Resilience of the larval slipper limpet *Crepidula onyx* to direct and indirect-diet effects of ocean acidification

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Ocean acidification (OA) is known to directly impact larval physiology and development of many marine organisms. OA also affects the nutritional quality and palatability of algae, which are principal food sources for many types of planktonic larvae. This potential indirect effect of OA via trophic interactions, however, has not been fully explored. In this study, veligers of *Crepidula onyx* were exposed to different pH levels representing the ambient (as control) and low pH values (pH 7.7 and pH 7.3) for 14 days, and were fed with *Isochrysis galbana* cultured at these three respective pHs. pH, diet, nor their interactions had no effect on larval mortality. Decrease in pH alone had a significant effect on growth rate and shell size. Structural changes (increased porosity) in larval shells were also observed in the low pH treatments. Interactions between acidification and reduced diet quality promoted earlier settlement. Unlike other calcifying molluscs, this population of slipper limpets introduced to Hong Kong in 1960s appeared to be resilient to OA and decreased algal nutritional value. If this robustness observed in the laboratory applies to the field, competition with native invertebrates may intensify and this non-native snail could flourish in acidified coastal ecosystems.

Anthropogenic emission of carbon dioxide (CO₂) to the atmosphere has been increasing and leads to the elevation of CO₂ partial pressure (pCO₂) in the ocean. This process of ocean acidification (OA) has driven the average ocean pH to drop by 0.1 since pre-industrial times and is predicted to drive a further drop of 0.2–0.4 units by the turn of this century. Continuing OA represents a major threat to a wide array of marine organisms. Early life stages of marine invertebrates, especially calcifying larvae are particularly sensitive to OA. Known effects of OA include reduced survivorship, changes in physiological processes, and decreased calcification rates. Despite the overall negative impacts of OA, little is known about its indirect effects, including changes in ecological interactions.

The potential effect of OA on food abundance and quality is one example of such important yet little studied ecological interactions. Food quality and quantity are known to affect survival, growth and larval competence of several marine invertebrates. It is reported that OA was shown to alter the stoichiometry of algae. Increase in pCO₂ enhances inorganic carbon uptake, which in turn increases the C:N ratio, and thus, decreases algal nutritional quality. Elevated pCO₂ can also reduce or alter the polyunsaturated fatty acids (PUFAs) composition of algae, where PUFAs are critical for enzyme activity, stress resistance, growth and survival for various marine organisms. These OA-induced changes in algal nutritional value was shown to affect feeding preference and growth rate of the amphipod *Orchestoidea tuberculata*. Whether this indirect, diet quality effect of OA translates to other species remains unknown. However, increased food quantity has been suggested to help ameliorate the negative impacts of increased pCO₂, and this effect was experimentally demonstrated in the mussel *Mytilus edulis* and the barnacle *Amphibalanus improvisus*. These changes in algal abundance and nutritional content under OA condition may alter the herbivore’s feeding strategies and have yet to be fully investigated.

Some invasive species were resilient to exposure to OA alone. However, it is unclear whether the above-mentioned changes in trophic interaction under future climate condition could negatively affect their ability to colonize and spread to new habitats. One such invasive species is *Crepidula fornicata*, their survivorship under acute exposure to high temperatures was unchanged regardless of rearing pCO₂ levels (550, 750, 1,000 μatm).

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Global climate change and ocean acidification are expected to alter both phytoplankton abundance and nutrient availability in the ocean. Increased CO2 partial pressure (pCO2) has been shown to negatively affect calcification in many calcium carbonate secreting organisms, particularly those utilizing aragonite as their primary calcifying matrix. However, the interactive effects of OA and diet quality are still largely unknown. Here we exposed larvae of the slipper limpet, Crepidula onyx, to acidified rearing conditions and/or algal diets and tested if this invasive species is robust towards OA stress and could continue to flourish in its non-native habitat.

### Results

#### Seawater carbonate chemistry and algal C:N ratio.

Seawater carbonate chemistry from the three experimental trials were pooled after an initial test showed no significant differences between trials (pCO2; Kruskal–Wallis test, H2,42 = 0.310, p = 0.856, ΩAr; Kruskal–Wallis test, H2,42 = 0.938, p = 0.626, and ΩCa; Kruskal–Wallis test, H2,42 = 0.931, p = 0.628). For the larval rearing bottles, pCO2 (Kruskal–Wallis test, H2,42 = 35.880, P < 0.0001), ΩAr (Kruskal–Wallis test, H2,42 = 35.880, P < 0.0001) and ΩCa (Kruskal–Wallis test, H2,42 = 35.880, P < 0.0001) varied significantly between pH treatments (Table 1). Temperature between the three trials showed small and negligible (< 0.2 °C) difference (Kruskal–Wallis test, H2,42 = 12.643, p = 0.002).

