

### The effect of environmental factors on shell growth and repair in Buccinum undatum

Colvin, Charlotte; Hollyman, Philip; Richardson, Christopher; Chenery, Simon; Waggitt, James; McCarthy, Ian

### Journal of Experimental Marine Biology and Ecology

DOI: 10.1016/j.jembe.2022.151720

Published: 01/06/2022

Peer reviewed version

Cyswllt i'r cyhoeddiad / Link to publication

Dyfyniad o'r fersiwn a gyhoeddwyd / Citation for published version (APA): Colvin, C., Hollyman, P., Richardson, C., Chenery, S., Waggitt, J., & McCarthy, I. (2022). The effect of environmental factors on shell growth and repair in Buccinum undatum. *Journal of Experimental Marine Biology and Ecology, 551*, [151720]. https://doi.org/10.1016/j.jembe.2022.151720

Hawliau Cyffredinol / General rights Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

• Users may download and print one copy of any publication from the public portal for the purpose of private study or research.

- You may not further distribute the material or use it for any profit-making activity or commercial gain
   You may freely distribute the URL identifying the publication in the public portal ?

Take down policy If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

1 The effect of environmental factors on shell growth and repair in *Buccinum undatum* 

- 3  $^{1}$ School of Ocean Sciences, College of Natural Sciences, Bangor University, Menai Bridge, Anglesey LL59 5AB, UK
- 4 <sup>2</sup>British Antarctic Survey, High Cross, Madingley Road, Cambridge, CB3 0ET, UK
- <sup>3</sup>Centre for Environmental Geochemistry, British Geological Survey, Keyworth, Nottingham NG12 5GG, UK
- 6 \*Corresponding author c.n.colvin@bangor.ac.uk (C.N. Colvin)
- 7
- 8 Abstract

9 The processes and factors which affect shell growth and repair in molluscs are poorly 10 understood. In this study, the capabilities of shell growth and repair in the marine gastropod 11 Buccinum undatum were investigated experimentally by implementing laboratory-controlled 12 mechanical damage to the shell margin/lip. Three key factors, life stage (juvenile or adult), 13 seawater temperature (5-15°C) and food availability (unfed, weekly, or daily feeding), were 14 investigated in a series of controlled laboratory experiments to establish their roles in the 15 processes of shell growth and repair. Significant differences in rates of shell growth and repair 16 between food and temperature regimes were observed, with the greatest difference occurring 17 with different life stages. Rates of shell growth in non-damaged whelks were slightly faster but 18 not significantly different from damaged individuals in any of the experiments. Tank-reared 19 juveniles maintained in the highest seawater temperature regime (15°C) displayed significantly 20 faster rates of shell repair (F=6.47, p<0.05) than conspecifics held at lower seawater 21 temperatures. Through characterising both biological and environmental factors affecting shell 22 growth and repair, it is demonstrated that there are multiple aspects influencing shell growth 23 and shell repair. It is important to be able to understand and establish differences in rates of 24 growth to better manage this commercial species.

25 Keywords

26 Gastropod mollusc; shell growth; *Buccinum undatum;* fishery; shell damage; repair

<sup>2</sup> C.N. Colvin<sup>1\*</sup>, P.R. Hollyman<sup>2</sup>, C.A. Richardson<sup>1</sup>, S.R.N. Chenery<sup>3</sup>, J.J. Waggitt<sup>1</sup>, I.D. McCarthy<sup>1</sup>

28 1. Introduction

29 Growth is a fundamental process for determining survival and reproductive success within all 30 taxa and is critical for population growth and resilience, and therefore, understanding what 31 impacts measurable growth is of key importance (Sebens, 1987; Pepin, 1991). Within living 32 organisms, growth is primarily defined as a measurable increase in length or mass (e.g., von 33 Bertalanffy, 1938; France et al., 1996; Ohnishi and Akamine, 2006). For some species, e.g., the 34 common whelk, Buccinum undatum, somatic and shell growth is continuous throughout life 35 (Kideys et al., 1993). As such, B. undatum displays a predictable relationship between size and 36 age, which has been used to determine growth rates in several studies (Hollyman et al., 2018; 37 Emmerson et al., 2020; Borsetti et al., 2021).

38 The molluscan shell is vitally important, providing protection and additional support to the 39 internal body in the form of a calcareous exoskeleton (Ruppert et al., 2004; Bonucci, 2007). As 40 such, the shell is required to increase as the internal soft tissue grows. The physical process of 41 shell growth occurs as a result of CaCO<sub>3</sub> secretions from the mantle epidermis (Findlay et al., 42 2011). Damage to the shell can lead to interruptions in growth, with incidents often recorded 43 in the structure of the shell in the form of shell scarring (Richardson, 2001; Ramsay et al., 2001; 44 Preston and Roberts, 2007), although there is a paucity of literature surrounding the effects of 45 shell damage and rates of regrowth in gastropod species. In addition to the normal process of 46 shell growth with age, it is necessary for gastropods to be able to quickly repair their shells 47 following predation attacks and to protect them from their local environment and reduce 48 further predation when vulnerable soft tissues are left exposed following damage. During 49 periods following damage shell growth can heavily impact on metabolic functioning of the

50 organism, as energy resources are redirected into shell repair (Ebert, 1968; Ruppert et al., 2004;

3

51 Melzner et al., 2011; Thomsen et al., 2013).

52 There are a range of factors that inflict damage to the shells of marine gastropods, including 53 storms and increases in turbulence and exposure (Sepùlveda et al., 2012; Solas et al., 2015). 54 For *B. undatum*, damage to the shell margin often occurs following unsuccessful predatory 55 attacks, both against and inflicted by whelks themselves (Checa, 1993; Mensink et al., 2000; 56 Ramsay et al., 2001). However, one of the key factors leading to shell damage is that caused 57 from fishing gear, both directly from impact with mobile towed fishing gear and indirectly 58 following capture. For example, the process by which whelk are graded for size on a ship's deck 59 can pose additional risk to the shells of under-sized specimens due to the riddling process. 60 Captured whelks are passed through a series of metal riddle bars which sort the catch and 61 retain whelks greater than the minimum legal landing size but allow undersized whelks passage 62 through where the shells may become broken or chipped (Bergman et al., 1994; Cadée et al., 63 1995; Fahy et al., 1995; De Vooys et al., 1998; Mensink et al., 2000).

