The impact of environmental acidification on the microstructure and mechanical integrity of marine invertebrate skeletons

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Ocean acidification (OA), from seawater uptake of anthropogenic CO2, has a suite of negative effects on the ability of marine invertebrates to produce and maintain their skeletons. Increased organism pCO2 causes hypercapnia, an energetically costly physiological stress. OA alters seawater carbonate chemistry, limiting the carbonate available to form the calcium carbonate (CaCO3) minerals used to build skeletons. The reduced saturation state of CaCO3 also causes corrosion of CaCO3 structures. Global change is also accelerating coastal acidification driven by land-run off (e.g. acid soil leachates, tannic acid). Building and maintaining marine biomaterials in the face of changing climate will depend on the balance between calcification and dissolution. Overall, in response to environmental acidification, many calcifiers produce less biomineral and so have smaller body size. Studies of skeleton development in echinoderms and molluscs across life stages show the stunning effect of OA. For corals, linear extension may be maintained, but at the expense of less dense biomineral. Conventional metrics used to quantify growth and calcification need to be augmented by characterisation of the changes to biomineral structure and mechanical integrity caused by environmental acidification. Scanning electron microscopy and microcomputed tomography of corals, tube worms and sea urchins exposed to experimental (laboratory) and natural (vents, coastal run off) acidification show a less dense biomineral with greater porosity and a larger void space. For bivalves, CaCO3 crystal deposition is more chaotic in response to both ocean and coastal acidification. Biomechanics tests reveal that these changes result in weaker, more fragile skeletons, compromising their vital protective roles. Vulnerabilities differ among taxa and depend on acidification level. Climate warming has the potential to ameliorate some of the negative effects of acidification but may also make matters worse. The integrative morphology-eomechanics approach is key to understanding how marine biominerals will perform in the face of changing climate.

Key words: Climate change, marine biominerals, corals, molluscs, sea urchins, serpulid worms

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Introduction

Ocean acidification (OA) resulting from increased ocean uptake of CO₂, driven by the increase in anthropogenic greenhouse gas emissions, is unprecedented on geological time scales (Zeebe et al., 2016). This uptake is causing major changes to ocean chemistry with a suite of negative effects on the ability of marine invertebrates to produce and maintain their skeletons. Firstly, increased organism pCO₂ causes hypercapnia, a stress that can affect many physiological processes. Hypercapnia can be energetically costly reducing the resources that animals need to calcify because maintenance of essential metabolic processes (e.g. acid-base homeostasis) takes priority over diverting energy to growth and calcification (Stumpp et al., 2012; Pan et al., 2015). The production and maintenance of calcium carbonate (CaCO₃) structures has been shown to be modulated by energy acquisition (Melzner et al., 2011). Secondly, OA reduces the saturation state of CaCO₃ minerals, limiting the carbonate available for calcification (IPCC, 2014). These minerals are the building blocks of biocalcification and include aragonite and calcite. Thirdly, the direct corrosive effect of OA can cause pitting and erosion of CaCO₃ structures, as reported for calcifiers that are resident in low pH CO₂ seep environments (Harvey et al., 2018). Calcite is less susceptible to dissolution at lower pH values than aragonite, unless it contains high levels of magnesium (Ries et al., 2009; Chan et al., 2012). However, calcite is more brittle compared to aragonite, making it mechanically weaker (Fitzer et al., 2015a; Meng et al., 2018).

Due to sea-level rise and changing weather patterns, global change is exacerbating coastal acidification in areas where freshwater runoff results in reduced pH due to leachate from acid sulphate soils and humic acids and tannic acids from groundwater (Amaral et al., 2011, 2012; Duarte et al., 2013; Jiang et al., 2017; Fitzer et al., 2018). Environmental acidification in a changing ocean is being caused by both ocean and coastal acidification. These two forms of acidification (CO₂ and land run off) differ in chemistry mechanisms (Fig. 1). Environmental acidification also occurs at CO₂ seeps (Foo et al., 2018; González-Delago and Hernández, 2018).

Marine invertebrate skeletons are complex and ancient structures (Wilkinson, 1979) produced through the superb biocatalyst control of mineral production, resulting in a great diversity of taxon-specific structures (Wilbur, 1972; Cavey and Markel, 1994; Von Euw et al., 2017). For a broad range of marine invertebrates (e.g. corals, molluscs, echinoderms) experimental acidification in laboratory studies and environmental acidification at natural CO₂ seeps and in low pH coastal waters (Amaral et al., 2011, 2012; Duarte et al., 2013; Jiang et al., 2017; Fitzer et al., 2018) show the negative effect of acidification on biominalisation. Faced with these challenges, building and maintaining calcified structures by marine invertebrates will depend on the balance between calcification and dissolution.

Calcified structures are expensive to produce (Comeau et al., 2017) and, in response to the physiological challenges presented by OA, many calcifiers produce smaller skeletons and so have smaller body sizes. This is seen in the stunting effect of OA at the larval and adult stage of molluscs and echinoderms (Parker et al., 2012, 2013; Byrne et al., 2013; Garilli et al., 2015; Dworjanyn and Byrne, 2018). Vulnerabilities with respect to the amount of biomineral produced, its mechanical properties and propensity for dissolution or etching in OA conditions differ greatly among taxa, even within the same phylum that have similar calcification mechanisms (Table 1). Most laboratory studies have involved translocation of animals that have calcified (i.e. ‘grown up’) in control conditions to OA conditions for short- or long-term exposure, an approach that shows the effects of OA on established skeleton and the ability to make new skeleton. For insight into the inherent differences of the production of biomineral in normal ambient and acidification conditions, many studies have reared larvae and juveniles under OA (Parker et al., 2012, 2013; Byrne et al., 2013; Dworjanyn and Byrne, 2018) or have availed of naturally low pH habitats (e.g. CO₂ seeps or low pH coastal waters) where biominalisation has occurred through life (Crook et al., 2013; Garilli et al., 2015; Fitzer et al., 2018; Foo et al., 2018; González-Delago and Hernández, 2018; Migliaccio et al., 2019).

The impact of OA on calcification depends on level of pCO₂, with near future projections (e.g. pH 7.8) having milder impacts than far future ones (e.g. pH 7.6 and lower) and also depends on the duration of exposure (see Table 1). For instance, many calcifiers can reside at a mean pH 7.7–7.9 at low pH vent sites zones through their life, but are absent at zones with a mean pH ≤ 7.6 (Calosi et al., 2013; Kroeker et al., 2013; Foo et al., 2018). The impact of environmental acidification also varies among regions and habitats with some species or populations appearing to be adapted or phenotypically adjusted to environmental acidification. This is seen in the resilience of sea urchins living in low pH upwelling zones and vent sites (Calosi et al., 2013; Hofmann et al., 2014; Migliaccio et al., 2019). Some molluscs can alter the amount of the form of CaCO₃ produced (e.g. aragonite or calcite) to a more favourable form making them more resilient to acidification (Fitzer et al., 2014a, 2016; Langer et al., 2014; Rühl et al., 2017).