| larval rearing | measured | calculated | C:N ratio |
|----------------|-----------|-------------|-----------|
| Control pH + Control diet | 23.44 ± 0.13 | 2155.71 ± 48.63 | 403 ± 19 | 2.88 ± 0.10 | 4.41 ± 0.15 |
| Medium pH + Control diet | 23.28 ± 0.17 | 2141.35 ± 67.07 | 963 ± 140 | 1.55 ± 0.15 | 2.38 ± 0.23 |
| Low pH + Control diet | 23.33 ± 0.05 | 2164.23 ± 40.11 | 1822 ± 89 | 0.95 ± 0.04 | 1.46 ± 0.06 |
| Control pH + Medium diet | 23.48 ± 0.19 | 2165.19 ± 39.79 | 455 ± 13 | 2.68 ± 0.06 | 4.10 ± 0.10 |
| Control pH + Low diet | 23.46 ± 0.16 | 2180.33 ± 23.75 | 441 ± 2 | 2.72 ± 0.06 | 4.17 ± 0.09 |
| Medium pH + Medium diet | 23.27 ± 0.21 | 2174.60 ± 72.64 | 979 ± 123 | 1.56 ± 0.16 | 2.39 ± 0.25 |
| Low pH + Low diet | 23.24 ± 0.14 | 2190.68 ± 29.08 | 2034 ± 162 | 0.85 ± 0.11 | 1.30 ± 0.17 |

**Table 1.** Seawater carbonate chemistry parameters throughout the experiment. Seawater total scale pH, temperature and total mean alkalinity (mean TA: 2167.55 μmol.kg⁻¹) were used to calculate CO2 partial pressure (pCO2), aragonite and calcite saturation states (respectively ΩAr and ΩCa) for a salinity of 32.0 psu using the package seacarb for R. All values are expressed as mean ± SD. Values with different letters are significantly different from each other.

| Algal culture | measured | calculated | C:N ratio |
|---------------|-----------|-------------|-----------|
| Control | 27.64 ± 0.49 | 2172.78 ± 26.80 | 31 ± 35 | 9.90 ± 1.55 | 14.11 ± 2.34 | 0.0367 | 0.0043 | 8.48 |
| Medium pH | 27.42 ± 1.55 | 2154.06 ± 39.47 | 903 ± 260 | 1.99 ± 0.62 | 2.99 ± 0.93 | 0.0464 | 0.0049 | 8.53 |
| Low pH | 27.52 ± 1.49 | 2141.54 ± 37.30 | 2146 ± 573 | 0.96 ± 0.24 | 1.45 ± 0.36 | 0.0490 | 0.0036 | 13.57 |

**Table 2.** Seawater carbonate chemistry parameters in the algal cultures and percent carbon and nitrogen content of Isochrysis galbana cultured at 3 different pH conditions. Seawater total scale pH, temperature and total mean alkalinity (mean TA: 2156.13 μmol.kg⁻¹) were used to calculate CO2 partial pressure (pCO2), aragonite and calcite saturation states (respectively ΩAr and ΩCa), for a salinity of 35.0 psu using the package seacarb for R. All values are expressed as mean ± SD. Values with different letters are significantly different from each other.

Increased pCO2 only negatively affected calcification in C. fornicata but did not affect the other physiological rates, e.g., respiration and ammonia excretion. It is unclear whether its sister species, Crepidula onyx, which has invaded the South China coast, shares the same level of resilience.

Food quality and quantity are known to affect larval development. Both C. fornicata and its sister species C. onyx have reduced growth rates at low food concentrations. For both Crepidula spp., short-term starvation and nutritional stress during the larval stage reduced post-metamorphic growth in juveniles. Larval growth of C. fornicata also varied with diet quality: larvae grew faster on a diet of Isochrysis galbana, which is rich in essential fatty acids. Food quality and quantity are therefore critical in shaping the rate of larval development. Global climate change and ocean acidification are expected to alter both phytoplankton abundance and nutrient availability in the ocean.
pH and diet had no effect on the larval mortality and respiration rate. Larval mortality of C. onyx was not significantly affected by pH treatments ($p = 0.552$) nor diet ($p = 0.272$, Fig. 1, Table 3). In addition, pH and diet interactions had no effect on mortality ($p = 0.534$). Respiration rates (nmol O$_2$ hr$^{-1}$μm BL$^{-1}$) were not significantly affected by pH treatments, diet and pH and diet interactions (Fig. S1). However, sampling days showed significant effects on the respiration rates i.e. higher oxygen consumption rates in older larvae (Fig. S1).

