* at $17 \pm 0.5^{\circ}\text{C}$ (Jardillier et al., 2008). Therefore, the larval settlement rate (%) was evaluated using premetamorphic 96 hpf veliger larvae. Twenty-four glass aquaria (20 cm W \times 35 cm L \times 20 cm H) were filled with 12 L of seawater at the same temperature and pH as in the original tanks. Temperatures were maintained by using a water bath set at 19°C for half of the aquaria. The other 12 aquaria were left at the low temperature of 17°C . We used a closed system with no water renewal to prevent the larvae from escaping from the small aquaria. Therefore, the pH of the acidified treatment could not be adjusted during the 24-h settlement period. As a result, the pH_T increased from 7.69 ± 0.04 at the beginning of the test to 7.86 ± 0.03 at the end of the test. In each aquarium, we installed two 10 cm \times 15 cm vertical polycarbonate plates, one covered with the green microalga *Ulvelia lens*, and one without *U. lens*. *Ulvelia lens* is one of the major inductive cues for the settlement of abalone larvae (De Viçose et al., 2010, 2012; Daume et al., 2004), and the absence of such cues can substantially reduce larval settlement rates (Searcy-Bernal et al., 1992; Slattery, 1992; Daume et al., 1999). After estimating the density of the larvae, a 1-mL volume was distributed using a pipette into two aquarium pseudo-replicates ($n = 6$ per treatment). The volume was adjusted to create a similar density of the larvae in each aquarium. After about 24 h, we used a binocular microscope to record the number of larvae that had settled on each plate. The water in each tank was filtered using a 40- μm filter to collect any larvae that were unsettled or that had attached to the sides and bottom of the tank. We applied low-pressure seawater to remove any larvae that had attached to the tank. The larvae that were collected from each tank were preserved in 70% ethanol for later counting. Finally, we calculated the percentage of settled larvae per plate type (blank and *Ulvelia*-covered) per treatment using the average of the two pseudo-replicates per tank (formula 3).

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$$(3) = \frac{\text{number of settled larvae}}{\text{total larvae}} \times 100$$

401 One tank was lost for the pH-elevated and 17°C treatment, resulting in 2 instead of 3 replicates for
402 that treatment.

403

404 *Statistical analyses*

405 All statistical analyses were performed in R (version 4.0.5) and RStudio software (RStudio,
406 Boston, United States). Differences in hatching success, percentage of swimming larvae, shell
407 length, behavioral parameters, larval respiration rate, and larval settlement on *Ulvelia*-covered
408 plates were assessed using a two-way analysis of variance (ANOVA) with pH, temperature, and
409 the interaction between pH and temperature as fixed factors. For all of the aforementioned
410 parameters, for each sampling time, the data were averaged per tank (n = 12 in total). Subsequently,
411 our replicates are tanks (not individual larvae). The normality of the residuals was verified with a
412 Shapiro–Wilk’s test, and the homogeneity of variances was tested using Levene’s test. To study
413 the effects of pH and temperature on the settlement rate on blank plates, we applied a Wilcoxon
414 rank sum test with continuity correction due to the presence of many zero values, and because the
415 normality of the residuals could not be verified. To study the interaction between pH and
416 temperature, we used a four-treatment-level Kruskal-Wallis rank sum test followed by a post hoc
417 Wilcoxon rank sum test with continuity correction. The p and F values of two-way ANOVA are
418 presented in the results, unless otherwise indicated.

419 To assess the frequency of the morphological and developmental parameters, we performed a four-
420 treatment-level (low pH - low temperature; low pH - high temperature; high pH - low temperature;

high pH - high temperature) chi-squared (χ^2) test to evaluate the effects of pH and temperature on larval phenotypes at 24 and 72 hpf.

Results

Carbonate chemistry and physicochemical characteristics of seawater

The mean seawater temperature, salinity, and carbonate chemistry parameters are presented in Table 2. The salinity was 35.18 ± 0.04 psu in all experimental tanks, and remained stable over the course of the experiment. A_T measured in the experimental tanks was $2364 \pm 17 \mu\text{Eq}\cdot\text{kg}^{-1}$ and remained stable over the course of the experiment and between experimental aquaria. The mean pH_T values were 7.95 ± 0.04 and 7.68 ± 0.12 for the high and low-pH treatments, respectively. The average temperatures were $17.40 \pm 0.55^\circ\text{C}$ and $19.35 \pm 0.53^\circ\text{C}$ for the low and high temperature treatments, respectively (means \pm SD).

Table 2 Mean parameters of seawater carbonate chemistry during the experiment. Total-scale seawater pH (pH_T), temperature, salinity, and total alkalinity (A_T) were used to calculate the partial pressure of CO_2 (pCO_2 ; μatm), dissolved inorganic carbon (DIC ; $\mu\text{mol}\cdot\text{kg}^{-1}$ SW), HCO_3^- , CO_3^{2-} , aragonite saturation state (Ω_{ar}), and calcite saturation state (Ω_{calc}) using CO2SYS software. The pH_T and temperature values shown represent the average value for each treatment, measured daily over the 5 days of the experiment ($n = 15$ per treatment). Salinity was measured once a day over the 5 days of the experiment ($n = 5$ per treatment). Results are expressed as mean \pm SD.

Treatment name	Treatment (pH-T°C)	Temperature (°C)	pH _T	pCO ₂ (µatm)	DIC	HCO ₃ ⁻ (µmol·kg ⁻¹)	CO ₃ ²⁻ (µmol·kg ⁻¹)	Ω _{ar}	Ω _{calc}
High pH – low temperature	8.0–17°C	17.5 ± 0.6	7.99 ± 0.04	478 ± 49	2122 ± 16	1950 ± 24	156 ± 10	2.40 ± 0.15	3.72 ± 0.23
Low pH – high temperature	7.7–19°C	19.4 ± 0.6	7.66 ± 0.12	1161 ± 319	2245 ± 45	2118 ± 58	88 ± 24	1.36 ± 0.37	2.10 ± 0.56
High pH – high temperature	8.0–19°C	19.3 ± 0.5	7.91 ± 0.05	593 ± 74	2147 ± 23	1985 ± 35	142 ± 14	2.20 ± 0.22	3.39 ± 0.34
Low pH – low temperature	7.7–17°C	17.3 ± 0.5	7.71 ± 0.11	999 ± 263	2236 ± 39	2109 ± 51	91 ± 21	1.40 ± 0.32	2.173 ± 0.49

Hatching success

The larval hatching success did not differ by temperature, pH, or the interaction of the two ($p > 0.05$; Table S1).

Percentage of swimming larvae

At 24 hpf, 72 hpf and 96 hpf the percentage of swimming larvaewas significantly lower at 19°C than at 17°C (24h hpf : $F_{1,8} = 19.92$, $p = 0.002$, Fig. 4A; 72 hpf : $F_{1,8} = 5.52$, $p = 0.046$, Fig. 4C;

96 hpf : $F_{1,8} = 13.47$, $p = 0.006$, Fig. 4E). However, no pH effect (24h hpf : $F_{1,8} = 0.41$, $p = 0.539$, Fig. 4A; 72 hpf : $F_{1,8} = 3.11$, $p = 0.115$, Fig. 4C; 96 hpf : $F_{1,8} = 0.04$, $p = 0.845$, Fig. 4E) and no interaction between temperature and pH (24h hpf : $F_{1,8} = 0.09$, $p = 0.764$, Fig. 4B; 72 hpf : $F_{1,8} = 0.29$, $p = 0.602$; Fig. 4D; 96 hpf : $F_{1,8} = 0.3$, $p = 0.618$; Fig. 4F) were found.