In calcifying marine invertebrates, environmental carbon for calcification such as dissolved inorganic carbon (DIC) can be sourced in the form of CO₃²⁻ or hydrogen carbonate (HCO₃⁻) from ambient seawater. Under CO₂-driven acidification, this may impair shell growth as pH and CaCO₃ saturation are lowered resulting in reduced carbonate available for biominalisation (Doney et al., 2009). Therefore, under OA, where CO₃²⁻ becomes less available from seawater limited shell growth and increased abnormalities become a coping mechanism for continued biominalisation (Wittmann and Pörtner, 2013). However, respiratory CO₂ can also be a source of carbon for calcification where it is used to form
Table 1: Studies that have investigated the impacts of environmental acidification on the microstructure and/or mechanics of marine invertebrate skeletons and methods used. pH levels are listed as published on the total (pHT) or NBS (pHNBS) scales or no scale (pH*) if not indicated.

| Phylum/ species | Time | pH or Ω levels/location | Morph methods | Morph results | Biomechanics methods | Biomechanics results | References |
|-----------------|------|--------------------------|---------------|--------------|----------------------|----------------------|------------|
| Cnidaria        |      |                          |               |              |                      |                      |            |
| Balanophyllia europaea | Resident | pHT 7.7, 7.9, 8.1; Vent | μCT, SEM | pH 7.7, 21-31% ↑ in porosity & 7% ↓ in bulk density | Nanoindentation      | pH 7.7, hardness - no effect of pH; ↓ in stiffness | Fantazzini et al., (2015) |
| Favia fragum    | 8 d to juvenile** | Ω Ar 0.22-3.71; Lab | SEM | Ω Ar ≤ 1; change in crystal size, shape & orientation, thin septa | - | - | Cohen et al., (2009) |
| Porites astreoides | Resident | Ω Ar < 1-3.71; Coast | μCT | Ar < 2; ↓ density | - | - | Crook et al., (2013) |
| Stylophora pistillata | 12 mo | pHT 7.2, 7.4, 7.8, 8.0; Lab | μCT | pH 7.2-7.4, ↑ in porosity & ↓ in bulk density, thinner skeleton, no change in crystallography | - | - | Tambutté et al., (2015) |
| Stylophora pistillata | 14 mo | pH NBS 7.3, 7.6, 8.2; Lab | SEM, EBSD | pH NBS 7.3, 7.6, crystallographic changes, shorter less round fibre bundles | - | - | Coronado et al., (2019) |
| Brachiopoda     |      |                          |               |              |                      |                      |            |
| Magelania venosa | 335 d | pH+ 7.35-8.15 Lab, ASW | SEM | pH+ 7.35-7.6, ↑ pore size; ↑ thickness primary layer; ↓ thickness secondary layer; more organic rich shell | - | - | Ye et al., (2018) |
| Annelida        |      |                          |               |              |                      |                      |            |
| Hydroides elegans | 9-18 d** | pHNBS 7.4, 7.6, 7.8, 9.8, 8.0; Lab | μCT, SEM | pHNBS 7.4-7.8, ↑ porosity, irregular layers, surface pitting & erosion, thinner; change in mineral layers & crystallography | Nanoindentation, crushing test | pHNBS 7.4-7.8, ↓ hardness, elasticity & 62% ↓ in crushing force | Chan et al., (2012); Li et al., (2014; 2016) |
| Mollusca         |      |                          |               |              |                      |                      |            |
| Gastropoda      |      |                          |               |              |                      |                      |            |
| Austrocochlea constricta | 4 mo | pHNBS 7.8, 8.1; Lab | - | - | Vickers hardness, Elastic modulus | pHNBS 7.8, ↑ shell hardness, ↑ stiffness | Leung et al., (2017) |
| Austrocochlea adonis | 4 mo | pHNBS 7.86, 8.0; Lab | - | - | Vickers hardness, Elastic modulus | No effect of pH | Leung et al., (2017) |
| Phylum/species          | Time    | pH or Ω levels/location | Morph methods | Morph results          | Biomechanics methods                  | Biomechanics results                     | References                      |
|-------------------------|---------|-------------------------|---------------|------------------------|---------------------------------------|-----------------------------------------|----------------------------------|
| Austrocochlea porcata   | 95d     | pH_{NBS} 7.7, 7.9, 8.1; Lab | -             | No effect of pH        | Crushing test                         | pH_{NBS} 7.7, ↓ shell strength by 28%  | Coleman et al., (2014)           |
| Bulla quoyi             | 4 mo    | pH_{NBS} 7.9, 8.1; Lab   | -             | -                      | Vickers hardness, Elastic modulus      | No effect of pH                         | Leung et al., (2017)            |
| Charonia lampas         | Resident| pH_{T} 7.8, 8.1; Vent   | μCT           | pH_{T} 7.8, Two-fold ↓ in density and thickness | -                                     | -                                       | Harvey et al., (2018)       |
| Columbella rustica      | 3 mo    | pH_{NBS} 7.6, 8.1, Lab   | μCT           | pH_{NBS} 7.6, 0.8-8% ↓ in density depending on shell region | -                                     | -                                       | Chatzinikolaou et al., (2017) |
| Littorina littorea      | 5 mo (small and large) | pH_{NBS} 7.8, 8.0; Lab | -             | -                      | Crushing test                         | pH_{NBS} 7.8, Large snails, ↓ in crushing force, Small snails, no effect of pH | Landes and Zimmex (2012) |
| Nassarius nitidus       | 3 mo    | pH_{NBS} 7.6, 8.1, Lab   | μCT           | pH_{NBS} 7.6, 38-51% ↓ in density depending on shell region | -                                     | -                                       | Chatzinikolaou et al., (2017) |
| Nerita atraentosa       | 4 mo    | pH_{NBS} 7.8, 8.1; Lab   | -             | -                      | Vickers hardness, Elastic modulus      | No effect of pH                         | Leung et al., (2017)            |
| Nucella lapillus        | 14 mo (juveniles) | pH_{T} 7.8, 7.9, 8.0; Lab | μCT, SEM     | pH_{T} 7.8, 20-30% ↓ in density; surface erosion, lack of layering in shell, younger shells thinner, older shells thicker | -                                     | -                                       | Queirós et al., (2015); Rühl et al., (2017) |
| Nucella ostrina         | 6 mo.   | pH_{T} ~ 7.6, 8.1; Lab   | -             | -                      | Crushing test                         | 10% reduction in strength               | Barclay et al., (2019)          |
| Phasianella australis   | 4 mo    | pH_{NBS} 7.9, 8.1; Lab   | -             | -                      | Vickers hardness, Elastic modulus      | No effect of pH                         | Leung et al., (2017)            |
| Phorcus sauciatus       | Resident| pH_{NBS} 7.52-8.2 Vent   | -             | Vent site – corrosion and cracks in shell | Crushing test                         | Weaker shells                          | Viotti et al., (2019)          |
| Subnarella undulata     | 65 d    | pH_{NBS} 7.7, 7.9, 8.1; Lab | -             | pH_{NBS} 7.7, ↓ shell thickness | Crushing test                         | No effect of pH                         | Coleman et al., (2014)         |
| Tegula funebralis       | 6 mo.   | pH_{T} 7.4-8.1; Lab      | -             | -                      | Crushing test                         | 50% reduction in strength               | Barclay et al., (2019)          |
| Thalotia conca          | 4 mo    | pH_{NBS} 7.9, 8.1; Lab   | -             | -                      | Vickers hardness, Elastic modulus      | No effect of pH                         | Leung et al., (2017)            |
| Turbo undulatus         | 4 mo    | pH_{NBS} 7.86, 8.0; Lab   | -             | -                      | Vickers hardness, Elastic modulus      | No effect of pH                         | Leung et al., (2017)            |
| Phylum/species         | Time       | pH or Ω levels/location | Morph methods | Morph results | Biomechanics methods | Biomechanics results | References                  |
|------------------------|------------|-------------------------|---------------|---------------|----------------------|----------------------|---------------------------|
| **Bivalvia**           |            |                         |               |               |                      |                      |                           |
| Adamussium colbecki    | 1 mo       | pH T 7.6, 8.1; Lab      | SEM           | No effect on crystals | Nanoindentation      | No effect of pH     | Dell’Acqua et al., (2019) |
| Arctic islandica       | 3 mo       | pH NBS 7.7, 7.9, 8.0   | SEM           | No effect on shape or size of crystals | -                     | -                     | Stemmer et al., (2013)    |
| Cerastoderma edule     | 2 mo       | pH NBS, 6.4, 6.7, 7.0, 7.4, 7.8, Lab | SEM | pH NBS 6.4, 6.7 ↓ net calcification, no change in dissolution | Nanoindentation      | No effect of pH     | Milano et al., (2016)    |
| Crassostrea virginica  | 20 wk (juveniles) 2 wk (adults) | pH NBS 7.5, 8.2, Lab | FTIR SEM      | pH NBS 7.5 40% ↓ in shell mass in juveniles | Microindentation, fracture toughness | pH NBS 7.5 ↓ calcite hardness and fracture toughness in juveniles | Beniash et al., (2010) |
| Crassostrea virginica  | 11 wk      | pH NBS 7.97/8.1 8.11/8.36 Lab | SEM           | No difference in shell body mass | Microindentation, fracture toughness | pH NBS 7.9/8.1 ↓ hardness and fracture resistance at salinity | Dickinson et al., (2012) |
| Magallana angulata     | 35 d early juveniles | pH NBS 7.8, 8.1 Lab | SEM, EBSD μCT, pH NBS 7.8 ↑ porosity of microstructure, ↓ density | Nanoindentation | pH NBS 7.8 ↓ hardness and stiffness | Meng et al., (2018) |
| Magallana gigas        | 6 wk       | pH NBS, 7.7, 8.1, Lab   | -             | -             | Crushing test        | pH NBS 7.7 ↓ crushing force | Wright et al., (2018) |
| Mytilus californianus  | 8 d larvae** | pH NBS 7.8, 8.0, 8.1, Lab | SEM           | pH NBS 7.8 15% thinner | Crushing test        | pH NBS 7.8 15-20% weaker pH NBS 8.0 13-15% weaker | Gaylord et al., (2011) |
| Mytilus edulis         | 6 mo       | pH NBS 7.2, 7.3, 7.4, 7.5, 7.7, 8.1, 8.2, Lab | SEM, EBSD | pH NBS 7.2, 7.3, 7.4, 7.5, 7.7 disorganised crystals & altered layer structure | -                     | -                     | Fitzer et al., (2014b)   |
| Mytilus edulis         | 6 mo (juveniles) | pH NBS 7.2, 7.3, 7.4, 7.5, 7.7, 8.1, 8.2, Lab | SEM, EBSD | pH NBS 7.2, 7.3, 7.4, 7.5, thinner calcite layer and altered layer structure and crystallography | -                     | -                     | Fitzer et al., (2014a)   |
| Mytilus edulis         | 6 mo       | pH T 7.65, 8.0, Lab     | -             | -             | Crushing test        | pH T 7.65 ↓ flex before failure; Strength not affected | Mackenzie et al., (2014) |
| Mytilus edulis         | 6 mo       | pH NBS 7.2, 7.3, 7.4, 7.5, 7.7, 8.1, 8.2, Lab | -             | -             | Microindentation, fracture toughness, nanoindentation | pH NBS 7.4, 7.5, 7.7 ↓ calcite hardness and ↓ fracture toughness | Fitzer et al., (2014a) |
| Mytilus edulis         | 2 mo       | pH NBS 7.8, 8.1, Lab    | SEM           | Disordered crystals | Crushing test        | pH NBS 7.8 ↓ crushing strength 22-24% | Li et al., (2015) |