pH affected larval shell length, growth rate and shell integrity. Shell length (SL) on day 14 was significantly affected by pH treatments ($p = 0.0001$, Fig. 2a, Table 3). Post hoc test showed that larvae in the low pH treatment were significantly smaller the other two treatments (Tukey’s test, $p < 0.0006$). Newly settled juveniles on day 14 had a mean SL of 0.79 $\pm$ 0.04 mm in the control, 0.74 $\pm$ 0.08 mm in medium pH and 0.71 $\pm$ 0.06 mm in the low pH. However, diet had no significant effect on SL ($p = 0.468$), nor did the pH and diet interactions ($p = 0.329$). Larval shell area on day 14 also differed significantly between pH treatments ($p = 0.007$), ranging from 0.014 $\pm$ 0.002 mm$^2$ in the control pH to 0.010 $\pm$ 0.002 mm$^2$ in the low pH treatment. Growth rate (Fig. 2b) was also significantly affected by pH treatments ($p = 0.048$) and growth rate in the control was significantly higher than those in medium pH treatments (Tukey’s test, $p = 0.019$). Mean growth rate was 173.09 $\pm$ 16.60 µm log day$^{-1}$ in the control, 157.48 $\pm$ 20.69 µm log day$^{-1}$ in the medium pH and 162.85 $\pm$ 18.47 µm log day$^{-1}$ in the low pH treatment. Diet had no significant effect on the growth rate ($p = 0.139$), nor did the interactions between pH and diet ($p = 0.330$).

Results from scanning electron microscopy (Fig. 3) revealed minor but noticeable structural changes in the larval shells at low pH conditions. The growing outer edge of the nacreous layers differed between pH treatments: under control pH conditions, the shell exhibited clear nacreous layers (Fig. 3a) and a defined crystallites, granulated structures (Fig. 3b); medium pH resulted in perforated nacreous layers (Fig. 3c) and slightly eroded

| Parameters    | Source of variation | d.f. | $F$-value | $p$-value |
|---------------|---------------------|------|-----------|-----------|
| Mortality rate| pH                  | 2    | 0.61      | 0.55      |
|               | Diet                | 2    | 1.35      | 0.27      |
|               | pH x diet           | 2    | 0.64      | 0.53      |
| Shell length  | pH                  | 2    | 11.78     | 0.0001*   |
|               | Diet                | 2    | 0.78      | 0.47      |
|               | pH x diet           | 2    | 1.15      | 0.33      |
| Shell area    | pH                  | 2    | 7.90      | 0.006*    |
| Growth rate   | pH                  | 2    | 3.30      | 0.048*    |
|               | Diet                | 2    | 2.08      | 0.14      |
|               | pH x diet           | 2    | 1.14      | 0.33      |
| Settlement    | pH                  | 2    | 1.84      | 0.21      |
|               | Diet                | 2    | 3.08      | 0.10      |
|               | pH x diet           | 2    | 6.58      | 0.02*     |
| Clearance rate| pH                  | 2    | 2.40      | 0.11      |
|               | Diet                | 2    | 284.68    | <0.0001*  |
|               | pH x diet           | 4    | 2.26      | 0.09      |

Table 3. Results of the analysis of variance (ANOVA) for all larval traits measured in Crepidula onyx. *Significant results ($P < 0.05$).
periostracum layers (Fig. 3d), while low pH treatment led to nacreous layers with pits and perforations (Fig. 3e), and heavily eroded crystallite structures in the periostracum layer (Fig. 3f).

**pH and low diet quality enhanced larval settlement.** The percentage of larvae settled was not affected by pH \((p = 0.372)\) nor diet \((p = 0.367)\) alone, but was affected by pH and diet interactions \((p = 0.017)\). It appeared that larval settlement was enhanced with decreasing pH in combination with low diet quality (Tukey’s test, \(p < 0.05\), Fig. 4).

**Low diet quality enhanced clearance rate.** Clearance rate (CR) of *C. onyx* was not significantly affected by pH treatments \((p = 0.109)\) but was significantly affected by diet \((p < 0.0001\), Fig. 5, Table 3). Clearance rates

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**Figure 2.** Mean shell length at day 14 (a) and growth rate (b) of *Crepidula onyx* exposed to different pH levels and diets. pH had significant effects on both shell length and growth rate while diet nor the pH and diet interactions showed no significant effects. Due to uneven number of larvae measured, mean of means of shell length was used. Bar graphs with different letters are significantly different from each other. Error bars represent standard deviation \((n = 6)\).

**Figure 3.** Scanning electron micrographs showing the ultrastructures of day 14 *Crepidula onyx* shells exposed to control pH (a,b), medium pH (c,d) and low pH (e,f). Scale bars = 5 \(\mu\)m.
were significantly lower in the control diet than the other two lower diets (Tukey’s test, \( p < 0.001 \)). However, pH and diet interactions had no effect on the CR (\( p = 0.089 \)).

**Discussions**

Not only does ocean acidification impact larval physiology, it can also affect larval performance through indirect interactions, including the change in algal-prey nutritional value as OA reduces the nutritional quality of marine algae by elevating the C:N ratio. However, this reduction in food quality together with reduced pH (pH 7.3) did not affect the mortality and respiration rate of larval *Crepidula onyx*. Decrease in pH alone reduced *C. onyx* larval shell length, growth rate and shell structure. Low pH and low diet quality, however, appeared to promote larval settlement. Our work suggests that some species, including *C. onyx*, exhibit plasticity to cope with, if not are already well adapted to ocean acidification and low algal nutritional value.