64 This study investigates the impact of environmental factors, seawater temperature, food 65 availability and ontogeny on shell growth and repair in the common neogastropod whelk 66 Buccinum undatum. This whelk occurs in British coastal waters and has a widespread distribution throughout North Atlantic shelf waters with records from the Greenland Seas in 67 68 the north to the Bay of Biscay in the south (Golikov, 1968; De Vooys and van der Meer, 2010; 69 Hayward and Ryland, 2011; Magnúsdóttir et al., 2019). They are the largest edible marine 70 gastropod within the North Atlantic (Fahy et al., 2005) and are of commercial importance. 71 Between January and March 2020, >4000 tonnes were landed by U.K. vessels into U.K. ports 72 (price >£1,200 per tonne) (MMO, 2020). The U.K. shellfish industry has long been of importance 73 economically and historically has relied on sustainable whelk populations that are resilient to

environmental change and harvesting pressures (Clark et al., 2016). Any disruption to an organism's growth through repeated shell damage and repair could potentially result in impacts on the fishery (Hilborn and Minte-Vera, 2008; Biro and Sampson, 2015), for example through reduced growth and longer or failure to reach the minimum landing size (currently in the U.K. MLS = 45 mm McIntyre et al., 2015). Despite the increasing fishery demand for *B. undatum* and need for sustainable management strategies (Fahy et al., 2000), shell damage and repair are understudied and ecologically-important aspects of research.

The aims of this study are to understand both the environmental and biological factors that can affect shell growth rate and repair in *B. undatum*. It is hypothesised that during shell repair, shell growth rate (increase in length) is reduced following a re-direction of energy away from size increase towards shell repair and that this process will be affected by seawater temperature and food availability for whelk of different age (size) groups.

86 2. Materials and Methods

#### 87 2.1 Sample collection and holding

88 Newly-laid egg masses of *Buccinum undatum* attached to the pier pilings of a small jetty and 89 surrounding rocks at Brynsiencyn, North Wales (UK) 53°09′30.4″N 4°16′46.6″W, were collected 90 during low water of spring tides (November 2014 and November 2017). Large (>45 mm total 91 shell length (TSL), assumed to be adult) and small (<45 mm TSL, assumed to be juvenile) whelks 92 were collected (January & July 2019) using inkwell pots baited with Atlantic mackerel (Scomber 93 scombrus), deployed for 24h subtidally at 4m depth off Brynsiencyn. Whelk maturity was 94 probabilistically determined using a previously calculated size of maturity (SOM) estimation to 95 group specimens as either adult whelk (AW) or juvenile wild (JW) (see Haig et al., 2015). This 96 was achieved using the European MLS of 45 mm as opposed to larger, site-specific MLS outlined

97 in Haig et al. (2015). This smaller size was used to ensure juveniles were juveniles, as a larger 98 TSL could incorporate more mature individuals. Egg masses and whelks were maintained in 99 laboratory tanks, supplied with flowing ambient temperature seawater (8-10°C), and aerated. 100 Ten egg masses were placed in each of three 50L tanks with fine mesh over the outflow pipe 101 to retain any emerging juveniles (tank-reared juvenile whelk (TRJ)). After approximately one-102 month, juvenile whelks hatched and were transferred daily to smaller 8L tanks (mesh over 103 outflow) supplied with ambient temperature flowing seawater, where they were on-grown and 104 fed daily on a diet of freshly-shucked mussel (Mytilus edulis) tissue.

105 2.2 Experiment set up

106 A daily frequency of feeding continued throughout the experiments, unless stated otherwise 107 (see Table 1), to ensure food availability was not a limiting factor for shell repair and shell 108 growth. After hatching and collection, whelks were on-grown at a range of seawater 109 temperatures (5, 10 and 15°C) and feeding regimes (unfed, daily fed or weekly fed) (see Table1) 110 and following ten days acclimation, shell growth and repair were investigated under these 111 different environmental conditions. For all experiments carried out under ambient conditions, 112 wild-caught whelks were maintained in a flow-through system in 15L tanks, with TRJ whelk 113 suspended within these in 8L tanks with mesh inserts to prevent escape. For temperature 114 experiments, larger 175L temperature-controlled tanks were used for wild-caught whelk with 115 TRJ again suspended in mesh containers to allow water flow. To achieve acclimation at the 116 upper or lower temperatures, water temperature was raised or lowered incrementally before 117 experimental shell damage was undertaken ten days later. Each whelk was individually labelled 118 with a waterproof paper number adhered with superglue (see Figure 1) so that daily changes 119 in individual shell growth rates could be followed. Although experiments were run over several 120 different time periods (see Table 1), the size and age of tank-reared juvenile whelks were kept

121 consistent between years, along with all the environmental factors.

122 2.3 Controlled damage and measurements

123 Following several iterations of method development, the shells of the TRJ, JW and AW whelks 124 were experimentally damaged by cutting out a square portion of the shell lip of each whelk 125 using a Dremel 3000, with a cutting wheel attachment. Each cut was 40% of the aperture 126 diameter (see Figure 1). A proportion of whelks within each experiment were left undamaged 127 to serve as a control comparison. However, for experiments where food and seawater 128 temperature were controlled, due to a limited number of individuals, TRJ did not have un-129 damaged controls. Photographs of each damaged and undamaged whelk shell were taken 130 against a measurable scale and the area of shell removed and regrown and total shell length 131 (TSL) determined (to 3 d.p.) using ImageJ (see Figure 1). Measures of TSL were recorded to 132 monitor growth, not shell repair. The frequency of measurements depended on the rate of 133 visible shell growth and repair and was consequently different between different age groups. 134 Measurements were taken every 3 days for (TRJ), 6 days (JW) and 7 days (AW) respectively until 135 the end of the experiment period to limit disturbance during growth.