*Continued*
| Phylum/species     | Time     | pH or $\Omega$ levels/location | Morph methods           | Morph results                                                                 | Biomechanics methods | Biomechanics results | References                  |
|-------------------|----------|---------------------------------|-------------------------|-------------------------------------------------------------------------------|----------------------|----------------------|-----------------------------|
| Mytilus edulis    | 6 mo     | $pH_{NBS}$ 7.4, 8.1, Lab         | SEM-EBSD, XPEEM         | $pH_{NBS}$ 7.2, altered crystallography structure and more ACC.               | -                    | -                    | Fitzer et al., (2016)       |
| Mytilus edulis    | 9 months | $pH_{NBS}$ 7.2, 7.3, 7.4, 7.5, 7.7, 8.1, 8.2, 8.1, Lab | Shell shape analysis, shell thickness index (STI) | $pH_{NBS}$ 7.2, 7.3, 7.4, 7.5, 7.7, ↓ shell thickness and rounder flatter shells. | -                    | -                    | Fitzer et al., (2015b)      |
| Mytilus edulis    | 7 weeks  | $pH_{NBS}$ 7.2, 7.4, 7.7, 8.0, Lab | SEM                     | $pH_{NBS}$ 7.2, corrosion of internal aragonite layers                       | -                    | -                    | Melzner et al., (2011)      |
| Mytilus edulis    | 6 mo     | $pHT$ 7.65, 8.0, Lab             | -                       | Crushing test $pHT$ 7.65 $\downarrow$ flex before failure; Strength not affected | -                    | -                    | Mackenzie et al., (2014)    |
| Mytilus galloprovincialis | 68 d  | $pHT$ 7.25, 8.07; Vent transplant | SEM, EBSD               | $pHT$ 7.25, thinner shell, disturbed less ordered structure                    | -                    | -                    | Hahn et al., (2012)         |
| Mytilus galloprovincialis | 21 d - 5 mo | $pHT$ 6.8, 7.2, 7.8, Vent transplant to lab | SEM                     | Calcification continued in vents; Isolated shells, dissolution at low $pH$      | -                    | -                    | Rodolfo-Metalpa et al., (2011) |
| Pinctada fucata   | 28 d     | $pH_{NBS}$ 7.6, 7.8, 8.1         | SEM                     | Shell dissolution                                                              | Crushing test         | $pH_{NBS}$ 7.6, 25.9% and 26.8% weaker shells | Welldsen et al., (2011) |
| Saccostrea glomerata | 2 yr  | $pH_{NBS}$ 7.7, 8.1, Coast++      | SEM-EBSD                | $pH$ 7.7, disordered crystallographic structure                              | -                    | -                    | Fitzer et al., (2019b)      |
| Saccostrea glomerata | N/A   | $pH_{NBS}$ 6.6, 6.8, 6.9, 7.8, 7.9, Coast++ | -                       | -                                                                              | Crushing test         | pH $pH_{NBS}$ 6.6, 6.8, 6.9, Weaker shells | Amaral et al., (2012); Fitzer et al., (2019b) |
| Cephalopoda       |          |                                 |                         |                                                                                |                      |                      |                             |
| Argonauta nodosa  | N/A      | $pHT$ 7.35, 7.55, 7.8, 8.0, Lab   | SEM, EBSD               | Shell only $pHT$ 7.35-7.8, dissolution and etching, altered crystallography and structure | -                    | -                    | Wolfe et al., (2012, 2013a) |
| Echinodermata     |          |                                 |                         |                                                                                |                      |                      |                             |
| Diadema africanum | 100 d, juveniles | $pHT$ 7.6, 8.0, Lab | SEM                     | $pHT$ 7.6, Test plates thinner, spines, dissolution/etching                 | Crushing test (whole urchin dried) | $pHT$ 7.6, $\downarrow$ crushing force | Rodríguez et al., (2017)    |
| Echinometra sp.   | 343 d    | $pH_{NBS}$ 7.7, 8.1, Lab          | SEM                     | Test plates and sines, no dissolution                                         | -                    | -                    | Hazan et al., (2014)        |
| Phylum/species           | Time          | pH or Ω levels/location | Morph methods | Morph results                                                                 | Biomechanics methods                      | Biomechanics results                                    | References                           |
|-------------------------|---------------|-------------------------|---------------|--------------------------------------------------------------------------------|--------------------------------------------|----------------------------------------------------------|--------------------------------------|
| *Echinometra mathie*    | 13 mo         | pH T 7.65, 8.1, Lab     | -             | -                                                                               | Ambital and apical plate fracture force and elasticity | No effect of pH                                           | Moulin *et al.*, (2015)             |
| *Heliocidaris erythrogramma* | 2 wk, early juveniles | pH NBS 7.4, 7.6, 7.8, 8.1, Lab | SEM           | pH NBS 7.4, Spines, ↑ pore size and dissolution | -                                          | -                                                        | Wolfe *et al.*, (2013b)            |
| *Heliocidaris erythrogramma* | 9 mo     | pH NBS 7.6, 8.1, Lab     | SEM           | pH NBS 7.6, ↑ pore size apical but not ambital plates | Nanoindentation, elasticity                | pH NBS 7.6, ↓ hardness and elasticity                     | Johnson and Byrne (unpublished data) |
| *Lytechinus variegatus* | 3 mo, early juveniles | pH 7.8, 8.0, 8.1, Lab      | SEM           | pH 7.4, Spines, dissolution and malformation | -                                          | -                                                        | Albright *et al.*, (2012)          |
| *Lytechinus variegatus* | 59 d          | pH 7.47, 7.7, 7.9, Lab   | SEM           | pH 7.47, Spines, reduced barbs | Snap test                                  | pH 7.47, ↓ snap force                                     | Emerson *et al.*, (2017)           |
| *Paracentrotus lividus* | 1 mo (juveniles) | pH 7.7, 7.8, 8.0, 8.1, Lab | SEM           | pH 7.7, larger pore size in tooth, no change test plate thickness; no spines dissolution/etching | Crushing test (whole urchin dried) Fracture force plate | pH 7.7, ↓ crushing force; No effect of pH on test plate | Asnaghi *et al.*, (2013, 2014, 2019) |
| *Paracentrotus lividus* | 12 mo         | pH NBS 7.8, 7.9, 8.0, 8.1, Lab | -             | -                                                                               | Crushing tests, ambital and apical plate fracture force, nanoindentation, elasticity | No effect of pH                                           | Collard *et al.*, (2016)           |
| *Paracentrotus lividus* | Resident       | pH NBS 7.8, 8.2, Vent    | -             | -                                                                               | Crushing force, ambital and apical plate fracture force, nanoindentation, elasticity | No effect of pH                                           | Collard *et al.*, (2016)           |
| *Paracentrotus lividus* | 100 d, juveniles | pH T 7.6, 8.0, Lab       | SEM           | pH T 7.6, Test plates thinner; Spines, no dissolution/etching                  | Crushing test (whole urchin dried)         | pH T 7.6, ↓ crushing force                                | Rodríguez *et al.*, (2017)         |
| *Tripneustes gratilla*  | 146 d juvenile to adult | pH NBS 7.6, 7.8, 8.1 Lab | SEM           | pH NBS 7.6, thinner test, spines nodissolution/etching | Crushing test (live whole urchin) Fracture force, rupture at sutures not skeleton | pH NBS 7.6, ↓ crushing force | Byrne *et al.*, (2014)           |