The decrease in pH alone had a significant effect on shell size (10% decrease), growth rate (13% decrease), and shell structures. As diet did not affect shell length and growth rate, ultrastructure changes were assumed to be an effect due to pH alone. Similarly, larval shell deformities were observed in *C. fornicata* at high pCO\(_2\) (1400 µatm) condition, approximately pH\(_T\) 7.56\(^{39}\) and in *Mytilus edulis* at increasing pCO\(_2\) (1,120, 2,400 and 4,000 µatm)\(^{35}\). These changes in shell structures could imply a reduced calcification rate\(^{40}\) and/or increased dissolution. Future studies using radioactively labelled isotopes could help differentiate between these two processes. Reduction in biocalcification could indicate an energetic tradeoff\(^{38}\) as maintaining homeostasis is energetically costly, e.g., ion transport accounted for over 80% energy expenditure in larval urchin exposed to OA conditions\(^{38}\). The ecological consequence of such tradeoff, e.g., increased vulnerability to predator\(^{39}\), warrants additional studies.

Despite pH effects on shell growth and morphology, larval mortality rate and respiration of *C. onyx* were not affected by acidification. Contrary to other species that showed metabolic depression under OA conditions, e.g.,

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**Figure 4.** Percent spontaneously settled larval *Crepidula onyx* after being exposed to different pH levels and diets for ten days. No significant difference in the number of settled larvae on day 10 between pH and diet; but significant interactions between pH and diet was observed and letters indicate the post-hoc Tukey's test grouping. Error bars represent standard deviation (n = 6).

**Figure 5.** Larval clearance rate of *Crepidula onyx* estimated through incubating of known amount of larvae with known amount of algal cell for 2 hours. Significant differences in clearance rates were observed between diet qualities, with high clearance rates at low diet quality. Bar graphs with different letters are significantly different from each other. Error bars represent standard deviation (n = 4).
Littorina littorea and Mytilus chilensis, C. onyx together with C. fornicata experienced little pH impact on their respiration rate. Such resilience could be attributed on several factors. These factors include: 1) pre-exposure to low pCO2 condition in their habitats and inside brood chamber. Coastal environments in Hong Kong can experience seasonal low pH as low as 7.7 pH unit. Crepidula onyx brood their larvae in brood chambers for 2 to 4 weeks prior to release, and pH therein could be low, e.g., pH 7.0 in Ostrea chilensis and 6.4 in the calyptraeid Crepidatella dilatata; 2) presence of a large energy reserve from yolk, e.g., Gallager and Mann showed that survival of bivalve larvae positively correlated with lipid content in eggs and varied between broodstock. The Hong Kong population has relatively larger egg size (mean size of 181.75 μm) compared to the populations from California (172 μm) and Panama (157–159 μm). Further, starved larvae from this local population were able to settle though at significantly smaller sizes, which would suggests C. onyx larvae have relatively high energy reserve (Maboloc and Chan, pers. obs.); 3) presence of maternal transferred protein, e.g., larval oysters of Crassostrea sikamea collected from polluted sites are more resistant to trace metal stress due to maternal transferred metallothionein; 4) high efficiency ion transport through energetic trade-offs and or energy re-allocation (see above discussion); 5) changes in feeding behaviors (see below); or a combination of the factors above.

Low pH and low diet quality promoted larval settlement. Dooley and Pires reported a similar observation for C. fornicata, where larvae settled and metamorphosed at higher frequency at lower pH (pH 7.5 and pH 7.7) than the control (pH 8.0). Plasticity in settlement schedule could be an indication of larval stress or reduction of energy reserve as postulated by the "desperate larva hypothesis". Assuming little to no difference in post-settlement mortality this increase in settlement will likely have a positive implication on the number of individuals recruiting to the population. Alternatively, these earlier settlers could also suffer higher mortality as they could be less selective against poor settlement sites. Together with the lack of impact on larval mortality, OA and low algal nutritional value likely have little overall impact on the local population of C. onyx.

Diet quality alone has no significant effect on the larval development of C. onyx. Larvae could have met its metabolic requirements by increasing its feeding in response to low food quality as the food concentration was non-limiting in this study. Results from supplementary feeding experiments conducted with larvae from one of the females studied showed that larvae exposed to reduced pH and/or fed with low-pH grown algae, increased their clearance rates by three fold when compared to those in the control (Fig. 5, Table 3). Similar increase in clearance rates with the increasing pCO2 was also observed in C. fornicata. Blue mussel Mytilus edulis larvae also showed high feeding rates at pH 7.35. This observed increase in clearance rate could potentially incur energetic cost and affect energy allocation. However, of the small number of marine invertebrate studied, ciliary motion for both swimming (e.g., 0.5–1.5% larval energy stores h−1) and feeding (e.g., 0.502 calories day−1 in Menippe mercenaria) accounted only a small portion of their energy budget. It is however possible that under a food limiting condition, acidification-induced reduction in food quality could have negative impacts.

If the high individual resilience of C. onyx observed in the laboratory is realized as transgenerational plasticity in the field, this invasive species may have competitive advantage over the local species, which might eventually lead to shifts in community compositions under future ocean conditions. However, to fully understand invasive dynamics of C. onyx, further studies are needed to identify the impact of more realistic multiple stressors scenario and how long term exposure to these environmental changes affects its reproductive success, settlement and dispersal.