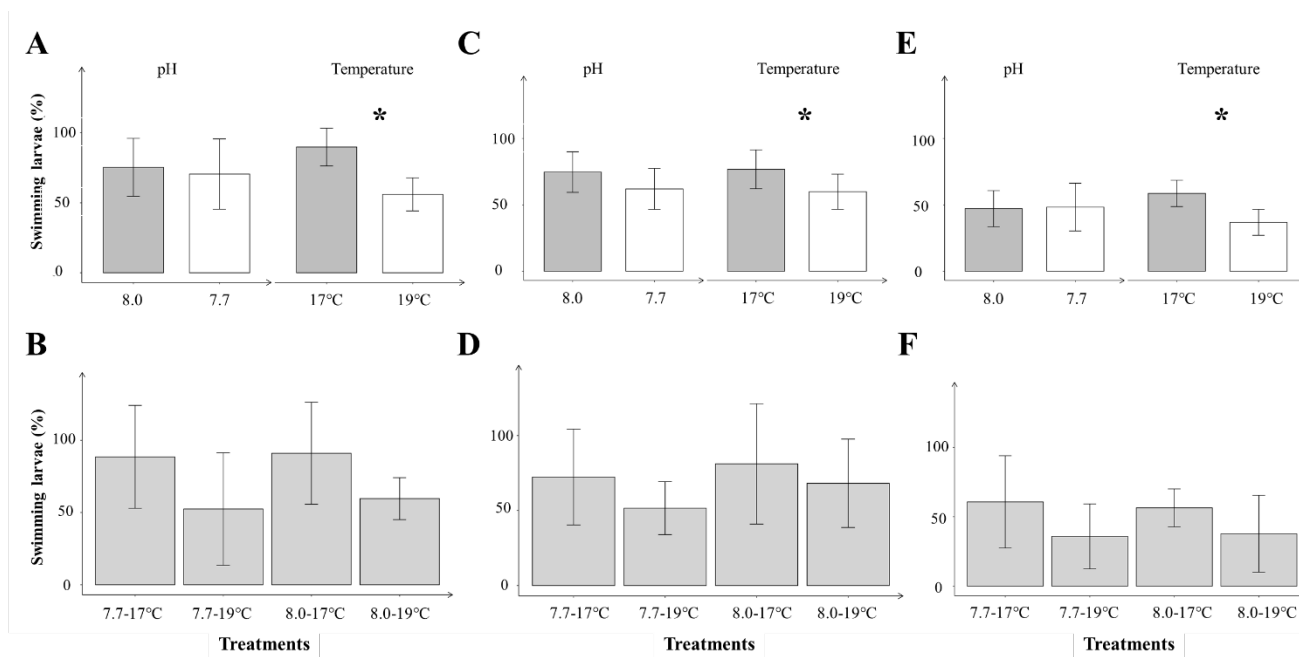


Fig. 4 Effects of pH and temperature on *Haliotis tuberculata* percentage of swimming larvae at **A)** 24 hours post-fertilization (hpf), **C)** 72 hpf, and **E)** 96 hpf ($n = 6$ tanks per treatment). Combined treatment effects of pH and temperature on *H. tuberculata* percent swimming larvae at **B)** 24 hpf, **D)** 72 hpf, and **F)** 96 hpf ($n = 3$ tanks per treatment). bar values represent means and error bars represent the standard deviation of the mean. Asterisks denote significant differences between temperature treatments by two-way ANOVA.

466 *Morphometric characteristics*

467 24 hpf larvae: Shell formation was significantly affected by temperature ($X^2 (2) = 54.59$, $p <$
468 0.001), pH ($X^2 (2) = 94.82$, $p < 0.001$), and the four treatment levels (low pH - low temperature,
469 low pH - high temperature, high pH - low temperature, high pH - high temperature); $X^2 (6) =$
470 130.71 , $p < 0.001$) (Fig. 5A; Table S2). Larvae exposed to pH 7.7 were 4.5 times less likely to
471 have normal shell formation compared with larvae exposed to pH 8.0 (9.9% at pH 7.7 versus 45%
472 at pH 8.0; Table S2). Larvae exposed to 19°C temperatures were 3 times as likely to have normal
473 shell formation compared with larvae exposed to 17°C temperatures (45% at 19°C versus 15% at
474 17°C; Table S2). The larvae exposed to the low pH and low temperature treatment had the lowest
475 proportion of normal shell formation, whereas larvae exposed to the high pH and high temperature
476 treatment had the highest proportion of normal shell formation (4% vs 59%, respectively) (Fig.
477 5A; Table S2).

478 Similarly, tissue organogenesis at 24 hpf was significantly affected by temperature ($X^2 (2) = 24.75$,
479 $p < 0.001$), pH ($X^2 (2) = 7.83$, $p = 0.020$), and the four treatment levels (low pH - low temperature,
480 low pH - high temperature, high pH - low temperature, high pH - high temperature; ($X^2 (6) =$
481 45.46 , $p < 0.001$) (Fig. 5B; Table S2). Larvae exposed to the high temperature had a lower
482 proportion of normally developed tissues than did larvae exposed to the low temperature (39% at
483 19°C versus 61% at 17°C; Table S2), whereas larvae exposed to both low-pH and high pH
484 conditions had similar tissue organogenesis (51% at pH 7.7 versus 54% pH 8.0).