*Table 1: Continued*
Table 1: Continued

| Phylum/species                  | Time  | pH or Ω levels/location | Morph methods | Morph results | Biomechanics methods | Biomechanics results | References                  |
|--------------------------------|-------|--------------------------|---------------|--------------|----------------------|----------------------|----------------------------|
| *Tripneustes ventricosus*      | 5 wk  | pH 7.4, 7.7, 8.1, Lab    | SEM           | pH 7.4, 7.7, spine etching Test plates, no dissolution/etching | Fracture force spines, two-point bending, elasticity | pH 7.4, 7.7, spine more brittle, ↓ fracture force, 35% and 16%, no effect of pH on elasticity | Dery et al. (2017) |
| *Strongylocentrotus droebachiensis* | 45 d  | pH 7.2, 7.7, 8.0, Lab    | -             | pH 7.2 ~ 7.2, test plates pitted; spine dissolution | Fracture force spines | pH 7.2 ~ 7.2, spines, ↓ fracture force; No effect of pH on test plates | Holtmann et al. (2013) |
| **Crustaceans**                |       |                          |               |              |                      |                      |                            |
| *Amphibalanus amphitrite*      | 86 d-20 wk | pH 7.4, 8.2, Lab         | -             | -            | Penetrometry, breaking test | pH 7.4, ↓ force to penetrate shells | McDonald et al. (2009) |
| *Amphibalanus improvisus*      | 86 d-20 wk | pH 7.3-7.9 Lab           | SEM           | Low pH increasing corrosion | Crushing test | No effect of pH | Panch et al. (2013, 2014) |

Morphology methods: EBSD, electron backscatter diffractometry; FTIR, Fourier-transform infrared spectroscopy; μCT, micro-computed tomography; SEM, scanning electron microscopy; XPEEM, photoemission electron microscopy. Biomechanics methods: crushing tests (force of rupture); nanoindentation (hardness; Young’s modulus of elasticity — a measure of stiffness); ACC, amorphous calcium carbonate; ASW, artificial seawater; Coast, = coastal runoff pH gradient; Lab, laboratory; Morph, morphology; Ω, saturation state calcium carbonate minerals; Time, exposure duration; Vent, CO₂ seep pH gradient; —, not investigated; ≈, larval to juvenile studies; all others are of adults; ***, includes pH flux conditions.

Despite concerns for the prospects for marine calcifiers in a changing ocean, the impacts of environmental acidification on the structure of the biomineral itself remain largely unexplored. We need to understand how acidification alters biomineral production and its microstructure and mechanical integrity to address uncertainties about how changes to marine skeletons exposed to environmental acidification (Table 1) can affect the survival and success of a vast diversity of marine species as they play essential roles in body support and protection. Corals and oysters, which provide structural foundation for habitat, follow a different course in this regard than more mobile species, as they are more likely to be vulnerable to OA (Hoegh-Guldberg et al., 2007; Kroeker et al., 2013; Dore et al., 2013; Dinelli et al., 2013; Byrne et al., 2016). The mineralogy of marine skeletons and their vulnerability to OA has also been reviewed (Rees, 2016; Smith et al., 2013, 2016).

The production and maintenance of biomineral structures, and the contribution of DIC sources required for calcification vary between calcifying species. Therefore, an understanding of how OA may be species specific. This is an important consideration in understanding the vulnerability of species and their calcification response in ocean and coastal acidification conditions.
Figure 1: Equations for the mechanisms of acidification from CO₂-OA and other forms of environmental acidification for example from acid sulphate soil leachates. Modified from Fitzer et al. (2018).