Materials and Methods
Adult collection and broodstock maintenance. Slipper limpets Crepidula onyx were collected from Victoria Harbor, Hong Kong (22°29′N, 114°17′E). Both adults and larvae acquired were reared to sexual maturity under laboratory conditions at the Coastal Marine Laboratory, Hong Kong University of Science and Technology (~7 months to first brooding). The animals were maintained in filtered (0.2 μm) seawater (pH 8.09 ± 0.10; T = 22 °C; S = 32; light/dark cycle: 12 h/12 h) with water changes every other day and with daily feeding of Isochrysis galbana at 4 × 10⁵ cells ml−1. Since adult slipper limpet C. fornicata can consume and digest larvae and zooplankton. To add more nutrients, C. onyx adults were supplemented with newly hatched Artemia nauplii at ~30 individual ml−1 twice a week. After ~7 months of rearing, egg capsules were being brooded and veligers were released into the water. The swimming veligers from each individual were collected immediately after their release through a 100 μm sieve and counted for use in subsequent experiments.

Experimental design, larval rearing, and seawater carbonate chemistry. To test the direct and diet-mediated indirect effects of ocean acidification on the development of C. onyx, the larvae were exposed to three pH levels (control pH 8.01, medium pH 7.8, and low pH 7.7) at three temperatures (22 °C, 23 °C, and 24 °C) and two salinities (31‰ and 34‰), as well as the predicted surface pH reduction of 1.4 units by 2300. A total of 7 treatments was tested: namely larvae reared at control pH and fed with algae grown in high, medium, and low pH (treatments 1–3); larvae reared at medium pH fed with algae grown in control and medium pH (treatments 4–5), and larvae reared at low pH fed with algae grown in control and low pH (treatments 6–7). There were duplicate rearing bottles for each treatment during each experimental trial. The experiment was repeated three times with larvae from different mothers. Each of the mothers was kept with two males.

Larvae were reared in 1.5 l filtered seawater at a density of 1 larva 5 ml−1 (~300 larvae per bottle) and maintained at a temperature of 23 ± 2 °C. Culture bottles were cleaned and water was completely changed with pre-equilibrated CO2 filtered seawater on day 4, day 8, and day 12 post hatching. The experiment was terminated on day 14. Larvae were fed every day (4 × 10⁵ cells ml−1) starting from day 0 with I. galbana cultured at 3 different pH conditions (see algal culturing). This ad libitum concentration was chosen based on Zhao et al. as low food concentration alone negatively affect larval growth and development. To check if food addition can cause
pH variations in the rearing bottles, pH was measured once before and after food addition. pH variations were minimal and within the experimental levels (e.g., before food addition, pH value at one low pH bottle was 7.361, after adding control food, pH value was 7.425). More importantly, all cultures were continuously aerated with a gas mixture at the experimental pCO₂ level, pH deviations stabilized quickly to the set level.

Each larval culture was continuously aerated and mixed through gentle air bubbling. The pH in the medium and low cultures were controlled by constant addition of a mix of compressed air and pure CO₂ controlled by a thermal mass flow controller (GFC 17 Aalborg, New York USA; ±1% FS accuracy). The pH, millivolt, and temperature of each culture bottle were monitored daily with Metrohm 826 pH meter and Unitrode (Herisau, Switzerland). Salinity was measured using a handheld refractometer. The pH was converted to the total scale (pHT) after calibration with TRIS (Tris/HCl) buffer solution with a salinity of 33.0 provided by the Dickson Lab at the Scripps Oceanographic Institute. Duplicate samples for total alkalinity (TA) were taken on day 4, day 8, day 12 and day 14 from all the cultures and from the newly filtered, pH equilibrated seawater used to refill the jars (n = 3). A computer-driven titration system (905 Titramounted with a glass electrode; Unitrode with Pt 1000; Herisau, Switzerland) was used to assess the TA of filtered samples (0.2 μm) with a Gran function, as described by Dickson et al. The carbonate system parameters (pCO₂, ΩAr and ΩCa) were calculated from these two measurements with the R package seacarb using the dissociation constants from Mehrbach et al. as refitted by Dickson and Millero.

Algal culturing. *Isochrysis galbana* was sub-cultured from the same algal starter and maintained in f/2 medium under 3 different pH₇ conditions (9.15 ± 0.39 as control/regular culture method, medium pH₇ 7.74 ± 0.13, and low pH₇ 7.39 ± 0.11). To achieve the medium and low pH levels, algal cultures were bubbled with pure CO₂ controlled by pH-stat systems (R-WP017 CO₂ Regulator, Easy-Aqua, Guangzhou, China). pH level in the medium was not fixed as the pH is increased during the experiment. Mortality rates (MR) were computed as the coefficient of significant linear regression system parameters (pCO₂, ΩAr and ΩCa) were calculated from these two measurements. Water volume in each culture jar was adjusted at every water change and sampling to maintain the initial density of the culture. For each culture, survival (S) was calculated for each day as the proportion of larval density divided by the maximum number of larvae counted during the experiment. Mortality rates (MR) were computed as the coefficient of significant linear regression between survival and time (% larvae day⁻¹, Table S1).