136 2.4 Statistical analysis

From the measurements of TSL and damage area, an individual cumulative value of TSL increase and percentage repair was calculated for the experiment period and daily rates calculated from these final values as percentage repair and growth per day (mm<sup>2</sup>.day<sup>-1</sup> and mm.day<sup>-1</sup> respectively). General Linear Models (GLM) were used to test whether growth and shell repair were significantly influenced by environmental variables. Depending on experimental treatment (Table 1), response variables were either percentage repair (%) or TSL growth day<sup>-1</sup> 143 (mm); explanatory variables were either age (TRJ, JW, AW), food (unfed, daily, or weekly) or 144 temperature (5, 10 or 15°C). Response variables were modelled as continuous variables, and 145 explanatory variables were modelled as categorical variables. Interactions between life stage 146 and food or temperature were included to test whether relationships between growth rates or 147 percentage repair and food/temperature varied amongst age-groups (TRJ, JW, AW). Statistical 148 significance (p-values) were obtained from F-tests, and backwards model selection based on p-149 values was applied (Zuur et al., 2009). When interaction terms were non-significant in the full 150 model, they were replaced with a non-interactive term and the process restarted. Diagnostic 151 plots of residuals were checked. For graphical representation, the cumulative values were 152 plotted in place of actual rates for a clearer comparative view of growth and shell repair 153 between treatment groups. TSL was presented graphically using an average cumulative growth, 154 taken as the average increase in shell length, and percentage repair as the average proportion 155 of shell repaired out of 100% for all individuals per measured day.

Due to differences in experiment length, with adult whelk studies run for longer periods than juveniles, and disparity in the degree of shell repair recorded, with some whelks attaining 100% repaired shell long before the end of the experiment, the values have not been taken as a final day measure. To resolve this, for statistical purposes, experiment duration for measures of repair rate have been taken as the day at which the first 75% of individuals from one single experiment have repaired to 100%.



162

Figure 1: Experimentally damaged TRJ *Buccinum undatum* (number 38) showing the area of shell removed at the shell lip at day 0 and then repaired by day 6 (black dashed lines). Percentage repair area measurement calculated through the remaining area to be deposited until a value of 0 was reached. Total shell length (TSL) is indicated by the yellow dashed line.

#### 167 2.5 Calcein staining

168 To investigate qualitative changes in shell structure during shell repair, whelks were marked 169 with the fluorescent dye Calcein to distinguish the original shell growth prior to damage 170 (Kaehler and McQuaid, 1999). Twenty whelks whose shell margins (lips) had been mechanically 171 damaged were stained in a 50 mgL<sup>-1</sup> Calcein-seawater solution for 24 hours, to mark the point 172 in growth when the shell was damaged. The whelks were then on-grown and fed for 7 days in 173 flowing ambient temperature seawater after which the edge of the shell lips of ten of the shells 174 that had repaired 50% of their shell were again damaged for a second time and then on-grown 175 for a further 7 days. At the end of the experiment, whelks were frozen at -20°C as a means of 176 dispatch and upon thawing, the flesh was removed avoiding damage to the shell margin. 177 Organic material was removed from the shell surface by immersion for 30 mins in a 0.01M 178 NaOH solution, rinsed in fresh water and air-dried before the complete shell was embedded in 179 Kleer-set<sup>™</sup> polyester casting resin (see Hollyman et al., 2020). The embedded shell lip and area 180 of shell damage were sectioned using a precision diamond saw blade (Beuhler isomet 4000) 181 and the cut surface ground on successively finer silicon carbide abrasive papers (FEPA P400 and 182 P1200 grade) before attaching the dry, cut surface to a microscope slide using superglue. Once 183 dry, the attached resin and shell were sectioned again using the saw to create a 1 mm thin 184 section. The thin section was ground with successively finer grades of silicon carbide paper and 185 polished to a final grade using a 3  $\mu$ m diamond suspension to achieve an approximate 0.2 mm 186 thick polished section. Low power (10x magnification) images of the shell structure were taken 187 under transmitted light, using a compound microscope. Matching (10x magnification) images 188 of the fluorescing Calcein marks were taken using a UV fluorescent light microscope and 189 photographed using an attached eyepiece camera. This produced comparative composite 190 images of the repaired shell and structure and Calcein marks.

191

# 192 Table 1: Summary of experimental design with key differences and experimental conditions outlined

Treatment	Timeframe	Experiment Length (days)	Life Stage	Tank arrangement	Food Availability	Seawater Temperature	
Life stage		39	Tank Reared Juvenile (TRJ)	8L within 15L tank 20 per tank	Daily		
	January 2019	48	Juvenile Wild (JW)	15L tank 20 per tank	Daily	_	
		70	Adult Wild (AW)	15L tank 10 per Daily tank		— Ambient (8-10°C)	
Food availability	July 2019	15	Tank Reared Juvenile (TRJ)	8L within 15L tank 15 per tank	Unfed, daily, once weekly	-	
		21	Adult Wild (AW)	15L tank 10 per tank	Unfed, daily, once weekly	_	
Temperature	January 2016	15	Tank Reared Juvenile (TRJ)	8L within 175L tank 20 per tank	Daily	5, 10 and 15°C	
	January 2019	21	Adult Wild (AW)	175L tank 40 per tank	Daily	5 and 10°C	

195 3.1 Effect of age on shell repair and growth.

196 The effect of life stage on shell repair rate is shown in Figure 2A. Mean repair rate with time 197 decreased with increase in life stage, with an overall range in rates of 2.1% day<sup>-1</sup> between the 198 three measured groups (see Table 2 for rates). Laboratory tank-reared whelks (TRJ) (TSL 11.5-199 21.25mm) repaired their shells rapidly and all shells were repaired within 30 days. Juvenile wild-200 caught (JW) (TSL <45mm) and adult wild-caught (AW) (TSL >45mm) whelks repaired their shells 201 more slowly so that by day 40 all the JW whelks had repaired their shells but none of the AW 202 whelks had completely repaired their shells by day 65. The cumulative increase in total shell 203 length with time (Figure 2B) for the three different whelk age groups showed a similar trend to 204 repair rate. Repair rate and increase in total shell length were significantly different between 205 age groups (both p< 0.001; Table 2). Compared with the experimentally-damaged whelks, the 206 undamaged control whelks generally achieved a larger total length (Figure 2B), although this 207 was not significantly different (p>0.05; Table 2).