Ocean acidification (OA) CO₂-acidification:
- Atmospheric CO₂
- CO₂ vents

\[ \text{CO}_2 + \text{H}_2\text{O} \leftrightarrow \text{H}_2\text{CO}_3 \leftrightarrow \text{HCO}_3^- + \text{H}^+ \leftrightarrow \text{CO}_3^{2-} + 2\text{H}^+ \]

Other forms of environmental acidification:
- Coastal freshwater runoff
- Sulphate soil coastal acidification:

\[ \text{FeS}_2\text{O}_7 + \text{7/2 O}_2\text{aq} + \text{H}_2\text{O} \rightarrow \text{Fe}^{3+}\text{aq} + 2 \text{ SO}_4^{2-}\text{aq} + 2 \text{H}^+\text{aq} \] (1)

\[ \text{Fe}^{2+}\text{aq} + \text{1/4 O}_2\text{aq} + \text{3/2 H}_2\text{O} \rightarrow \text{FeO}OH_{\text{aq}} + 2 \text{H}^+\text{aq} \] (2)

The list of studies (Table 1) provides an overview of the impacts of environmental acidification on skeletal structure and biomechanics and the pH levels and scales (e.g. pHT vs pHNBS) used are important to note. In the following text, we review research on the major marine calcifying taxa (tube worms, corals, molluscs, echinoderms) with a focus on those most studied, bivalves and sea urchins. Only a few studies have investigated the impact of habitat acidification and warming, and the interaction of these factors, on the structure and integrity of marine biomineral, and we highlight a few of these.

**Corals**

Corals produce an aragonite skeleton that is covered by the tissues of the coral polyps (Von Euw et al., 2017). Several studies have investigated the impact of OA on the microstructure of coral skeletons using SEM or μCT and one study has incorporated mechanical tests (Table 1). The number of studies appears surprisingly limited considering the ecological importance of reef-building corals. For this important taxon more research is needed on the impact of projected near-future acidification on skeletal structure and mechanical integrity and in context with habitat warming.

The temperate solitary cup coral *Balanophyllia europaea* living at a low pH vent site had a more porous and thinner skeleton than those living at nearby control sites (Fantazzini et al., 2015). Skeletal pore size increased and bulk density decreased, with decreasing pH (8.1, 7.9, 7.7). However, growth, with respect to size, was maintained by these corals along the vent pH gradient. Nanoindentation revealed that the skeleton of *B. europaea* living at low pH (pH 7.7) had a lower hardness and stiffness compared with corals residing at pH 8.1. While this species is able to maintain growth at reduced pH, the deposition of less biomineral is likely to increase susceptibility to physical damage (Fantazzini et al., 2015).

For colonies of the tropical coral, *Stylophora pistilla*, exposed to pH 7.2 for 12 months, SEM and μCT revealed that the biomineral produced had greater porosity and lower bulk density compared with conspecifics maintained at ambient pH (Table 1). Note the caveat that the low pH treatments used are well beyond OA scenarios (IPCC, 2014) (as the case for many of the studies listed in Table 1). Tambutté et al., (2015) suggest that these extreme treatments are useful to help identify trends. While the growth of *Stylophora pistilla*, as linear extension did not change in pH 7.2, the density of the biomineral did, potentially as a trade-off strategy under competition to maintain access to light and space with the compromise of a weaker skeleton (Tambutté et al., 2015). This change in the skeleton also reflects the decline in *S. pistilla* colonies at low pH sites in nature (Tambutté et al., 2015). A recent study of these species maintained at low pH for 14 months showed a change in skeletal chrolstallography (Coronado et al., 2019).

Colonies of *Porites astreoides* living in a naturally low pH environment from coastal run off also had a lower density skeleton (Crook et al., 2013) and juvenile *Favia fragum* exposed to CO₂ driven under saturation conditions (pH not provided) for 8 days had a thinner skeleton compared with controls (Cohen et al., 2009).
In these coral studies skeletal dissolution was not noted. It appears that the thin veneer of polyp/coralite tissue protects the underlying skeleton from dissolution. Importantly, the most universal measure of coral health and biomineralisation rate (linear extension) is unlikely to be a reliable metric of coral health and growth in the face of OA, with potential production of more fragile phenotypes (Tambutté, 2015).

**Tube worms**

Serpulid polychaetes are a major group of marine worms that produce a calcareous tube that varies in composition from being entirely aragonitic, to entirely high-Mg calcite to mixtures of these two (Smith, 2013). The serpulid, Hydroides elegans, is one of the most important species in tropical biofouling communities and the microstructure and mechanics of its calcareous tube has been investigated in the OA context (Chan, 2012; Li et al., 2014, 2016).

In studies where H. elegans was reared in experimental OA conditions across larval development through to tube formation, SEM and μCT revealed that at low pH (7.4–7.8), the tube had increased porosity, decreased thickness and had surface pitting and erosion. There was also a change in the mineral layers and crystallography. These changes reduced the hardness of the tube. The force needed to crush the tube decreased by 62%. The composition of aragonite in the tube is also influenced by OA with H. elegans depositing more calcite in the tube at low pH. This species seems to be able to adjust the mineralogy of its tube as a potential adaptation to the changes in water chemistry driven by OA, but this compromises the strength and elasticity of the tube (Li et al., 2014, 2016). In comparison, the serpulid, Spirobranchus triqueter was similarly fragile when reared under moderate (pH 7.7) and severe (pH 7.4) reductions in pH. Tube fracture toughness was not linearly related to the reduced pH, but was related to changes in porosity as found for H. elegans. Larger pores in S. triqueter resulted in thinner calcareous layers in the tubes and therefore reduced fracture toughness (Díaz-Castañeda et al., 2019).

In experiments where ocean warming was also considered, increased temperature (+6°C) had the opposite effect, increasing skeletal hardness and elasticity (Chan et al., 2012). Although the temperature increase used was more extreme than predicted under future climate scenarios, the outcome for this important polychaete indicated that in a future ocean biomineral production may be more dictated by climate warming than acidification.

**Gastropods**

Depending on the species and life stage, gastropods produce aragonitic or calcitic shells or a combination of these (Marin et al., 2008; Rühl et al., 2017). The impact of OA on shell microstructure of several species, maintained in experimental OA for various lengths of time has been investigated (Table 1). For the ecologically important species, Nucella lapillus and Nassarius nitidus, the reduction in shell density determined using μCT was marked (20–50%), while that for Columbella rustica was much less (0.8–8%) (Queirós et al., 2015; Chatzinikolaou et al., 2017). This shows differences in vulnerability of shells to OA between closely related gastropod species. For the two species greatly affected, the extent of biomineral reduction varied across locations in the shell which would make weaker regions a target for derothropic predators such as crabs. In a study of the mechanical integrity of the shells of Littorina littorea maintained in pH 7.8 for 5 months, the shells of larger snails were more vulnerable to the crushing action of a crab predator than those of smaller snails (Landes and Zimmer, 2012). In a study of 7 gastropod species, the mechanical properties of the shells of six of these were not affected by acidification (pH 7.8-7.9) (Leung et al., 2017). For Austrorhodibia constricta, shell hardness and stiffness increased. This species also produced a shell with higher calcite to aragonite and magnesium to calcium ratios under acidification conditions (Leung et al., 2017). In this study elevated temperature (+2.5–4.0 °C) was the more important factor on shell mechanics. For juvenile N. lapillus, grown under OA conditions, a 2°C warming appeared to counter the negative effect of OA, but as this differed between small and large snails, the trend was difficult to interpret (Rühl et al., 2017).

Gastropods resident at vents also have thinner, less dense shells as seen in μCT reconstructions (Garilli et al., 2015; Harvey et al., 2018). The shell surface of Charonia lampas exhibited the corrosive effects of low pH water even to the extent that the soft tissue was exposed (Harvey et al., 2018). Overall, gastropods living at CO2 vent sites tend to have a smaller adult body size, similar to the Lilliput effect seen in the fossil record for mollusc species that survived past low pH driven extinction events (Garilli et al., 2015). It is suggested that smaller body size is a physiological adaptation to environmental acidification in order to maintain calcification as well as to have the energy to repair shell dissolution (Garilli et al., 2015). For herbivorous gastropods living at vent sites, the enhanced algal food levels due to high CO2 can buffer, to a variable extent, the negative effect of OA on calcification (Doubleday et al., 2019).