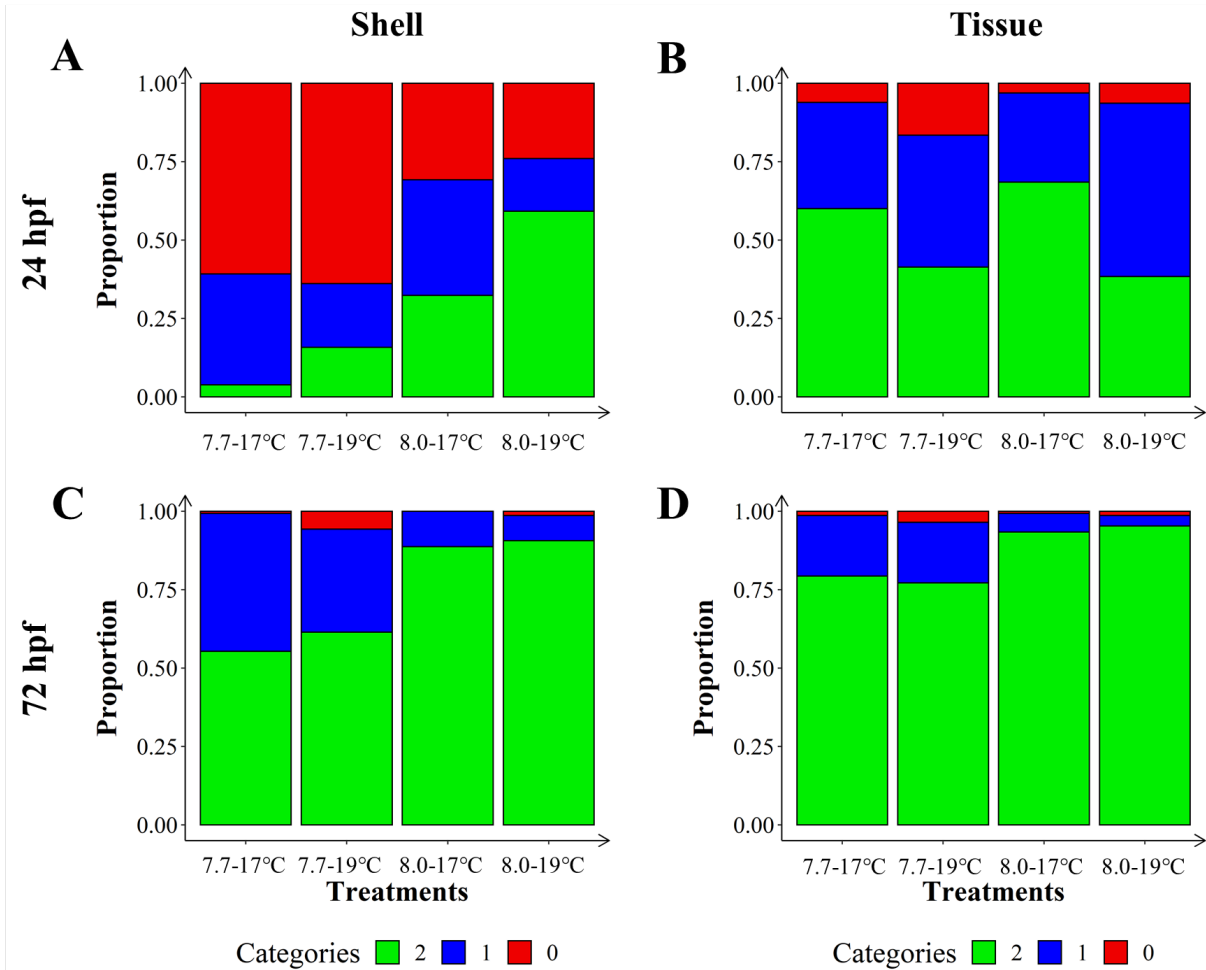


Fig. 5 Proportional distribution of larvae at 24 hours post-fertilization (hpf) (**A** and **B**) and at 72 hpf (**C** and **D**) according to their morphological categories. Colored bars indicate the degree of malformation for each variable. A and C: shell formation score (2 = normal shell; 1 = partially and/or abnormally developed shell; 0 = no shell). B and D: tissue organogenesis score (2 = normally developed tissues; 1 = partially or abnormally developed tissues; 0 = undeveloped tissues). The frequencies of each category are shown (n = 518 larvae in total).

72h larvae: Shell formation was significantly affected by temperature ($X^2(2) = 10.61$, $p = 0.005$), pH ($X^2(2) = 71.19$, $p < 0.001$), and the four treatment levels (low pH - low temperature, low pH -

high temperature, high pH - low temperature, high pH - high temperature; $X^2 (6) = 84.76$, $p < 0.001$) (Fig. 5C; Table S2). Larvae exposed to pH 7.7 had a significantly lower proportion of specimens with normal shell development compared with larvae exposed to the high pH of 7.9 (60% at pH 7.7 versus 90% at pH 8.0) (Fig. 5C; Table S2). The larvae in the high pH and high temperature treatment showed the highest proportion of specimens with normal shell development (92%), followed by those in the high pH and low temperature treatment (88%). Larvae exposed to the low-pH and low temperature treatment had the lowest proportion of specimens with normal shell development (56%; Fig. 5C; Table S2).

Tissue organogenesis at 72 hpf was significantly affected by pH ($X^2 (2) = 36.15$, $p < 0.001$) and by the four treatment levels (low pH - low temperature, low pH - high temperature, high pH - low temperature, high pH - high temperature; $X^2 (6) = 39.54$, $p < 0.001$) (Fig. 5D; Table S1), but was not significantly affected by temperature ($X^2 (2) = 2.70$, $p = 0.26$). Larvae exposed to the low-pH had a significantly lower proportion of specimens with normally developed tissues (78%) compared with larvae exposed to the high pH (95%; Table S2).

Shell length

The shell length of the 24-hpf larvae could not be measured in the low-pH treatments because there were very few larvae with normally developed shells (Fig. 5A). The shell length of the 72-hpf larvae was significantly shorter in the low-pH treatments compared with the high pH treatments ($F_{1,8} = 11.90$, $p = 0.008$; pH 8.0_{mean} = $265.01 \pm 6.69 \mu\text{m}$; pH 7.7_{mean} = $258.57 \pm 10.61 \mu\text{m}$). However, the temperature had no effect on larval shell length at 72 hpf ($F_{1,8} = 4.66$, $p = 0.063$; 17°C_{mean} = $264.83 \pm 8.49 \mu\text{m}$; 19°C_{mean} = $260.28 \pm 8.89 \mu\text{m}$). No interaction between temperature and pH was

517 observed ($F_{1,8} = 1.60$, $p = 0.24$; $8.0\text{--}17^{\circ}\text{C}_{\text{mean}} = 265.92 \pm 6.92 \mu\text{m}$; $8.0\text{--}19^{\circ}\text{C}_{\text{mean}} = 264.09 \pm 6.34$
518 μm ; $7.7\text{--}17^{\circ}\text{C}_{\text{mean}} = 263.00 \pm 10.38 \mu\text{m}$; $7.7\text{--}19^{\circ}\text{C}_{\text{mean}} = 254.44 \pm 9.09 \mu\text{m}$).

519

520 *Shell calcification*

521 The mean birefringence (number of grayscale pixels) at 24 hpf was significantly lower at pH 7.7
522 than at the high pH of 7.9 ($F_{1,8} = 8.55$, $p = 0.019$; Fig. 6A). However, birefringence (and
523 correspondingly, shell calcification) did not differ significantly by temperature ($F_{1,8} = 1.33$, $p =$
524 0.281 ; Fig. 6A) or by the interaction between temperature and pH ($F_{1,8} = 0.01$, $p = 0.920$; Fig. 6B).
525 The mean birefringence at 72 hpf was significantly lower in the pH 7.7 compared with pH 8.0 ($F_{1,8}$
526 $= 12.21$, $p = 0.008$) and 19°C compared with 17°C ($F_{1,8} = 17.83$, $p = 0.003$) treatments, indicating
527 reduced shell calcification in low pH treatment and high temperature treatment compared with the
528 ambient pH and temperature respectively (Fig. 6C). However, the interaction between temperature
529 and pH did not significantly affect shell birefringence at 72 hpf ($F_{1,8} = 0.08$, $p = 0.784$; Fig. 6D).