Figure 2: *Buccinum undatum*: A) average cumulative % shell repair (± 1S.E. ribbon) with time
and B) average cumulative increase in total shell length growth (± 1S.E. ribbon) with time for
three different age groups during a 70-day (AW – adult wild-collected), 48-day (JW – juvenile
wild-collected) and 39-day (TRJ – tank-reared juvenile) laboratory experiment.

213 3.2 Effect of food ration on shell repair and growth.

Damaged juvenile (TRJ) whelks repaired their shells significantly faster than AW whelks 214 215 (p<0.00.1 Table 2) but there were no significant effects of ration or the interaction between 216 age and ration on repair rate (both p>0.05; Table 2). A plot of cumulative repair rate (Figure 217 3A) showed juveniles that had no food or were fed daily had a faster repair rate than those fed 218 once a week. AW whelks fed once a week repaired their shells at a similar rate to TRW 219 individuals fed a similar ration. However, those AW whelks fed daily or unfed repaired their 220 shells more slowly (Figure 3A). When the effects of ration on cumulative total shell length 221 (Figure 3B) were investigated statistically, TRJ whelks grew significantly faster (p<0.001; Table 222 2) than damaged and control AW whelks (Figure 3B). Generally, the AW control whelks grew 223 faster than the damaged AW whelks, although the exceptions were the AW control whelks and 224 AW whelks fed daily which grew the slowest, 0.028 mm.day<sup>-1</sup> and 0.027 mm.day<sup>-1</sup> respectively 225 (Table 2 and Figure 3B). Whilst shell growth rate was depressed in AW damaged and control 226 whelks, growth rate was promoted in whelks fed once a week, particularly in TRJ whelks (Figure 227 3B). The effect of ration and the interaction between ration and age were both significant 228 (p<0.001 and p<0.05 respectively; Table 2).



Figure 3: *Buccinum undatum* grown during a 21-day (AW) and 15-day (TRJ) laboratory experiment with different feeding regimes. A) average cumulative % shell repair (± 1S.E. ribbon) with time for two different groups, tank-reared juveniles (TRJ) and wild-collected adult whelks (AW) provided with three different rations (unfed, daily, weekly) and B) average cumulative increase in total shell length growth (± 1S.E. ribbon) with time for three different groups, tankreared juveniles (TRJ), wild-collected adult whelks (AW) and undamaged wild-collected adult whelks (control) (AW control) provided with three different rations (unfed, daily, weekly).

237 3.3 Effect of Temperature on shell repair and growth.

229

Experimentally-damaged juvenile whelks (TRJ) repaired their shells faster than adult whelks (AW) at all three seawater (SW) temperatures (p<0.001; Table 2; Figure 4A). TRJ whelks maintained at 15°C repaired their shells at an average of 10.4% day<sup>-1</sup> with >95% of shells being repaired by day 9. A faster initial increase in repair rate (between days 0 and 12) was observed for TRJ whelks maintained at 5°C compared to TRJ whelks maintained at 10°C. At 15°C juveniles repaired their shells completely in 14 days with a repair rate more than double the rate of TRJ whelks held at 10°C and 5°C which repaired 70% of their shell over the same time-period (Figure 245 5A). Adult whelks reared at 10°C and 5°C repaired 75% and 50% of their shell respectively by 246 day 20 (Figure 4A). The effects of SW temperature and the interaction between SW 247 temperature and age were both significant (p<0.001 and p<0.05 respectively; Table 2). When 248 the effect of SW temperature on cumulative total shell length (Figure 4B) was investigated 249 statistically (Table 2), TSL increased in TRJ whelks independent of SW temperature (Figure 4B). 250 In AW whelks TSL generally increased independent of SW temperature for the first 5 days 251 whereupon TSL increase slowed down (Figure 4B). Adult whelks increased in TSL the fastest at 252 10°C (AW and AW control) whereas TSL increase in adult whelks was slow at 5°C (Figure 4B). 253 The effect of SW temperature on TSL was significant (F=3.41, p<0.05) but the interaction 254 between SW temperature and age was non-significant (p>0.05; Table 2).



Figure 4: Shell repair in *Buccinum undatum* grown during a 21-day (AW), 15-day (TRJ) laboratory experiment at different seawater temperatures. A) Average cumulative % shell repair (± 1S.E. ribbon) with time for juvenile (TRJ) and adult (AW) whelks grown at 5, 10 and 15°C (TRJ) and 5 and 10°C (AW). All adult whelks died at 15°C. B) Average cumulative increase in total shell length (± 1S.E. ribbon) with time for juvenile (TRJ) and adult (AW) (damaged and control) whelks

grown at 5, 10 and 15°C (TRJ) and 5 and 10°C (AW). More than 75% mortality was observed
within adults in the 15°C experiment within the first 5 days, so it was not continued. Tankreared juveniles (TRJ), wild-collected adult whelks (AW).

264 3.4 Structure of shell growth

265 Figure 5 shows the appearance of *B. undatum* thin shell lip sections viewed in transmitted light 266 (A and C) and the same sections illuminated with U.V. light (B and D). The point where the shell 267 was damaged is a truncated cut perpendicular to the growing shell (red arrow), with the 268 subsequently deposited thinner post-repaired shell (black arrow). Following exposure to 269 Calcein, post-shell damage, a bright fluorescent line is visible under U.V. light (small white arrow 270 Figure 5B). A similar response to shell damage and Calcein incorporation is visible in shells that 271 were damaged twice followed by Calcein exposure. Figure 5C shows two truncated cuts (red 272 arrows) and subsequent thinner shell re-growth (black arrows). The same section viewed in 273 U.V. shows two fluorescent lines corresponding to the two periods of Calcein exposure (small 274 white arrows Figure 5D). The thin sections show that following shell damage, new shell 275 deposition begins rapidly following Calcein incorporation on the inner surface of the crossed 276 lamellar layer with newly forming outer prismatic layer developing approximately 1mm inside 277 the point of shell damage (Figure 5A). Shell extension occurs rapidly as the removed shell is 278 quickly replaced to repair the missing shell at the lip and is thickened on the inner surface with 279 crossed lamellar shell structure. The extension of repaired shell following the second damage 280 incident however appears to be less than when the shell was first damaged (Figure 5C).