**Bivalves**

Bivalve skeletons are comprised of two shells joined at a hinge and vary in their composition of calcite and aragonite (Marin et al., 2008; Wehrmeister et al., 2011). There has been extensive research on the impacts of OA on these animals as they form the basis of significant aquaculture productivity. They are also ecologically important as being major prey for many species and their filter feeding activity strongly influences water clarity and quality.

Many bivalve studies report reduced shell growth, reduced shell thickness and mechanically weaker shells under OA.
conditions (Gazeau et al., 2007; Ries et al. 2009; Beniash et al., 2010; Dickinson et al., 2012; Fitzer et al., 2015a, b, 2018). Reduced shell growth and thinning of shells under experimental CO₂ acidification have been observed using SEM at both the larval (Gazeau et al., 2010; Parker et al., 2012; Fitzer et al., 2014a) and adult stages (Gazeau et al., 2013; Fitzer et al., 2014b). However, no effect on shape or size of crystals was observed in the clam Arctica islandica or the scallop Adamussium colbecki under elevated CO₂ (Stemmer et al., 2013; Dell’ Acqua et al., 2019). The abnormalities and changes to shell growth can affect bivalves at the microstructure level of the forming phases of aragonite and calcite (Wehrmeister et al., 2011), and can also impact the amorphous calcium carbonate (ACC) that is an important precursor of crystalline carbonate minerals (Addadi et al., 2003).

Microstructural observations of bivalves exposed to acidification conditions have been determined using a range of microscopy techniques including SEM coupled with electron backscatter diffraction (EBSD) (Melzner et al., 2011; Fitzer et al., 2014a, b), Fourier-transform infrared (FTIR) spectroscopy (Beniash et al., 2010) and X-ray photo emission electron microscopy (XPEEM) (Fitzer et al., 2016) to characterize crystallography and the form of calcium carbonate. µCT has been used to examine skeletal density (Meng et al., 2018).

In the mussel Mytilus edulis, which contains an outer calcite and an inner aragonite layer (Fig. 2), experimental OA resulted in a thinning of the shell (Fitzer et al., 2015b), with disordered crystallography as revealed by SEM-EBSD as well as a corroded aragonite layer (Melzner et al., 2011; Fitzer et al., 2014a), in favour of a disordered calcite layer (Fitzer et al., 2014a, b). Examination of the shells of M. edulis exposed to CO₂ acidification with XPEEM revealed that more ACC was produced (Fitzer et al., 2016). This was suggested to aid repair to the disordered calcite layer in the shell (Fitzer et al., 2016).

While M. edulis has been a focal case study species (Table 1), many bivalves are similarly affected by OA with commonly observed thinning of the shells and disorder in the crystallography of the calcite layers (Figs 2 and 3). An SEM-EBSD investigation of Mytilus galloprovincialis transplanted to a CO₂ vent (~pH = 7.2–7.8) revealed that they grew a thinner shell with disordered crystallography (Rodolfo-Metalpa et al., 2011; Hahn et al., 2012). This was also the case the shells of the oysters Magallana angulata (pH 7.2, 7.5) and M. bongkongensis (pH 7.3) grown in laboratory (CO₂ dosing) (Meng et al., 2018, 2019), and Saccostrea glomerata farmed in coastal acidified environments (pH 7.6–7.8) (Fitzer et al., 2018, 2019b). The disordered crystallography in S. glomerata wild type oysters grown in coastal acidified environments was as a result of altered biominerisation pathways shown by changing shell carbon isotopes (Fitzer et al., 2019b). Oysters such as the pearl oyster Pinctada fucata that have an internal layer of aragonite, dissolution occurred internally under OA conditions (Welladson et al., 2011).

The impacts of OA on the biomechanics of bivalve shells have been studied using a variety of techniques, in particular microindentation of the shell surface to calculate hardness. The resultant cracks propagating from the indents can also be applied to calculate fracture toughness (Mackenzie et al., 2014; Fitzer et al., 2015a). The changes to the shell microstructure in M. edulis grown in OA conditions resulted in a mechanically weaker shell as indicated by

Figure 2: A schematic representation of the use of SEM-EBSD as an analytical tool to assess the effects of acidification on shell microstructure. (A) M. edulis has been a focal study species to understand the impacts of OA on marine biomineral. (B) Section imaged using SEM is a cross section through a shell grown under OA (pH NBS 7.5). (C) The same section imaged using EBSD and analysed for crystallographic orientation displayed as a crystallographic orientation map. (D) Calcite and (E) aragonite crystals at higher magnification from the same cross section of the M. edulis shell. (F) Disordered microstructure of the calcite layer and (G) dissolution of the aragonite tablets (edges more rounded compared to panel E and tablets are less tightly packed) of the shells grown in OA. Images adapted from Fitzer et al. (2014a).
nanoindentation. The calcite produced was harder and more brittle. Microindentation indicated reduced fracture toughness (Fitzer et al., 2015a) and reduced bend/flex before failure (Mackenzie et al., 2014).

While OA (pH 7.5) as a single stressor reduced shell hardness and stiffness in *M. edulis*, when this level of acidification is used in combination with moderate warming (+2°C), these negative effects were reduced (Fitzer et al., 2015b). Temperature alone did not alter shell size of thickness; however, a combination of +2°C warming with acidification (pH 7.5) reduced the aragonite layer thickness in *M. edulis* (Fitzer et al., 2015a). A +4°C warming reduced the maximum crushing load of mussel shells in both ambient and reduced pH (pH 7.6) (Mackenzie et al., 2014).

The mechanical properties of bivalve shells that have different forms of CaCO₃ differ in response to OA. In the shell of *M. edulis*, which contains both calcite and aragonite, the calcite becomes harder or more brittle when exposed to OA. For the calcitic shell of the oyster *Crassostrea virginica* grown under experimental OA, microindentation revealed that shell hardness is reduced along with a resultant reduced fracture toughness (Beniash et al., 2010; Dickinson et al., 2012). Similarly, the shells of juvenile *Magallana angulata* grown under experimental OA (pH 7.2, 7.5) had reduced hardness, as a result of increased porosity as visualized by μCT (Meng et al., 2018). In the more resistant *M. bongkongensis*, juvenile shells were similarly affected, but at a much lower pH (pH 7.3) (Meng et al., 2019). For this species, shell hardness and crystallography remained unchanged at pH 7.6 (Meng et al., 2019). Oysters such as *Pinctada fucata* that have an internal layer of aragonite had a weaker shell when grown under experimental OA conditions (Welladsen et al., 2011) than oysters that have a calcitic internal layer. The shells of *S. glomerata* growing in estuarine habitats with sulphate soil acidification (pH 6.6, 6.8, 6.9) were also significantly weaker in crushing tests (Amaral et al., 2012). Although these pH levels are much lower than predicted OA, this level of acidification occurs in acid sulphate regions during increased rainfall (Amaral et al., 2012). Reduced growth of *S. glomerata* during such extreme events may be offset by positive growth in dry periods (Amaral et al., 2012).

Overall, OA affects the microstructure of bivalve shells through reduced shell growth with thinner, more porous shells that have reduced fracture toughness. Trends appear similar in mussels and oysters in terms of microstructural responses to OA regardless of whether laboratory CO₂ acidification or environmental acidification from CO₂ or coastal run off are the source of acidification (Table 1).