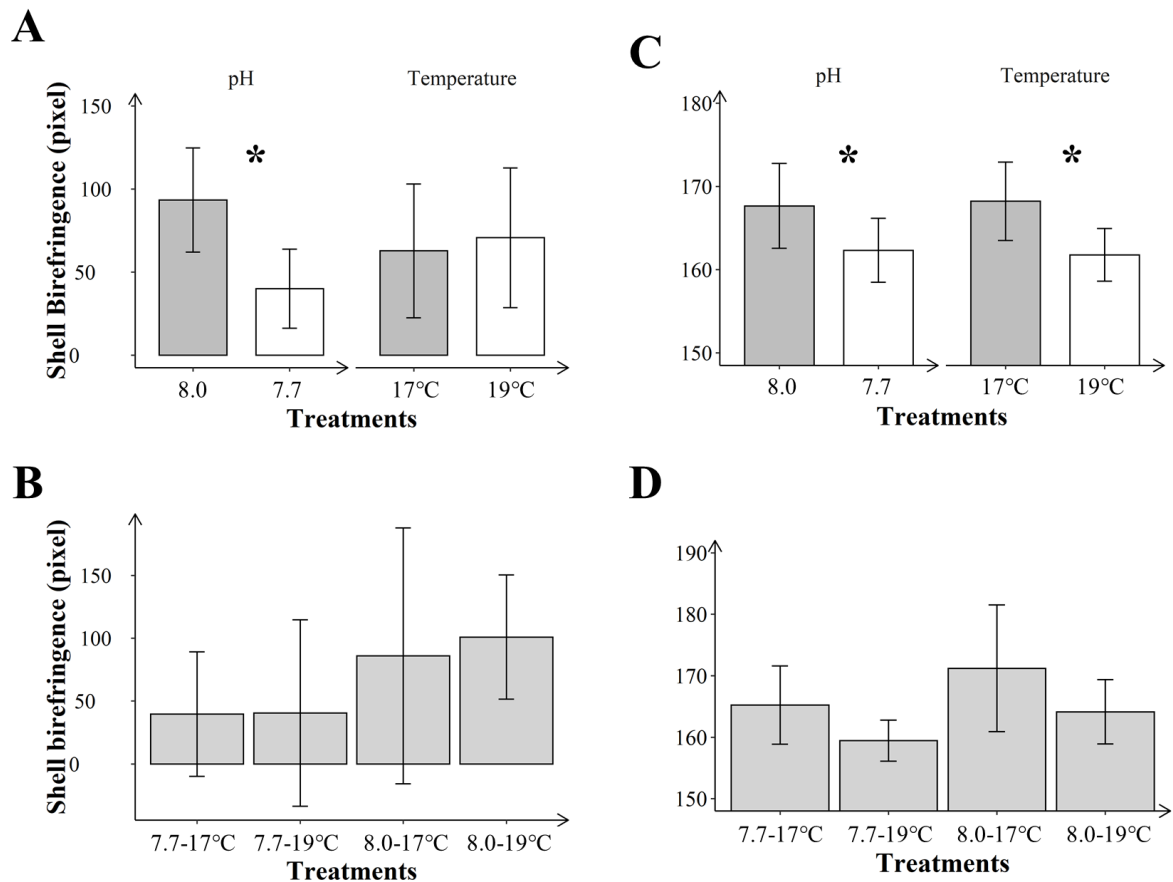


Fig. 6 Individual effects of pH and temperature on mean shell birefringence of larvae at **A)** 24 hpf and **C)** 72 hpf (n = 6 tanks per treatment). Combined effects of pH and temperature on mean shell birefringence of larvae at **B)** 24 hpf and **D)** 72 hpf (n = 3 tanks per treatment). bar values represent means and error bars represent the standard deviation of the mean. Asterisks indicate significant differences between either temperature or pH treatments, as determined by two-way ANOVA.

540 *Scanning electron microscopy*

541 In the high pH treatments (7.9), the shell surface of larvae observed at 24 hpf showed a normal
542 granular texture with an alveolar pattern (Fig. 7A, B, E, F). However, the larvae reared in the low-
543 pH treatments (7.6) showed signs of shell deformation and microfractures (Fig. 7C, D). Details of
544 the shell surface boxed in Figures 7C and 7D revealed irregularities and mineralization defects that
545 were probably due to shell dissolution. pH 8.0The shell surface of larvae observed at 72 hpf were
546 of similar shape and pattern in both the low (7.7) and high (8.0) pH treatments (Fig. 8A–D). At
547 higher magnification, the shell surfaces of high pH larvae revealed a homogeneous granular texture
548 and alveolar pattern (Fig. 8E, F). In the low-pH treatments, the details of the shell surface boxed
549 in 8C and 8D revealed surface heterogeneities and small holes within the alveolar network
550 suggesting mineral dissolution. (Fig. 8G, H)

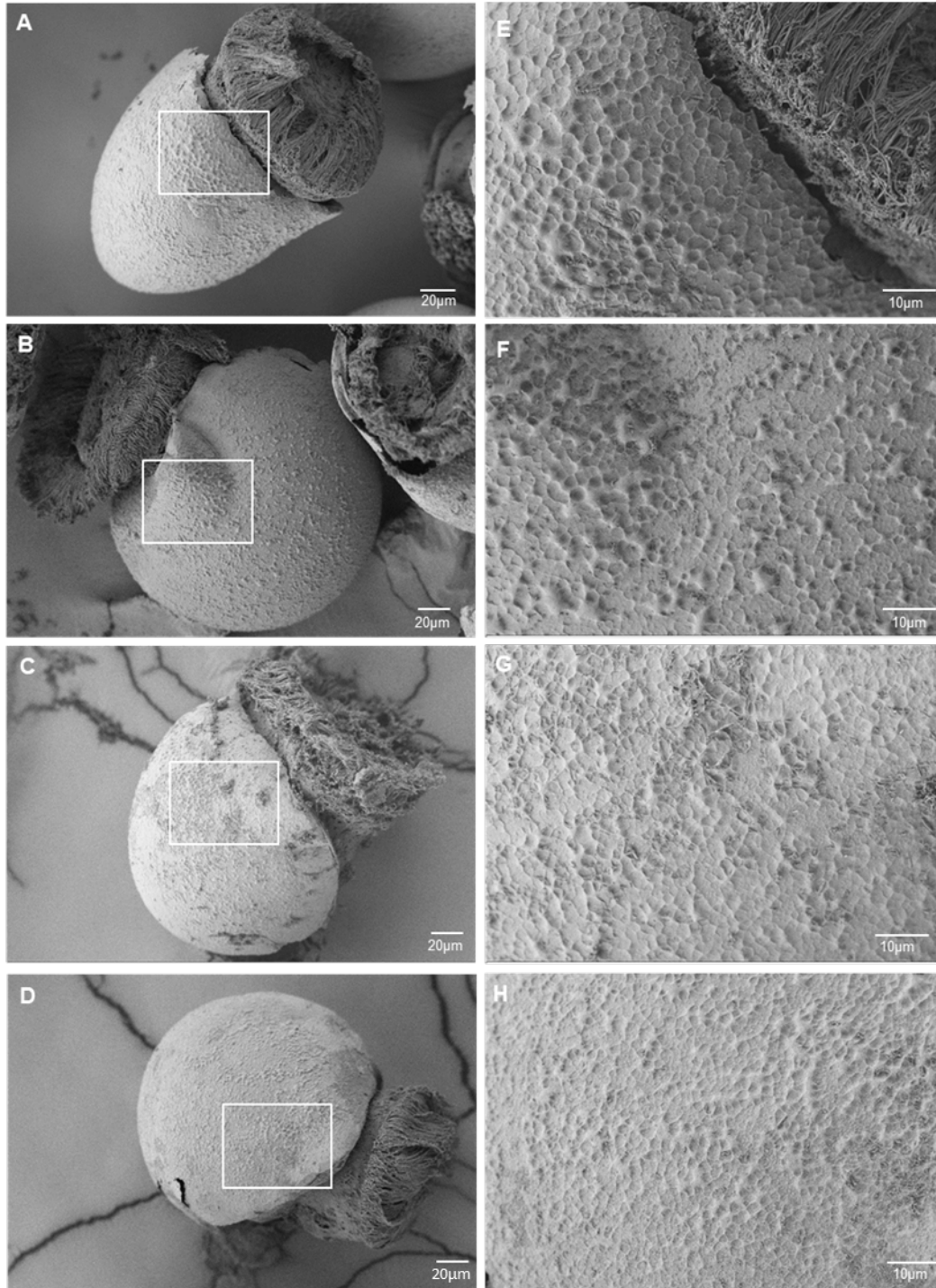
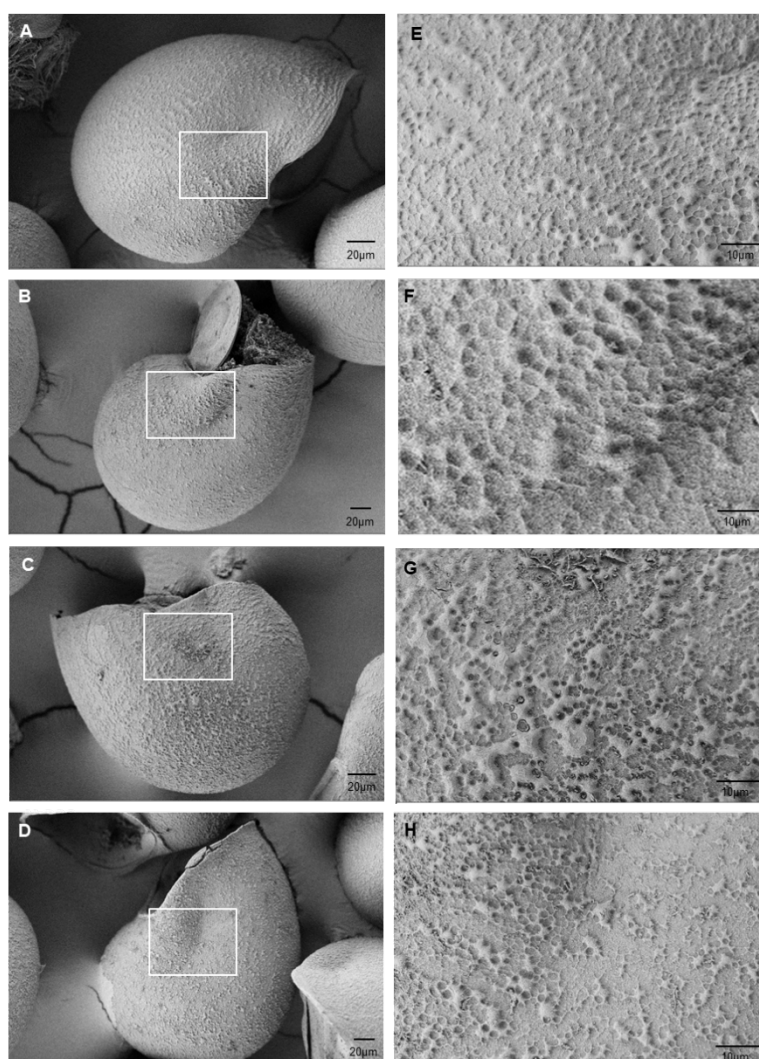