## Table 2: Summary of repair rates (mm<sup>2</sup>.day<sup>-1</sup>) and growth rates (mm.day<sup>-1</sup>) of experimentally-damaged *Buccinum undatum* grown under different food

## 292 rations and seawater temperatures.

Life Stage	Mean ± Standard Error									ANOVA	Where within Group	Fig.
Group	Tank Reared Juvenile (TRJ)			Juvenile Wild (JW)		Adult Wild (AW)						
	Dama	aged	Control	Damag	ged	Control	Damag	jed	Control	-		
Repair Rate	3.9 ±0.0	51 )71	-	2.04 ±0.01	4 .7	-	1.93 ±0.03	3 9	-	- Group: F= 865, p<0.001	All	2A
Growth Rate	0.0 ±0.0	91 )04	0.102 ±0.009	0.05 ±0.00	0 )5	0.062 ±0.006	0.02 ±0.00	7 12	0.029 ±0.005	Group: F= 90.6, p<0.001	All	2B
Food ration		Mean ± Standard Error								ANOVA	Where within Group	Fig.
Group	Unfed			Weekly		Daily						
	TRJ Damaged	AW Damaged	AW Control	TRJ Damaged	AW Damaged	AW Control	TRJ Damaged	AW Damaged	AW Control	-		
Repair Rate	5.765 ±0.025	4.756 ±0.002	-	5.405 ±0.226	4.760 ±0.001	-	5.954 ±0.203	4.761 ±0.001	-	Age: F= 54.6, p<0.001 Group: F= 1.87, p>0.05 Age*Group: F= 2.01, p>0.05	None	ЗA
Growth Rate	0.079 ±0.003	0.084 ±0.010	0.075 ±0.012	0.093 ±0.003	0.065 ±0.013	0.079 ±0.026	0.078 ±0.003	0.027 ±0.009	0.028 ±0.014	Age: F= 17.2, p<0.001 Group: F= 11.1, p<0.001 Age*Group: F= 6.53, p<0.05	Daily/ Unfed Daily/ Weekly	3B
Temperature		Mean ± Standard Error									Where within Group	Fig.
Group	5°C			10°C					15°C			
	TRJ Damage	TRJ AW Damaged Damaged		AW Control	TRJ Damaged	AW Dama	V ged	AW Control	TRJ Damaged	-		
Repair Rate	4.527 ±0.266	2.5 ±0.	545 179	-	4.605 ±0.269	3.76 ±0.1	56 86	-	10.397 ±0.253	Age: F= 296, p<0.001 Group: F= 183, p<0.001 Age*Group: F= 6.47, p<0.05	All	4A
Growth Rate	0.098 ±0.005	0.0 5 ±0.	026 006	0.050 ±0.014	0.105 ±0.004	0.06 ±0.0	51 09	0.045 ±0.010	0.106 ±0.009	Age: F= 72.7, p<0.001 Group: F= 3.41, p<0.05 Age*Group: F= 1.05, p>0.05	All	4B

295 This study used a series of controlled laboratory experiments to investigate the rate of shell 296 repair and shell growth in juvenile and adult common whelk, Buccinum undatum following 297 damage to the shell lip and examined how these rates are affected by food availability and 298 seawater temperature. The responses to shell damage recorded help provide an insight into 299 the factors involved in the recovery process of damaged individuals in wild populations 300 following shell damage as a result of storms, predation or demersal fishing activities. Our 301 findings reveal significant variation in responses between juvenile and adult *B. undatum*, with 302 these differences synchronous under all tested conditions for both rates of shell repair and 303 growth (i.e., TSL increase). Responsiveness to changing temperature and food availability 304 presented further differences in growth and repair rates, suggesting that there are multiple 305 factors influencing these rates in gastropod molluscs. The study therefore provides insight into 306 the key environmental drivers influencing shell repair in the wild.

307 Results from calcein staining showed that a damaged area did not only fill in with new shell 308 growth but continued the build-up of new shell layers on the inner surface of the shell and back 309 into the shell whorls throughout periods of damage and repair. The re-building of shell layers 310 by depositing beneath existing layers, although strengthening the repaired section, is likely to 311 result in heavier shells occurring within natural populations of areas with increased damage 312 incidents (Thomas and Himmelman, 1988; Ramsay et al., 2001; Stafford et al., 2015). Such 313 variation is often observed within wild whelk populations, with shell thickening and scarring 314 occurring more in certain geographical areas than others (Ramsay et al., 2001; Preston and 315 Roberts, 2007). In addition, this variation in shell scarring due to fishery disturbance has been 316 noted for other species, for example in the dog cockle *Glycymeris glycymeris*, with areas of 317 higher fishing seeing higher rates of damage and consequent repair (Kaiser et al., 2000;

318 Mensink et al., 2000). Additionally, predation pressures and wave action see further shell 319 variation in scarring events, as seen within the painted top shell Calliostoma 320 zizyphinum (Preston and Roberts, 2007) and the hydrothermal whelk species Buccinum 321 thermophilum (Martell et al., 2002), through incidents of shell chipping. Although the 322 thickening of the shell would increase resistance to a re-break, the process itself leaves the 323 individual vulnerable to its surroundings immediately post-damage, due to the large metabolic 324 implications of shell growth along with the damaged shell leaving the soft tissues exposed 325 (Palmer, 1992; Frieder et al., 2017).