**Echinoderms**

Echinoderms produce a unique endoskeleton laid down as a three-dimensional mesh-like calcite lattice (Fig. 4), with cells and connective tissue living in the void space (Cavey and Markel, 1994). The impacts of OA on the production and maintenance of the skeleton are extensively investigated for echinoids (sea urchins) across their planktonic and benthic life phases (Byrne et al., 2013; Dubois, 2014). These animals are ecologically and economically important grazers across world oceans (Lawrence, 2013). The sea urchin skeleton is composed of Mg-calcite, ~3–16 wt% MgCO₃, where Mg²⁺ is substituted for Ca²⁺ during calcification (Chave, 1954; Smith et al., 2016). As a result of this chemical composition, the echinoderm skeleton is one of the most soluble forms of CaCO₃, and so is vulnerable to dissolution in OA conditions (Andersson et al., 2008; McClintock et al., 2011; Dubois, 2014).
Despite the echinoderm skeleton being a form that is vulnerable to dissolution, the epithelium largely protects the living skeleton from direct exposure to low pH conditions (Dubois, 2014). The exceptions are the tips of the spines, potentially due to their thin epithelial cover. Overall, the established sea urchin skeleton does not appear to be vulnerable to direct dissolution under OA conditions (Table 1). Surface pitting of the test has only been reported at very low pH (Holtmann et al., 2013). Interestingly, the cidaroids, a group of sea urchins with many species living in deep water below the isocline, produce spines that do not have an epithelial cover, but have a calcitic cortical layer that makes them resistant to dissolution in OA conditions, as shown for 15 species, including in tests where species were maintained in pH 7.2–7.4 for 3–5 weeks (Dery et al., 2014, 2017, 2018).

With respect to production of biomaterial, calcification in sea urchins is constrained by OA. This is shown by the comparatively smaller skeletons of larvae and juveniles (Figs 5 and 6) reared in OA conditions through development and the smaller tests of adult sea urchins (Fig. 7) grown in long-term OA experiments (Byrne et al., 2013, 2014; Dubois, 2014; Dworjanyn and Byrne, 2018). It appears that sea urchins have a reduced ability to deposit biomineral under OA conditions (Figs 4 and 5). This may be due to energetic constraints associated with the higher metabolic costs of life at low pH and the priority to maintain acid-base balance (Stumpp et al., 2012; Carey et al., 2016). The impacts of OA on the skeleton may be mitigated by near future warming. In Tripneustes gratilla, a +3°C warming mitigated the negative effects of low pH (pH 7.6, 7.8) on test growth (Fig. 7). Further warming made matters worse (Byrne et al., 2014; Dworjanyn and Byrne, 2018).

In addition to the smaller amount of skeleton produced under OA conditions, skeletal microstructure may also be altered. Several SEM and/or μCT investigations have compared the pore size, the calcitic trabecular connections and the void space of the skeleton of sea urchins exposed to OA with these traits in the skeleton formed under ambient pH (Table 1). Larger pore size is reported for the spines of juvenile Heliocidaris erythrogramma (Fig. 6), but only at very low pH (pH 7.4) (Wolfe et al., 2013b) and for the teeth of Paracentrotus lividus where the tips were regenerated under OA (pH 7.7) (Asnaghi et al., 2013).

For H. erythrogramma maintained on OA conditions for 9 months, the mid body ambital test plates produced prior to exposure to OA exhibited no change in skeletal structure. In contrast, the younger apical plates from the active growth region of the test, that had likely grown over the 9-month exposure at low pH had thinner trabeculae and a greater void space compared to those from control urchins as visualized with SEM (Fig. 4A and B) and μCT (Johnson, 2019).

The μCT study of Strongylocentrotus fragilis that had likely grown from the juvenile stage in chronic low pH (pH 7.69–7.57) and oxygen saturation on the Californian continental shelf, showed that that skeletal porosity and pore size is higher than that for conspecifics living at higher pH. For this species, skeletal porosity increased with decreasing pH and oxygen saturation, but it is not known how hypoxia and low pH interact with respect to the differences in the skeleton (Sato et al., 2018).

The impact of OA on the mechanical properties of the test plates has been documented by nanoindentation and in
crushing and bending tests (Table 1). In *P. lividus* maintained in OA (pH 7.8, 7.9) for 12 months, there was no effect of pH on the hardness or breaking force of the test plates (Collard et al., 2016), as also the case for *Echinometra mathaei* (Moulin et al., 2014). Exposure to low pH reduced the hardness of the apical plates of *H. erythrogramma* that had likely been produced in OA conditions, but the previously established ambital plates were not affected (Johnson and Byrne, unpublished data). There was also no change in the mechanical properties of the test of *P. lividus* resident at a vent site (pH 7.7) compared with conspecifics from ambient conditions.

In studies where the whole test was crushed as might occur in an attack by a predatory fish (Guidetti and Mori, 2005), the force required to crush the dried tests of juvenile *P. lividus* and *Diadema africanum* maintained in OA was lower compared to juveniles from control treatments (Asnaghi et al., 2013; Rodriguez et al. 2017). However, as dried tests were used, the ecological relevance of these results is not clear. The force required to crush the test was lower in live *Tripneustes ventricosus* reared in OA (pH 7.6) from juvenile to adult in a test designed to mimic the ram force of a predatory fish, but the break occurred at the suture lines, not in the skeleton itself (Byrne et al., 2014). These tests need to be repeated across species with live urchins being most relevant test subjects, to incorporate the elasticity of the ligaments that bind the test plates (Ellers et al., 1998), and with application of forces modelled with respect to the jaw crushing or ramming action of predators, although these forces are poorly characterized.

With respect to the spines of sea urchins maintained at pH 7.4 and 7.7, the thinner, more brittle spines formed in these conditions by *Tripneustes ventricosus* and *Eucidaris*...
**A. improvises** shell was weaker (McDonald et al., 2009) through the larval stage, but the adult (+3°C) warming mitigated the negative effects of low pH (pH 7.6, 7.8), but further warming was deleterious. From Dworjanyn and Byrne (2018).

**Tripneustes gratilla** reared in three pH and three temperature levels in all combinations from the early juvenile (5.0 mm test diameter) for 146 days. A +3°C warming mitigated the negative effects of low pHNBS (pH 7.6, 7.8), but further warming was deleterious. From Dworjanyn and Byrne (2018). **Figure 7:**

**tribuloides** are weaker in bending tests, but not the spines of *Prionocidaris baculosa* (Dery et al., 2017). While the spines are the most vulnerable skeletal element in sea urchins to OA, these structures are designed to break in a predatory attack, and regenerate quickly (Dery et al., 2017). But this entails an energetic cost (Haga et al., 2016).

Overall, OA affects the material properties of the sea urchin skeleton through changes in growth rate with less biomineral produced. While it appears that there is weakening of the sea urchin skeleton in response to OA, thereby compromising its protective roles, more studies are needed where characterisation of the impacts of OA on biomineral microstructure is investigated in tandem with biomechanical tests and in skeletal elements that have been completely produced at low pH. Finally, the 3D void space of the sea urchin endoskeleton is important for the resident cells and tissues. Any change in the stereotypic arrangement of the skeleton is likely to have impacts for organism function beyond the skeleton itself.

**Crustaceans**

The crustacean exoskeleton is made of chitin and calcite and is comparatively tolerant of acidification due to the lower degree of calcification (Whiteley, 2011), the exception being adult barnacles, with several studies on these animals (Table 1). For *Amphibalanus amphitrite*, there were no effects of low pH (to pH 7.4) through the larval stage, but the adult shell was weaker (McDonald et al., 2009). Corrosion of the shell of *A. improvises* was evident at low pH (Pansch et al., 2014). Increased temperature (+4°C) resulted in an increase in shell strength of this species, but there was no effect of pH (Pansch et al. 2013, 2014). For *Balanus improvises*, low pH (stable pH 7.7, fluctuating pH 7.5–7.9) reduced the strength of the shell (Eriander et al., 2016).