Fig. 7 Scanning electron microscopy (SEM) images of *Haliotis tuberculata* larvae at 24 hpf in the four experimental treatments **A)** pH 8.0–17°C; **B)** pH 8.0–19°C; **C)** pH 7.7–17°C; and **D)** pH 7.7–

554 19°C. Lateral views of the larvae show the protoconch shell and the velum. **E–H** show magnified
555 views of the corresponding white squares in A–D.

556

557 The shell surface of larvae observed at 72 hpf were of similar shape and pattern in both the low
558 (7.7) and high (8.0) pH treatments (Fig. 8A–D). The magnified areas of the shells in all treatments
559 revealed a typical granular texture and alveolar pattern (Fig. 8E–H). However, the surface
560 heterogeneities and small holes within the alveolar network in the low-pH treatments (Fig. 8G, H)
561 suggest mineral dissolution.



562

Fig. 8 Scanning electron microscopy (SEM) images of *Haliotis tuberculata* larvae at 72 hpf in the four experimental treatments **A)** pH 8.0–17°C; **B)** pH 8.0–19°C; **C)** pH 7.7–17°C; and **D)** pH 7.7–19°C. Lateral views of the larvae show the well-developed protoconch covering the larval body. In **A**, the protoconch is well developed and covers the larval body. In **B**, the operculum closes the shell aperture after complete retraction of the veliger. **E–H** show magnified views of the corresponding white squares in A–D.

Larval behavior

None of the larval behavioral parameters (Distance moved (total cm⁻¹), mean velocity (cm·ms⁻¹), mean meander (deg·cm⁻¹) differed significantly due to pH, temperature, or the interaction between pH and temperature (Table S3).

Respiration rate

The respiration rate of larvae did not differ significantly between pH ($F_{1,8} = 1.54$, $p = 0.250$; two-way ANOVA), or temperature ($F_{1,8} = 2.57$, $p = 0.15$; two-way ANOVA). No interaction was found between pH and temperature for larval respiration rate ($F_{1,8} = 5.66$, $p = 0.045$).

584 *Larval settlement*

585 The larval settlement on *Ulvelia*-covered plates was significantly higher in the high temperature
586 (19°C) treatments than in the low temperature (17°C) treatments ($F_{1,7} = 14.02$, $p = 0.007$; Fig. 9A).
587 No significant differences were observed between pH treatments ($F_{1,7} = 1.40$, $p = 0.275$; Fig. 9A)
588 or from the interaction between pH and temperature ($F_{1,7} = 0.47$, $p = 0.52$; Fig. 9C).

589 On blank plates, we found significant effects from temperature ($W = 0$, $p = 0.005$; Wilcoxon rank
590 sum test with continuity correction; Fig. 9C) and from the four treatment comparisons ($H(3) =$
591 8.70 , $p = 0.033$; Kruskal–Wallis) (Fig. 9D). Larval settlement on the blank plates was significantly
592 higher in the high temperature (19°C) treatments than in the low temperature (17°C) treatments
593 (Fig. 9B). No significant differences were found between the low-pH and high pH treatments (W
594 $= 14$, $p = 0.923$; Wilcoxon rank sum test with continuity correction; Fig. 9C).