326 Total shell length was found to continue to increase throughout periods of repair and, although 327 a significant difference in the rate of TSL increase was not observed, shell damage did result in 328 some deviation in daily growth trends of repairing animals, when compared with control 329 groups. It was expected that due to the high metabolic cost of shell growth and 330 biomineralization, the rate of TSL increase would dramatically reduce due to the shell repair 331 process in damaged whelks (Ebert, 1968; Ruppert et al., 2004; Melzner et al., 2011; Thomsen 332 et al., 2013). However, the results indicate the importance of continual shell growth even when 333 the shell is damaged, with little difference in TSL between the control and damaged groups. 334 This is likely an evolutionary development to ensure that whelks continue to increase in size so 335 that they reach a spatial size refuge from a range of predators as quickly as possible. Although 336 there has been limited research into this, continual shell growth has recently been observed in 337 the brachiopod Liothyrella uva (Cross et al., 2015). This build-up occurring, with the complete 338 re-growing of new shell layers could further be an influencing factor in the continued TSL 339 increase due to increased shell deposition and biomineralization.

The influence of age on an individual's growth trajectory has long been studied across a range
of animal taxa (von Bertalanffy, 1938; Richardson, 2001; West et al., 2001; Sibley et al., 2015).

342 As expected from theory, the results of this study show ontogenetic differences in rates of shell 343 growth and repair, with slower rates for both processes observed in older individuals; tank-344 reared juvenile whelks repairing their damaged shell at a rate almost twice that of adult whelks. 345 These responses have often been seen as a result of constraints to body size and development, 346 with larger older individuals seeing reduced rates of growth (Tanabe, 1988; Richardson, 2001). 347 With an increased body size, the trade-off between additional metabolic costs, including 348 maturation and reproductive success, outweighs the requirement of further shell production 349 resulting in a further decline in growth rates, as energy allocation is shifted from growth to 350 reproduction (Tanabe, 1988; Palmer, 1990; Kideys, 1996; Richardson, 2001; Thomsen et al., 351 2013). In addition, an increased need for protection at a smaller size would further require 352 faster rates of growth and repair, as an individual needs to grow until they reach a size 353 threshold at which predation vulnerability is reduced, i.e., a size refuge is reached (Chase, 1999; 354 Karythis et al., 2020). Nonetheless, the capacity for retraction into and protection from the shell 355 is of key importance for increased rapid escape capabilities from a surrounding threat (Tanabe, 356 1988; Palmer, 1990; Seed and Hughes, 1995; Kideys, 1996).

357 Metabolic cost is a fundamental part of growth, with energy consumption vital for development 358 and survival. It is interesting to note that the results of this study indicate that food availability 359 had no significant effect on shell repair rates with unfed individuals and those fed daily showing 360 similar rates of shell repair. However, differences in growth (as indicated by changes in TSL) 361 were observed between the different feeding groups. Under unfed or daily feeding conditions, 362 juvenile whelks displayed faster repair rates than those offered food once per week, whereas 363 for TSL growth, weekly food conditions for juvenile and non-damaged adult whelks saw the 364 fastest overall rates. Although the response to a lack of food did not follow expected responses, 365 whereby for many marine organisms, energy is often saved for increased survival capabilities,

366 or processes are reduced until more favourable conditions arise (Tomanek and Somero, 1999; 367 Stillman, 2003; Melzner et al., 2011). The observed response in our experiments likely suggests 368 that the need to repair the shell is overwhelmingly important for protection and growth. Palmer 369 (1983) however, showed that starved gastropods used body reserves and supplementary 370 energy sources to fulfil the needs of shell formation, often through protein catabolism, a 371 process that continues throughout normal feeding but is relied upon more heavily when food 372 becomes scarce or unavailable. Without knowledge of the whelk's feeding behaviour in the 373 wild prior to capture, we do not know whether sufficient energy reserves may still have been 374 available to the unfed whelks or they were using protein catabolism to provide energy to repair 375 their shells and increase their TSL during their laboratory experimental conditions. Our initial 376 hypothesis was that there would be a link between food supply and shell repair rate, i.e., whelks 377 fed daily would repair their shells faster than those fed weekly and those with no food supply. 378 Our results demonstrated that there was no difference in repair rates between the three 379 different feeding regimes although there was a significant difference in growth rate between 380 the three groups of whelks. The daily fed whelks grew the slowest possibly because they over 381 ate as they had access to an unlimited food supply and were subsequently stressed. However, 382 throughout all daily feeding conditions no sign of distress to the whelks was observed and the 383 highest number of mortalities were in the unfed whelk groups and not those fed daily. We 384 postulate that during the summer the daily fed whelks may have allocated surplus energy from 385 their food intake to gametogenesis, a few months prior to when individuals would be engaging 386 in reproduction in the autumn, at the expense of increasing their TSL. Likewise, shell deposition

and rates of biomineralization are further affected through gene expression, and furtherthrough temperature changes (Joubert et al., 2014).

389 Temperature is a key environmental factor, with fluctuations in seawater temperature 390 dramatically affecting physiological processes within marine organisms, with a range of 391 responses and metabolic adjustments seen (Page and Hubbard, 1987; Prosser, 1991; Sokolova 392 and Portner, 2003; Harley et al., 2006). Benthic organisms have an increased susceptibility to 393 thermal stress (Foster, 1971; Harley et al., 2006) with greater impacts/effects observed in 394 juvenile/ smaller individuals (Pechenik et al., 2019; Levinton, 2020), a response observed within 395 this study. Temperature was found to have a significant effect on both rates of growth and 396 repair with juvenile whelks reared at 15°C presenting the highest rates observed. A similar 397 result was observed for adult whelks that showed increased rates at 10°C compared to 5°C but 398 could not be tested at 15°C due to their mortality rates increasing as their thermal limit was 399 reached and exceeded. It appears that juvenile whelk can handle much warmer seawater 400 temperatures demonstrated by the 15°C degree TRJ whelks that showed the fastest growth. 401 The observed responses would allow for early life development of whelks at shallower depth, 402 although as adult whelks (AW) are generally found in deeper, cooler waters, this could be 403 further reflective of the metabolic changes occurring (Valentinsson et al., 1999; McIntyre et al., 404 2015).