At least five larvae (8 ± 2 larvae per sampling point and treatments) from the fixed samples were photographed under the microscope (Nikon H600L, Japan) and shell length (SL) was measured with ImageJ. Growth rates (GR) were calculated for each culture as the regression coefficient of the significant logarithmic relationship between measured SL and time (Table S2). A subsample from each of the cultures was also collected on day 3, day 7 and day 14 for larval respiration measurements (supplementary method).

Nine days post hatching, three subsamples of 20 larvae from the duplicate rearing jars in each treatment were collected and placed in six-well plates with 5 ml of respective pre-equilibrated CO₂ to assess spontaneous settlement (n = 6). After 24-hours, the number of larvae swimming and settled or attached were counted.

Shell integrity and larval shell area. For shell integrity and morphology, a subsample of fixed larvae from each treatment (n = 5) were collected on day 14 and rinsed three times in phosphate buffer saline, cleared through a gradient of ethanol until 100% concentration and freeze dried. The samples were then mounted on a stub, viewed and photographed with scanning electron microscope (JEOL JSM - 6390 MA, USA) at the Materials Characterization and Preparation Facility at HKUST. Larval shell area was then measured with ImageJ.

Clearance rate. Maternal half-sibling larvae from one brood were collected from the 3 different pH cultures (control pH ~ 8.02, medium pH ~ 7.77 and low pH ~ 7.37). In each feeding experiment, 5 *C. onyx* larvae were placed into a 50 ml Falcon tube with 25 ml pre-equilibrated pH treated seawater. The larvae were acclimated for an hour before feeding and placed onto a custom made plankton rack for mixing. For each treatments, larvae were fed with 2 × 10⁵ cells ml⁻¹ of *Isochrysis galbana* cultured at 3 respective pHs. The treatments and blank procedural controls (without the larvae) were incubated for 2 hours then fixed with 4% buffered formalin. The algal concentration before and after the experiment was measured with a Beckman Z2 Coulter® Particle Count and Size Analyzer (California, USA). The clearance rate following the formula of Ginger et al. was calculated from the decrease in the algal concentration during the 2-hour feeding period ([(ln B₁ – ln B₂) – (ln C₀ – ln C₀)] V/t)/n; B₁ = blank control, initial algal concentration, B₂ = blank control, final algal concentration, C₀ = treatment, initial algal concentration, C₀ = treatment, final algal concentration, V = 25 ml, t = 2 h, n = 5 larvae and expressed in milliliters per larva per hour (ml larva⁻¹ h⁻¹). There were 4 replicates per pH and diet treatment.

Data analyses. All the statistical analyses were carried out with the software Statistica 7 and significance levels of α = 5%. All data were checked for normality (Shapiro-Wilk test) and homogeneity (Levene’s test). Data from all the experiments were pooled after an initial test (Univariante test) showed no significant differences.
References