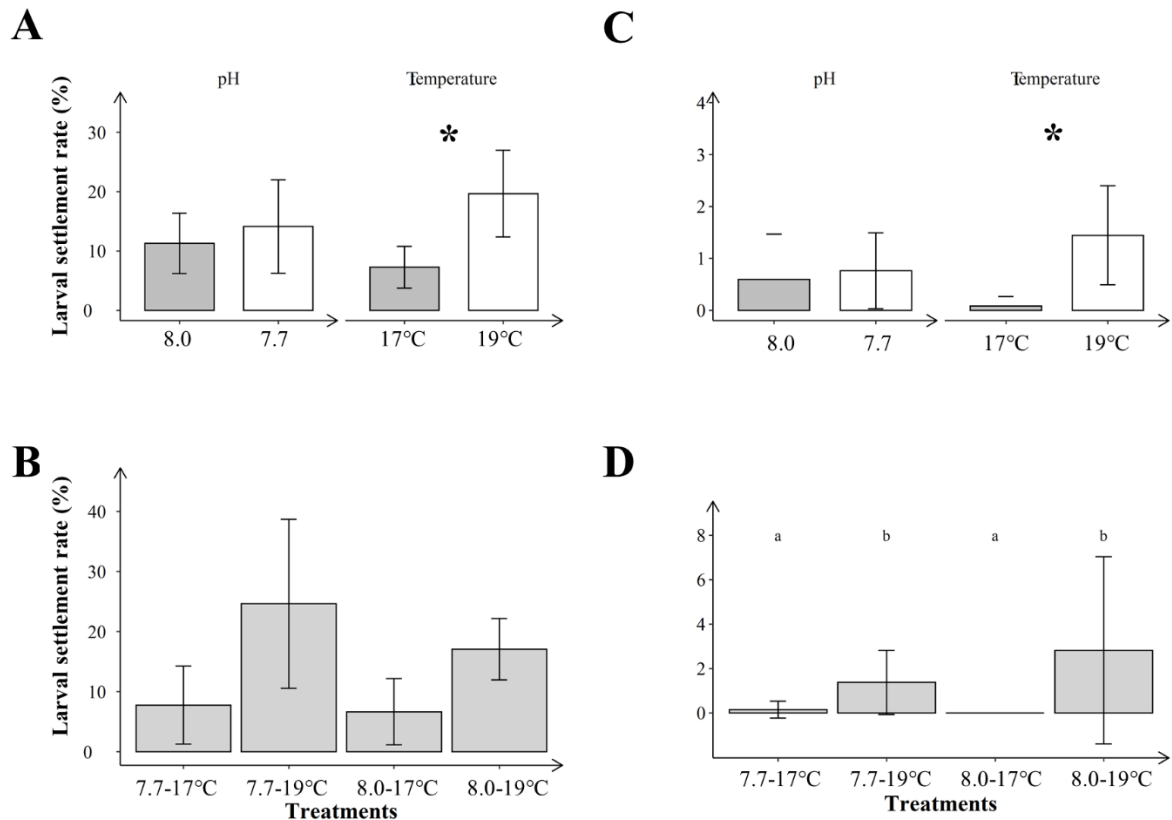


Fig. 9 Effects of pH and temperature on larval settlement on **A)** plates covered with the green microalga *Ulvella lens* (n = 6 tanks per treatment; two-way ANOVA) and **B)** blank plates (n = 6 tanks per treatment; Wilcoxon rank sum test with continuity correction). Combined effects of pH and temperature on larval settlement on **C)** *Ulvella*-covered plates and **D)** blank plates at 96 hours post-fertilization (hpf) (n = 3 tanks per treatment). bar values represent means and error bars represent the standard deviation of the mean. Different lowercase letters indicate significant differences among treatments. Asterisks indicate significant differences between either temperature or pH treatments.

Discussion

In this study, we exposed *H. tuberculata* larvae to the individual and combined effects of OA and OW. We found that a 0.3-unit decrease in pH induced decreases in or no effects on larval responses, whereas a 2°C increase in temperature caused broader effects, resulting in either increased/enhanced, decreases/diminished larval responses or had no impacts. However, no significant interactive effects of temperature and pH were observed for the variables measured. Despite that, some treatment effects were recorded for the parameters that were not analyzed using ANOVA (Table 3). Among the biological parameters investigated in this study, shell formation, length, and calcification appeared to be the most sensitive to the effects of temperature and pH. The results of this study lend further support to the results from recent meta-analyses suggesting that the interactive effects of OA and OW on calcifying species, including mollusks, are not as ubiquitous as OA's and OW's individual effects (Kroeker et al., 2013; Przeslawski et al., 2015).

The hatching success of trochophore larvae of *H. tuberculata* was not significantly affected by temperature, pH, or the interaction of the two. These results are in line with those of Pedroso (2017), who found no impact on the hatching success of *H. asinina* in response to raising the ambient temperature of 29°C by 2°C. The lack of significant effects by pH or by the interaction of pH and temperature was likewise similar to previous findings. Guo et al. (2015) found no effect of pH_{NBS} 7.94 on the hatching success of *H. diversicolor* and *H. discus*, whereas both species experienced reduced hatching success when the pH value was decreased to 7.71 or lower, compared with the high pH of 8.15. In two separate studies on the hatching success of *H. discus hannai*, Kimura et al. (2011) and Li et al. (2013) found no impact of exposure to pH_{NBS} 7.6 and higher, compared with pH of 8.02. Likewise, Tahil and Dy (2016) found that pH_{NBS} 7.78 did not affect the *H. asinina* hatching success compared to the pH of 7.97; however, pH treatments of 7.60

and 7.40 negatively affected the hatching success. Our results show that *H. tuberculata* larvae can hatch at the lower pH values and high temperatures expected in the coming decades. Such tolerance to OA and OW in the early larval stages has already been reported for other marine species (Przeslawski et al., 2015), and might originate from protective factors in the egg (Hamdoun and Epel, 2007).

An increased number of larvae with tissue abnormalities was observed under high temperature treatments. This was accompanied by significant declines in the percentage of swimming larvae in the high temperature treatments compared to the low temperature treatments at 24, 72 and 96 hpf. Low pH, also, reduced the normal tissue development of the larval stages. A study on *H. asinina* found < 1% of normal trochophores at the reduced pH of 7.60, compared with the ambient pH of 7.97 (Tahil and Dy, 2016). Parker et al. (2011) also reported slower larval development in the Sydney rock oyster, *Saccostrea glomerata*, when the ambient pH (pH_{NBS} 8.2) was reduced by 0.3 pH units. A study by Wessel et al. (2018) on *H. tuberculata* showed a significant increase in the number of 30-hpf larvae with abnormal and delayed development at pH_T 8.0 compared with larvae at pH_T 7.7 and 7.6. These findings suggest that a difference of only a few hours of exposure to stressors can influence their effects during larval development.

Larval shell formation was also impaired under low pH, similar to previous studies on *H. coccoradiata* (pH_{NBS} = 7.8 and 7.6 compared with the ambient pH of 8.2; Byrne et al., 2011); *H. kamtschatkana* (reported as pCO₂ values of 800 µatm and 1800 µatm compared with a control of 400 µatm; Crim et al., 2011); *H. asinina* (pH_{NBS} = 7.85 and 7.65 compared with the ambient pH of 8.15; Santander-De Leon et al., 2018); and *H. tuberculata* (pH_T = 7.68 and 7.58 compared with the ambient pH of 8.00; Wessel et al., 2018). Shell calcification of *H. tuberculata* larvae was also reduced under low pH, as shown by the decrease in birefringence intensity measured at 24 and 72

hpf. Additional SEM observations revealed differences in larval shell texture and the presence of numerous small holes on the outer shell surface, suggesting that CaCO_3 dissolution may be responsible for the reduced calcification seen in larvae raised at a lower pH. The lack of calcification observed under cross-polarization microscopy and SEM is consistent with previous results from Wessel et al. (2018), and provides further evidence that abalone larvae deposit less CaCO_3 and produce more fragile and thinner shells when exposed to OA. Our results confirm, as reported for numerous mollusk species, that the growing calcified shells of larval and juvenile abalones are highly sensitive to OA (Gazeau et al., 2013; Przeslawski et al., 2015). Naturally occurring *H. tuberculata* populations may be at greater risk from future pH reduction, as early shell alterations may impair larval development and recruitment.

In this study, the impaired process of shell formation induced by low pH was mitigated by high temperature, resulting in a higher proportion of 24- and 72-hpf larvae with normal shell development as shown in Fig. 5A and 5C. Observations of temperature reducing the negative impacts of OA on shell calcification and development have been previously reported for larval and juvenile mollusks (Davis et al., 2013; Ko et al., 2014).