405

In some gastropod species certain environmental conditions (e.g., a combined change in pH and elevated seawater temperature) have been observed to reduce morphological size and have a negative effect on growth, as observed, for example, in the periwinkle (*Littorina saxatilis*) and gold-ringed cowry (*Monetaria annulus*) (Sokolova and Pörtner, 2000; Irie and Fischer, 2009; Melatunan et al., 2013). Observations from our study support previous work into 411 metabolism and temperature, with increased temperatures causing an increased rate of growth, up until the thermal tolerance is reached (Emmerson et al., 2020; Borsetti et al., 2021). 412 413 In these studies, elevated temperature resulted in increased rates of calcification, due to 414 heightened enzyme activity associated with the biomineralization process and consequently 415 raised uptake of Ca<sup>2+</sup> ions (Bevelander and Nakahara, 1969; Pons et al., 2002; Thomsen et al., 416 2010; Lervik et al., 2013). Although the results from our study showed a difference in response 417 to changing temperatures. Plasticity to surrounding temperature has been seen to be 418 dependent on pH, and external stimuli in other marine molluscs (Melatunan et al., 2013).

419 In conclusion, our study has shown that shell growth and shell repair rates in B. undatum are 420 strongly influenced by factors such as age, seawater temperature, and food availability. The 421 growth and repair of shells of marine gastropods can vary greatly both within and between 422 species (Palmer, 1990). This variability is not only a result of phenotypic differences, such as 423 shell thickness and shape (Brookes and Rochette, 2007; Magnúsdóttir et al., 2018), but also as 424 a response to individual genotype (Magnúsdóttir et al., 2019; Goodall et al., 2021) and 425 environmental cues such as predator presence and seawater temperature changes (Zdelar et 426 al., 2018). It is important to understand how a combination of these factors impact the 427 populations of *B. undatum*, particularly given their current commercial importance, and 428 interest in fishery management strategies. In understanding how shell growth and repair occurs 429 in the whelk, and its impacts on the organism, we can better understand its resilience to both 430 direct and indirect fishery impacts.

431 Acknowledgements

This work was funded by the family of C.N.C. and Bangor University, and we would like to thankMark and Lina Colvin for their support and reviewing of this manuscript. We are grateful to

- 434 Berwyn Roberts for collecting *Buccinum undatum* and for his invaluable help with maintaining
- 435 aquaria and animal husbandry. We also thank Stef Kraft, Helèna Bonici-Strohmer, and George
- 436 Collinson for their assistance during this project.

Bergmann, M., Beare, D.J., Moore, P.G., 2001. Damage sustained by epibenthic invertebrates
discarded in the Nephrops fishery of the Clyde Sea area, Scotland. Journal of Sea Research,
440 45(2), pp. 105-118.

- Bevelander, G., Nakahara, H., 1969. An electron microscope study of the formation of the
  nacreous layer in the shell of certain bivalve molluscs. Calcified Tissue Research, 3(1), pp. 8492.
- Biro, P.A., Sampson, P., 2015. Fishing directly selects on growth rate via behaviour: implications
- 445 of growth-selection that is independent of size. Proceedings of the Royal Society B: Biological
- 446 Sciences, 282(1802), p.20142283.
- 447 Bonucci, E., 2007. Biological calcification. Normal and pathological processes in the early stages.
  448 Springer-Verlag, Berlin, Germany.
- Borsetti, S., Hollyman, P.R., Munroe, D., 2021. Using a sclerochronological approach to
  determine a climate-growth relationship for waved whelk, *Buccinum undatum*, in the US MidAtlantic. Estuarine, Coastal and Shelf Science, 252, p.107255.
- Brookes, J.I., Rochette, R., 2007. Mechanism of a plastic phenotypic response: predatorinduced shell thickening in the intertidal gastropod *Littorina obtusata*. Journal of Evolutionary
  Biology, 20(3), pp.1015-1027.
- 455 Cadée, G.C., Boon, J.P., Fischer, C.V., Mensink, B.P., Ten Hallers-Tjabbes, C.C., 1995. Why the
  456 whelk (*Buccinum undatum*) has become extinct in the Dutch Wadden Sea. Netherlands Journal
  457 of Sea Research, 34(4), pp. 337-339.

458 Chase, J.M., 1999. Food web effects of prey size refugia: variable interactions and alternative

459 stable equilibria. The American Naturalist, 154(5), pp.559-570.

- 460 Checa, A., 1993. Non-predatory shell damage in recent deep-endobenthic bivalves from Spain.
  461 Palaeogeography, Palaeoclimatology, Palaeoecology, 100(3), 309-331.
- 462 Clark, M.R., Althaus, F., Schlacher, T.A., Williams, A., Bowden, D.A., Rowden, A.A., 2016. The
  463 impacts of deep-sea fisheries on benthic communities: a review. ICES Journal of Marine
  464 Science: Journal Du Conseil, 73(suppl 1), i51-i69.
- 465 Cross, E.L., Peck, L.S., Harper, E.M., 2015. Ocean acidification does not impact shell growth or
- 466 repair of the Antarctic brachiopod *Liothyrella uva* (Broderip, 1833). Journal of Experimental
- 467 Marine Biology and Ecology, 462, pp.29-35.
- 468 De Vooys, C., Van der Meer, J., 1998. Changes between 1931 and 1990 in by-catches of 27
  469 animal species from the southern North Sea. Journal of Sea Research, 39(3-4), pp. 291-298.
- 470 De Vooys, C.G.N., Van Der Meer, J., 2010 The whelk (*Buccinum undatum* L.) in the western
- 471 Dutch Wadden Sea in the period 1946–1970: Assessment of population characteristics and472 fishery impact. J Sea Res 63:11–16.
- 473 Ebert, T.A., 1968. Growth rates of the sea urchin *Strongylocentrotus purpuratus* related to food
  474 availability and spine abrasion. Ecology, 49(6), pp. 1075-1091.
- Emmerson, J.A., Hollyman, P.R., Bloor, I.S.M., Jenkins, S.R., 2020. Effect of temperature on the
  growth of the commercially fished common whelk (*Buccinum undatum*, L.): a regional analysis
  within the Irish Sea. Fisheries Research, 223, p.105437.