**Discussion**

The morphological and biomechanical properties of marine invertebrate skeletons are integral to animal function and individual fitness which in turn affects population dynamics and community structure (Kroeker et al., 2014). In a climate change world, it is important to understand how calcification systems, and the biomineral that is produced, respond to environmental acidification. The combination of detailed morphology using advanced imaging techniques and biomechanics, is emerging as a leading-edge approach to quantify the vulnerability of marine skeletons. This approach is also key to detecting the potential for phenotypic adjustment in biomineralisation as a means to produce and maintain biomineral in the face of environmental acidification.

Although the field of marine climate change has focused on anthropogenic CO2-driven acidification, many important calcifying species live in shallow coastal habitats that are vulnerable to acidification from increasing run off due to pressures from sea level rise and precipitation (Amaral et al., 2011; Fitzer et al., 2018). The chemistry of ‘ocean’ and ‘coastal’ acidification, and how they impact the carbonate system, the basis for calcification, are fundamentally different (Fig. 1). For bivalves, despite these differences, the resulting morphology and biomechanics of the biomineral produced under the two forms of acidification are similar (Beniash et al., 2010; Rodolfo-Metalpa et al., 2011; Hahn et al., 2012; Fitzer et al., 2018; Meng et al., 2018). Coastal waters have a more variable chemistry than the ocean and understanding how this chemistry interacts with CO2-driven acidification, remains a significant challenge. This is especially important for socioeconomically important molluscs resources that are typically fished and cultured in coastal bays and estuaries.

In response to environmental acidification, the biomineral produced by marine invertebrates across many taxa, is more porous and less dense. Increased skeletal porosity in diverse species has been revealed by SEM and/or high-resolution 3D reconstructions generated by μCT. Indeed, some skeletal malformations are only evident through application of these methods (Fitzer et al., 2014a, b, 2015a, b, 2018; Rühl et al., 2017). There may be a balance between the absolute amount of mineral that can be physiologically deposited in the skeleton in order to maintain its key physiological functions and the energy that can be devoted to this function under OA. For molluscs and echinoderms, constraints in biomineral production result in smaller body size, similar to the Lilliput effect seen in the fossil record when smaller mollusc species survived global warming extinction events (Garilli et al., 2015). In contrast for corals, growth, as measured...
by linear extension of the skeleton, the standard metric of coral health appears to be maintained under CO2-driven acidification, but the skeleton produced has a lower density and is thus more vulnerable to physical damage (Tambutté et al., 2015). OA also impacts biomineral directly through dissolution and is this most commonly noted for molluscs that lack an outer protective cover of concholin (Ries et al., 2009; Rodolfo-Metalpa et al., 2011; Harvey et al., 2018). Echinoderm skeletons are protected from corrosion by their epithelial cover, and in corals, the polyp tissue protects the skeleton.

Within each of the groups considered here, environmental acidification impacts skeletal microstructure and biomechanics in different ways, reflecting the diverse calcification biology of the species investigated. In bivalves, where skeletal CaCO3 varies in composition of calcite and aragonite, the responses to CO2-driven acidification are often similar within the same family. For example, the mytelid (Mytilus sp) have harder, brittle shells prone to fracture under OA, compared with the Ostreidae (Crassostrea sp.) where the shells have reduced hardness. Despite these different responses, the outcome with regard to the mechanical integrity of the shell of mussels and oysters is similar with reduced fracture toughness of shells formed in OA conditions.

Volcanic CO2 vent systems have been used as natural laboratories to investigate the impacts of near and far future OA (Foo et al., 2018; González-Delago and Hernández, 2018). The CO2 gradient at these sites provides a pH range from very low ∼pH 6 to ambient levels in nearby habitats not affected by CO2 venting. Investigation of marine calcifiers resident along these CO2 gradients where all or most of their skeleton has formed under low pH, has been an important means to understand how acidification impacts calcification. Overall, the results of laboratory and vent studies, at similar pH/acidification levels, are similar with respect to the response of marine invertebrate calcification. For herbivorous species resident at vents, higher levels of algal resources promoted by the CO2 helps to ameliorate the energetic stress caused by acidification (Uthicke et al., 2016).

Biomechanical tests of the biomineral produced under experimental and natural acidification indicate that the shells and skeletons of many species are weaker than those produced in ambient conditions. This has been largely shown for mussels with some studies of tube worms and corals, although the shells of many gastropod species are not affected by acidification (Leung et al., 2017). For echinoderms, biomechanics tests indicated that the spines are most affected by acidification (Collard et al., 2016; Dery et al., 2017) and that the newly formed apical plates were weaker (Johnson and Byrne, unpublished data). Overall, the shift to producing weaker skeletons with increased porosity decreased density under acidification conditions in many species will compromise the function of these structures in protection against physical stress and defence against predators.

It is key to consider the fitness consequences of observed change. Change to animal size, strength or defence will have effects to marine communities beyond the fitness of the individual. Changes to predator–prey interactions may change predation rates of vulnerable species and thereby alter population dynamics and community structure in the future (Kroeker et al., 2014). Weaker shells change predator–prey dynamics as durophagous predators such as crabs optimize their foraging to preferentially target weaker-shelled individuals or weaker regions of the shell of their molluscan prey (Kroeker et al., 2014). Predatory mollusc and boring琵bions also target the weaker, less dense areas of shells to drill and bore through the biomineral. Many molluscs also exhibit an inducible defence behaviour producing stronger, more calcified shells in response to the presence of a predator or in response to hydrodynamic conditions and the latter also applies to sea urchins (Bibby et al., 2007; Collard et al., 2016). For the gastropod Littorina obtusata, the inducible defence response expressed in the presence of a crab predator was inhibited at low pH, albeit at a rather extreme level (pH 6.6, ~14 000 μatm pCO2) (Bibby et al., 2007). The thinner shells of the oyster Ostrea lurida reared from the larval stage in OA appears to have made them more vulnerable to predatory gastropods (Sanford et al., 2014). In a study of M. edulis and the drilling gastropod Nucella lapillus maintained in the same acidification conditions, the thinner mussel shells produced in OA conditions made them more vulnerable to predation by the gastropod (Sadler et al., 2018). The outcomes for marine calcifiers in the face of OA will also be influenced by other environmental factors such as food levels. Mussels and barnacles are able to maintain a higher-level calcification in the presence of high food levels (Melzner et al., 2011; Pansch et al., 2014). Sea urchins with a higher level of calcified algae in their diet produce stronger skeletons (Asanaghi et al., 2013). Species adapted to fluctuations in intertidal habitats are also more tolerant of acidification (Wolfe et al., 2013b; Leung et al., 2017).

For several marine invertebrate groups across a variety of biomineralized structures, the lack of studies of the microstructure and mechanics stands out as gaps to address. In particular, the response of the crustacean chitin-calcite exoskeleton to acidification is poorly studied. The impact of acidification on jellyfish calcium sulfate (basanite) statoliths (Mooney and Kingsford, 2016; Sötje et al., 2017) is not known. Understanding the effects of acidification on the organic matrix of marine skeletons is also an important gap in knowledge, especially as shell matrix proteins are essential for crystal nucleation and other key aspects of skeletogenesis (Marin et al., 2008).

Finally, while the integrated morphology-mechanics approach has great potential to characterize the impacts of environmental acidification on marine invertebrate skeletons and advance our understanding of the biological consequences of climate change, this approach has only been applied to a relatively small number of species (Table 1). There is great interest in how calcification and the biomineral
produced will be altered in a changing ocean and it is clear that application of morphology and ecomechanics, in context with use of realistic pH levels, will be particularly informative.

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