Shell length of *H. tuberculata* larvae was significantly reduced by low-pH conditions, but no temperature or interaction effects were observed. Crim et al. (2011) reported a 5% reduction in larval shell size in the northern abalone *H. kamtschatkana* after an 8-day exposure to acidified conditions (reported as CO_2 levels 400 μatm vs 800 μatm), which is consistent with our results (4% reduction at 24 hpf and 6% reduction at 72 hpf). Similarly, the larvae of *H. asinina* ($\text{pH}_{\text{NBS}} = 7.85$ and 7.65 compared with the ambient pH of 8.15; Santander-De Leon et al., 2018); *H. diversicolor*; and *H. discus hannai* ($\text{pH}_{\text{NBS}} = 7.94$ and lower compared with the high pH of 8.15; Guo et al., 2015) experienced reductions in shell length at low-pH compared with high pH

conditions. In a study by Onitsuka et al. (2018), larvae of *H. discus hannai* at pH_{NBS} 7.79 did not show any significant reduction in shell length compared with larvae at the high pH_{NBS} of 7.99.

In our study, the lack of significant effects on shell size from temperature or from the interaction between temperature and pH is consistent with results obtained for other marine mollusks. For example, Thiagarajan and Ko (2012) reported no impact of high temperature (30°C compared with 24°C), or temperature–pH interaction on the shell size of the Portuguese oyster *Crassostrea angulata*. On the other hand, Talmage and Gobler (2011) recorded decreased larval size in two bivalves (*Mercenaria mercenaria* and *Argopecten irradians*) resulting from both a reduced pH_{NBS} of 7.8 (compared with the ambient pH of 8.08) and an high temperature of 28°C (compared with the ambient temperature of 24°C). These contrasting effects confirm a high variability in the responses of marine mollusk larvae to both isolated and combined stressors.

The absence of direct effects of pH and temperature on larval physiology and behavior does not rule out the existence of their indirect effects through cellular and molecular mechanisms that were not measured in the present study. Indeed, larval developmental stages are characterized by huge changes (in physiology and sensitivity) over a short period of time (5 days) and the measurements of physiological and behavioral variables do not account for the whole developmental processes that may be impacted by the stressors. Complementary studies at the molecular level (i.e., gene expression, protein and enzyme synthesis, etc.) will be needed to fully understand the fine developmental changes induced by the climate change drivers (Portner et al. 2010).

Larval settlement was greater at high temperatures than at the low temperature on both *Ulvela*-covered and blank plates. The shorter larval duration that corresponded with high temperature in

697 this study is consistent with previous studies of related *Haliotis* species, such as *H. sorenseni*, *H.*
698 *rufescens*, *H. corrugata*, and *H. fulgens* (Leighton, 1972, 1974). This suggests that high
699 temperatures resulting from global warming reduce the duration of the planktonic larval stage and
700 accelerate the settlement rate (Byrne et al., 2011). This may be an advantage (Byrne et al., 2011),
701 as a longer planktonic stage may reduce the chances of survival due to increased risk of predation
702 and exposure to other environmental stresses (Parker et al., 2013). We did not record any
703 significant effects of pH on *H. tuberculata* settlement, which is in line with the findings of another
704 study on *H. kamtschatkana* (Crim et al., 2011). However, the literature contains contradictory
705 results, with a tendency toward a lower settlement rate of the New Zealand abalone, *H. iris*, in
706 low-pH conditions (Espinel-Velasco et al., 2020) as well as for the donkey's ear abalone, *H.*
707 *asinina*, in three low-pH treatments (Tahil and Dy, 2015).

708 Our results showed that in every treatment, the settlement rate of *H. tuberculata* larvae on *Ulvela-*
709 covered plates was higher than the settlement rate on blank plates by a factor of at least 15. This
710 supports the notion that settlement cues increase marine invertebrate larval settlement (Hadfield
711 and Paul, 2001; Pawlik, 1992; Rodriguez et al., 1993). However, lowered pH may cause indirect
712 effects, due to the effect of low pH on the settlement substrate. For example, O'Leary et al. (2017)
713 reported that the settlement rate of *H. rufescens* under both present-day and low-pH treatments
714 increased from 11% in tanks with no crustose coralline algae, to ~70% in the presence of crustose
715 coralline algae. This raises another concern regarding the indirect impacts of OA and OW on
716 abalone larval settlement. Coralline algae are common settlement cues for abalone larvae (De
717 Viçose et al., 2010, 2012; Williams et al., 2008; Roberts et al., 2004; Roberts and Nicholson, 1997),
718 and they are decreasing due to OA and OW (Kuffner et al., 2008; Martin and Gattuso, 2009; Diaz-
719 Pulido et al., 2012; Hofmann et al., 2012). Therefore, even in the absence of the direct impacts of

720 OA and OW on abalone larvae, such environmental changes may indirectly reduce larval
721 settlement by reducing their settlement cues. On the other hand, if *Ulvela* species, like many non-
722 calcifying macroalgae, can benefit from high temperatures and reduced pH (Koch et al., 2013;
723 Hofmann et al., 2012), then abalone larvae may be able to maintain their normal settlement rates.
724 This presents an area requiring further examination.

725 None of the other behavioral responses (Distance moved (total cm^{-1}), mean velocity ($\text{cm} \cdot \text{ms}^{-1}$),
726 mean meander ($\text{deg} \cdot \text{cm}^{-1}$),) differed as an effect of pH, temperature, or the interaction between the
727 two. These outcomes parallel the responses of 2-day-old larvae of the Pacific oyster, *Crassostrea*
728 *gigas*, when exposed to high pH and low (-0.3 units) pH conditions (Valentini, 2019). Although
729 the method used in this research to study larval behavior is very common, it is noted that applying
730 such a two-dimensional method may not allow the abalone larvae to express natural swimming
731 behavior. Therefore, developing three dimensional methods are desired for future studies.

732 The lack of significant changes in larval behavior may partly justify the lack of changes observed
733 in larval respiration. Larval respiration of *H. tuberculata* did not differ due to the individual or
734 combined factors of pH and temperature. Although very low pH and highly high temperatures (i.e.,
735 out of the projected ranges for the year 2100) are known to amplify respiration (Campanati et al.,
736 2018; Waldbusser et al., 2015; Liu and He, 2012; Padilla-Gamiño et al., 2013), some studies have
737 confirmed that respiration was not impacted by a -0.3 unit pH reduction and/or a +3°C temperature
738 increase above the ambient conditions (Frieder et al., 2017). For example, the respiration rate of
739 the larvae of *Mytilus californianus* did not change when the ambient pH was reduced from 8.3 to
740 7.8 (Waldbusser et al., 2015). Similarly, Campanati et al. (2018) found no impact on the oxygen
741 consumption of *Reishia clavigera* larvae when the ambient pH was reduced from 8.1 to 7.6. High
742 pCO_2 in seawater likely increases the maintenance costs of acid–base homeostasis, the intracellular

ion balance required for protein folding and pH-sensitive physiological processes (Rivest and Hofmann, 2014; Lefevre, 2016). Therefore, our observation in this study of a lack of increase in *H. tuberculata* metabolic rates under acidified conditions may suggest that the larvae were able to use existing pools of ion pumps to conserve their acid–base homeostasis (Portner and Reipschlager, 1996; Guppy and Withers, 1999).