1. Caldiera, K. & Wickett, M. E. Anthropogenic carbon and ocean pH. Nature 425, 365 (2003).
2. Ramajo, L. et al. Biominerization changes with food supply confer juvenile scallops (Argopecten purpuratus) resistance to ocean acidification. Glob. Change Biol. 22, 2025–2037 (2016).
3. Bopp, L. et al. Multiple stressors of ocean ecosystems in the 21st century: projections with CMIP5 models. Biogeosciences 10, 6225–6245 (2013).
4. Riebesell, U. et al. Reduced calcification on marine plankton in response to increased atmospheric CO2. Nature 407, 364–367 (2000).
5. Fine, M. & Tchernov, D. Scleractinian coral species survive and recover from decalcification. Science 315, 1811 (2007).
6. Kurihara, H. Effects of CO2–driven ocean acidification on the early developmental stages of invertebrates. Mar. Ecol. Prog. Ser. 373, 275–284 (2008).
7. Byrne, M. Impact of ocean warming and ocean acidification on marine invertebrate life history stages: vulnerabilities and potential for persistence in a changing ocean. In: Oceanography and Marine Biology: An Annual Review (eds Gibson, R. N., Atkinson, R. J. A., Gordon, J. D. M., Smith, I. P., Hughes, D. J.), pp. 1–42, CRC Press, Boca Raton, Florida, United States of America (2011).
8. Dupont, S., Ortega-Martínez, O. & Thorndyke, M. Impact of near-future ocean acidification on echinoderms. Ecotoxicology 19, 449–462 (2010).
9. Reynaud, S. et al. Interacting effects of CO2, partial pressure and temperature on photosynthesis and calcification in a scleractinian coral. Glob. Change Biol. 9, 1660–1668 (2003).
10. Leung, J. Y. S., Russel, B. D., Connell, S. D., Ng, J. C. Y. & Lo, M. M. Y. Acid dulls the senses: impaired locomotion and foraging performance in a marine mollusc. Anim. Behav. 106, 223–235 (2019).
11. Gaylord, B. et al. Ocean acidification through the lens of ecological theory. Ecology 96, 3–15 (2015).
12. Powell, E. N., Bohschen, E. A., Klink, J. M. & Hofmann, E. E. Influence of food quality and quantity on the growth and development of Crassostrea gigas larvae: a modeling approach. Aquaculture 210, 89–117 (2002).
13. Hettenger, A. et al. The influence of food supply on the response of Olympia oyster larvae to ocean acidification. Biogeosciences 10, 6629–6638 (2013).
14. Riebesell, U. et al. Enhanced biological carbon consumption in a high CO2 ocean. Nature 450, 545–U10 (2007).
15. Benitez, S. et al. Ontogenetic variability in the feeding behavior of a marine amphipod in response to ocean acidification. Mar. Pollut. Bull. http://dx.doi.org/10.1016/j.marpolbul.2016.07.016 (2016).
16. Duarte, C. et al. Ocean acidification induces changes in algal palatability and herbivore feeding behavior and performance. Oecologia 180, 453–462 (2016).
17. Trimborn, S., Wolf-Gladrow, D., Richter, K. U. & Rost, B. The effect of pCO2 on carbon acquisition and intracellular assimilation in four marine diatoms. J. Exp. Mar. Biol. Ecol. 376, 26–36 (2009).
18. Mercado, J. M., Javier, F., Gordillo, I., Niel, F. X. & Figueras, F. L. Effects of different levels of CO2 on photosynthesis and cell components of the red alga Porphyra leucosticta. J. Appl. Physiol. 11, 455–461 (1999).
19. Reinfelder, J. R. Carbon dioxide regulation of nitrogen and phosphorus in four species of marine phytoplankton. Mar. Ecol. Prog. Ser. 466, 57–67 (2012).
20. Rossoll, D. et al. Ocean acidification-induced food quality deterioration constrains trophic transfer. PLoS ONE 7, e47377, https://doi.org/10.1371/journal.pone.0047377 (2012).
21. King, A. L. et al. Effects of CO2 on growth rate, C:N:P, and fatty acid composition of seven marine phytoplankton species. Mar. Ecol. Prog. Ser. 537, 59–69 (2015).
22. Sanders, M. B., Bean, T. P., Hutchinson, T. H. & Le Quene, W. J. F. Juvenile king scallop, Crassostrea gigas. J. Exp. Mar. Biol. Ecol. 154, 9–20 (1990).
23. Thomsen, J. et al. Larval diet alters larval growth rates and post-metamorphic performance in the marine gastropod, Crepidula fornicata. J. Exp. Mar. Biol. Ecol. 252, 255–279 (2000).
24. Thomsen, J. & Melzner, F. Moderate seawater acidification does not elicit long-term metabolic depression in the blue mussel Mytilus edulis. Mar. Biol. 157, 2667–2676 (2010).
25. Kelly, M. W. & Hofmann, G. E. Adaptation and the physiology of ocean acidification. Funct. Ecol. 27, 980–990 (2013).
26. Lanning, G., Eilers, S., Portner, H. O., Sokolova, I. M. & Bock, C. Impact of ocean acidification on energy metabolism of oyster, Crassostrea gigas – changes in metabolic pathways and thermal response. Mar. Drugs 8, 2318–2339 (2010).
27. Pan, T. C. F., Applebaum, S. L. & Manahan, D. T. Experimental ocean acidification alters the allocation of metabolic energy. Proc. Natl. Acad. Sci. USA 112, 4696–4701 (2015).
28. Sanford, E. et al. Ocean acidification increases the vulnerability of native oysters to predation by invasive snails. Proc. R. Soc. B 281, 20132681 (2014).
40. Bibly, R., Cleall-Harding, P., Rundle, S., Widdicombe, S. & Spicer, J. Ocean acidification induced defences in the intertidal gastropod Littorina littorea. *Biol. Lett.* **3**, 699–701 (2007).

41. Navarro, J. M. et al. Impact of medium-term exposure to elevated pCO2 levels on the physiological energetics of the mussel *Mytilus chilensis*. *Chemosphere* **90**, 1242–1248 (2013).

42. Pacquet, A., Dorey, N. & Chan, K. Y. S. Ocean acidification increases larval swimming speed and has limited spawning and settlement of a robust fouling bryozoan, *Bugula neritina*. *Mar. Poll. Bull.* [https://doi.org/10.1016/j.marpollbul.2017.02.057] (2017).

43. Wang, L., Li, Q., Bi, H. & Mao, X. Human impacts and changes in the coastal waters of South China. *Sci. Total Environ.* **562**, 1017–1027 (2016).

44. Montory, J. A., Chaparro, O. R., Cubillos, V. M. & Pechenik, J. A. Isolation of incubation chambers during brooding: effect of reduced pH on protoconch development in the estuarine gastropod *Crepidula fornicata* (Calyptraeidae). *Mar. Ecol. Prog. Ser.* **374**, 157–166 (2009).