Table 3 Full results summary showing the effects of temperature, pH, and their interaction on the various parameters observed during this study. Arrow direction indicates whether these factors decreased/diminished (↓), increased/enhanced (↑), or had no significant effect (ns) on the measured parameters of *H. tuberculata* at 24, 72, and 96 hours post-fertilization (hpf). Empty cells indicate that the corresponding results were not applicable because they were not measured. Where nonparametric tests were used (i.e., for shell formation and tissue formation), treatment impacts are shown instead of interaction effects.

Parameter	Temperature effect			pH effect			Interaction effect		
	24 hpf	72 hpf	96 hpf	24 hpf	72 hpf	96 hpf	24 hpf	72 hpf	96 hpf
Hatching success	ns			ns			ns		
Percentage of swimming larvae	↓	↓	↓	ns	ns	ns	ns	ns	ns
Shell formation	↑	↓		↓	↓		8.0 - 17°C = ambient 7.7 - 19°C = ↓ 8.0 - 19°C = ↑	8.0- 17°C = ambient 7.7 - 19°C = ↓	

							7.7 - 17°C = ↓	8.0 - 19°C = ns 7.7 - 17°C = ↓	
Tissue formation	↓	ns		↓	↓		8.0 - 17°C = ambient 7.7 - 19°C = ↓ 8.0 - 19°C = ↓ 7.7 - 17°C = ns	8.0 - 17°C = ambient 7.7 - 19°C = ↓ 8.0 - 19°C = ns 7.7 - 17°C = ↓	
Shell length		ns			↓			ns	
Shell calcification	ns	↓		↓	↓		ns	ns	
Respiration rate		ns			ns			ns	
Larval swimming behaviors		ns			ns			ns	
Larval settlement			↑			ns			ns

756

757 In conclusion, the majority of the responses of *H. tuberculata* larvae were affected by low pH and
758 high temperature, but not by their interactions as summarized in Table 3. We did not detect any
759 significant interaction between the high temperature and low-pH stressors. However, such
760 interactions may exist, but we were not able to detect them using non-parametric tests. . For
761 example, although high temperature accelerated shell formation at 24 hpf, this was partially
762 mitigated by low pH. High temperature, however, turned into a threat by increasing the larval
763 tissue malformation at 24 hpf. High temperature decreased the percentage of swimming larvae at
764 all stages. It also declined shell calcification at 72 hpf despite the fact that it did not affect that of
765 larvae at 24 hpf. By the end of the larval stage at 96 hpf, high temperature accelerated the larval
766 settlement rate, which might reduce its detrimental effects on larval survival. Low pH, however,

did not have any impact on the larval settlement. Low pH impaired all shell formation related parameters in all stages. However, like high temperature, it did not affect processes such as hatching success, respiration rate and swimming behaviors. Even though the projected rates of pH decrease and temperature increase over the coming decades may not seem to be an obstacle for *H. tuberculata* larval production, it seems that under OA and OW, the majority of this species' responses are susceptible to changes in either temperature or pH. Thus, *H. tuberculata* larvae will have to cope with both stressors. In nature, this may happen through adaptation. In abalone farms, this can occur using various strategies. For example, first, by monitoring of the daily changes in seawater pH, it can be arranged to and tadjustment of avoid pumping low pH seawater into the farms whenduring the hours when the CO₂ concentrations in the seawater is at its peak ion (Murie and Bourdeau 2020). If farms have access to large-volume storage tanks, adding kelp species to the tanks may reduce CO₂ concentrations as kelp canopies have the ability to alter local seawater chemistry (Murie and Bourdeau 2020). Inclusion of an integrated multi-trophic aquaculture system based on algae might be helpful (Bolton et al. 2009) Addition of chemicals to increase seawater alkalinity and/or pH might be another solution for the hatcheries with a recirculating system. The selection of tolerant broodstocks to OA and GW can be also undertaken. However, due to the limitations of each of the aforementioned methods, the combinations of them may lead to more sustainable abalone productions.

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Table S1 Hatching success (%) under high temperature, low pH, and their interaction. Values are written as mean \pm standard deviation. Two-way ANOVA.

pH _{mean}		Temperature _{mean}		Treatment _{mean}				pH effect		Temperature effect		Interaction effect of pH x temperature	
8.0 (n = 6)	7.7 (n = 6)	17°C (n = 6)	19°C (n = 6)	8.0-17°C (n = 3)	7.7-19°C (n = 3)	8.0-19°C (n = 3)	7.7-17°C (n = 3)	F _{1,8}	p	F _{1,8}	p	F _{1,8}	p
82.50 \pm 2.70	81.13 \pm 4.71	82.70 \pm 3.55	80.93 \pm 4.02	81.71 \pm 3.32	78.57 \pm 4.29	83.29 \pm 2.31	83.68 \pm 4.19	0.43	0.529	0.72	0.421	2.57	0.147

Table S2 Number of larvae at 24 and 72 hours post-fertilization (hpf) representing each category of shell formation (Unshelled, partially shelled, and fully shelled) and soft tissue morphogenesis (Highly abnormal, abnormal, and normal) at two pH levels (7.7 and 8) and temperatures (17°C and 19°C), and within four experimental treatments (7.6–17°C; 7.6–19°C; 7.9–17°C; and 7.9–19°C).

24 hpf larvae						
Factor	Shell formation			Soft tissue morphogenesis		
pH	Unshelled	Partially shelled	Fully shelled	Highly abnormal	Abnormal	Normal
7.7	164	73	26	30	100	133
8.0	70	69	116	12	106	137
Temperature						
17	161	98	47	18	101	187

19	73	44	95	24	105	83
pH*temperature						
7.7-17°C	79	46	5	8	44	78
7.7-19°C	85	27	21	22	56	55
8.0-17°C	40	48	42	4	37	89
8.0-19°C	30	21	74	8	69	48
72 hpf larvae						
Factor	Shell			Tissue		
pH	<i>Unshelled</i>	<i>Partially shelled</i>	<i>Fully shelled</i>	<i>Highly abnormal</i>	<i>Abnormal</i>	<i>Normal</i>
7.7	9	104	168	7	55	219
8.0	2	27	265	3	12	279
Temperature						
17	1	76	215	3	38	251
19	10	55	218	7	29	247
pH*temperature						
7.7-17°C	1	59	82	2	29	111
7.7-19°C	8	45	86	5	26	108
8.0-17°C	0	17	133	1	9	140
8.0-19°C	2	10	132	2	3	139

1091

1092

1093 **Table S3** Behavioral parameters measured and analyzed using EthoVision XT 13.0 software
1094 (Noldus Information Technology, Wageningen, the Netherlands). Values are written as mean ±
1095 standard deviation. Two-way ANOVA.

	pH		Temperature		pH effect		Temperature effect		Interaction effect of pH × temperature	
	8.0 (n = 3)	7.7 (n = 3)	17°C (n = 6)	19°C (n = 6)	F _{1,8}	p	F _{1,8}	p	F _{1,8}	p

Distance moved (total cm ⁻¹)	0.53 ± 0.29	0.57 ± 0.47	0.65 ± 0.48	0.46 ± 0.25	0.02	0.890	1.98	0.196	2.37	0.162
Mean velocity (cm·ms ⁻¹)	0.00 ± 0.00	0.00 ± 0.00	0.00 ± 0.00	0.00 ± 0.00	0.01	0.922	2.29	0.167	2.82	0.131
Mean meander (deg·cm ⁻¹)	-38 ± 505	-161 ± 24	-229 ± 53	27 ± 91	0.68	0.432	3.95	0.082	0.82	0.391

1096