45. Gallager, S. M. & Mann, R. Growth and survival of larvae of *Meretrix mercenaria* (L.) and *Crassostrea virginica* (Gmelin) relative to broodstock conditioning and lipid contents of eggs. *Aquaculture* **56**, 165–121 (1986).

46. Collin, M. H. Worldwide patterns in the mode of development in calyptraed gastropods. *Mar. Ecol. Prog. Ser.* **247**, 103–122 (2002).

47. Collin, R. & Spangler, A. Impacts of adelphephagic development on variation in offspring size, duration of development, and temperature-mediated plasticity. *Biol. Bull.* **223**, 268–277 (2012).

48. Weng, N. & Wang, W. X. Improved tolerance of metals in contaminated oyster larvae. *Aquat. Toxicol.* **146**, 61–69 (2014).

49. Doyle, L. & Pires, A. The effect of pH on natural settlement and metamorphosis in the invasive limpet, *Crepidula fornicata*. Blinks – NSF REU – BEACON Research Fellowship 2015 (2015).

50. Botello, G. & Krug, P. J. ‘Desperate larvae’ revisited: age, energy and experience affect sensitivity to settlement cues in larvae of the gastropod *Alderia* sp. *Mar. Ecol. Prog. Ser.* **312**, 149–159 (2006).

51. Elkin, C. & Marshall, D. J. Desperate larvae: influence of deferred costs and habitat requirements on habitat selection. *Mar. Ecol. Prog. Ser.* **335**, 143–153 (2007).

52. Rodríguez, S. R., Ojeda, F. P. & Inestrosa, N. C. Settlement of benthic marine invertebrates. *Biol. Bull.* **189**, 44–54 (1995).

53. Botello, G. & Krug, P. J. ‘Desperate larvae’ revisited: age, energy and experience affect sensitivity to settlement cues in larvae of the gastropod *Alderia* sp. *Mar. Ecol. Prog. Ser.* **312**, 149–159 (2006).

54. Wendt, D. E. Energetics of larval swimming and metamorphosis in four species of *Bugula* (Bryozoa). *Biol. Bull.* **198**, 346–356 (2000).

55. Mootz, C. A. & Epifanio, C. E. An energy budget for Menippe mercenaria larvae fed Artemia nauplii. *Biol. Bull.* **146**, 44–55 (1974).

56. Feely, R. A., Sabine, C. L., Hernandez-Ayon, J. M., Ianson, D. & Hales, B. Evidence for upwelling of corrosive “acidified” water onto the continental shelf. *Science* **320**, 1490–1492 (2008).

57. Collin, R. & Spangler, A. Impacts of adelphephagic development on variation in offspring size, duration of development, and temperature-mediated plasticity. *Biol. Bull.* **223**, 268–277 (2012).

58. Weng, N. & Wang, W. X. Improved tolerance of metals in contaminated oyster larvae. *Aquat. Toxicol.* **146**, 61–69 (2014).

59. Pecquet, A., Dorey, N. & Chan, K. Y. K. Ocean acidification increases larval swimming speed and has limited spawning and settlement of a robust fouling bryozoan, *Bugula neritina*. *Mar. Poll. Bull.* [https://doi.org/10.1016/j.marpollbul.2017.02.057] (2017).

60. Lavigne, H. & Gatusso J. P. Seacarb: Seawater Carbonate Chemistry with R. R Package Version 2.4. [https://CRAN.R-project.org/package=Seacarb] (2011).

61. Mehrbach, C., Culberso, C. H., Hawley, G. & Pytkowic, R. M. Measurements of apparent dissociation constants of carbonic acid in seawater at atmospheric pressure. *Limnol. Oceanogr.* **18**, 897–907 (1973).

62. Dickson, A. G. & Millero, F. J. A comparison of the equilibrium constants for the dissociation of carbonic acid in seawater media. *Deep-Sea Res. I* **34**, 755–766 (1987).

63. Zimmermann, C. F., Keefe, C. W. & Bashee, J. Determination of carbon and nitrogen in sediments and particulates of estuarine / coastal waters using elemental analysis. In: Method 440.0, National Exposure Research Laboratory, Office of Research and Development, United States Environmental Protection Agency, pp. 440.0 - 1–440.0 - 10. Cincinnati, Ohio (1997).

64. Schneider, C. A., Rasband, W. S. & Eliceiri, K. W. NIH image to ImageJ: 25 years of image analysis. *Nat. Methods* **9**, 671–675 (2012).

65. Ginger, K. W. K. et al. Larval and post-larval stages of Pacific oyster (*Crassostrea gigas*) are resistant to elevated CO2. *PloS ONE* **8**, e64147, [https://doi.org/10.1371/journal.pone.0064147] (2013).

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Author Contributions

E.A.M. and K.Y.K.C. conceived and designed the experiments. E.A.M. conducted the experiments and both E.A.M. and K.Y.K.C. analyzed the data and wrote the manuscript.

Additional Information

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