Fahy, E., Carroll, J., Hother-Parkes, L., O'Toole, M., Barry, C., 2005. Fishery associated changes
in the Whelk *Buccinum undatum* stock in the southwest Irish Sea, 1995-2003. Dublin: Fisheries
Science Services.

Fahy, E., Masterson, E., Swords, D., Forrest, N., 2000. A Second Assessment of the whelk fishery *Buccinum undatum* in the southwest Irish Sea with particular reference to its history of
management by size limit. Dublin: Fisheries Science Services.

Fahy, E., Yalloway, G., Gleeson, P., 1995. Appraisal of the whelk *Buccinum undatum* fishery of
the southern Irish sea with proposals for a management strategy. Irish Fisheries Investigations,
Series B (Marine), 42.

Findlay, H.S., Wood, H.L., Kendall, M.A., Spicer, J.I., Twitchett, R.J., Widdicombe, S., 2011.
Comparing the impact of high CO2 on calcium carbonate structures in different marine
organisms. Marine Biology Research, 7(6), pp. 565-575.

490 Foster, B., 1971. On the determinants of the upper limit of intertidal distribution of barnacles

491 (Crustacea: Cirripedia). The Journal of Animal Ecology, 40(1), pp. 33-48.

492 France, J., Dijkstra, J., Thornley, J.H., Dhanoa, M.S., 1996. A simple but flexible growth
493 function. Growth, development, and aging: GDA, 60(2), pp.71-83.

494 Frieder, C.A., Applebaum, S.L., Pan, T.C.F., Hedgecock, D., Manahan, D.T., 2017. Metabolic cost
495 of calcification in bivalve larvae under experimental ocean acidification. ICES Journal of Marine
496 Science, 74(4), pp.941-954.

497 Golikov, A. N., 1968. Distribution and variability of long-lived benthic animals as indicators of498 currents and hydrological conditions. Sarsia. 34, pp. 199-208.

- Goodall, J., Westfall, K.M., Magnúsdóttir, H., Pálsson, S., Örnólfsdóttir, E.B., Jónsson, Z.O., 2021.
  RAD sequencing of common whelk, *Buccinum undatum*, reveals fine-scale population
  structuring in Europe and cryptic speciation within the North Atlantic. Ecology and
  evolution, 11(6), pp.2616-2629.
- Harley, C.D., Randall Hughes, A., Hultgren, K.M., Miner, B.G., Sorte, C.J., Thornber, C.S.,
  Rodriguez, L.F., Tomanek, L., Williams, S.L., 2006. The impacts of climate change in coastal
  marine systems. Ecology Letters, 9(2), pp. 228-241.
- 506 Hayward, P.J., Ryland, J.S. (eds.), 2011 (reprinted). Handbook of the Marine Fauna of North-

507 West Europe, Chapter 10: Molluscs, p. 535. Oxford: Oxford University Press.

- Hilborn, R., Minte-Vera, C.V., 2008. Fisheries-induced changes in growth rates in marine
  fisheries: are they significant? Bulletin of Marine science, 83(1), pp.95-105.
- Hollyman, P.R., Chenery, S.R., Leng, M.J., Laptikhovsky, V.V., Colvin, C.N., Richardson, C.A.,
  2018. Age and growth rate estimations of the commercially fished gastropod Buccinum
  undatum. ICES Journal of Marine Science, 75(6), pp.2129-2144.
- Hollyman, P.R., Leng, M.J., Chenery, S.R., Sloane, H.J., Richardson, C.A., 2020. Calibration of
  shell δ180 from the common whelk *Buccinum undatum* highlights potential for
  palaeoenvironmental reconstruction. Palaeogeography, Palaeoclimatology,
  Palaeoecology, 560, p.109995.
- 517 Irie, T., Fischer, K., 2009. Ectotherms with a calcareous exoskeleton follow the temperature-518 size rule-evidence from field survey. Marine Ecology Progress Series, 385, pp.33-37.

29

520 Gueguen, Y., 2014. Temperature and food influence shell growth and mantle gene expression

- 521 of shell matrix proteins in the pearl oyster *Pinctada margaritifera*. PLoS One, 9(8), p.e103944.
- 522 Kaehler, S., McQuaid, C.D., 1999. Use of the fluorochrome calcein as an in situ growth marker523 in the brown mussel *Perna perna*. Marine Biology 133:455-460.
- Kaiser, M.J., Ramsay, K., Richardson, C.A., Spence, F.E., Brand, A.R., 2000. Chronic fishing
  disturbance has changed shelf sea benthic community structure. Journal of Animal
  Ecology, 69(3), pp.494-503.
- Karythis, S., Cornwell, T.O., Noya, L.G., McCarthy, I.D., Whiteley, N.M., Jenkins, S.R., 2020. Prey
  vulnerability and predation pressure shape predator-induced changes in O<sub>2</sub> consumption and
  antipredator behaviour. Animal Behaviour, 167, pp.13-22.
- Kideys, A.E, Nash, R.D.M., Hartnoll, R.G., 1993. Reproductive cycle and energetic cost of
  reproduction of the neogastropod *Buccinum undatum* in the Irish Sea. Journal of the Marine
  Biological Association of the United Kingdom, 73(02), pp. 391-403.
- Kideys, A.E., 1996. Determination of age and growth of *Buccinum undatum* L. (Gastropoda) off
  Douglas, Isle of Man. Helgoländer Meeresuntersuchungen, 50(3), pp. 353-368.
- Lervik, A., Bedeaux, D., Kjelstrup, S., 2013. Active transport of the Ca2+- pump: introduction of
  the temperature difference as a driving force. European Biophysics Journal, 42(5), pp. 321-331.
- Levinton, J., 2020. Thermal stress: The role of body size and the giant major claw in survival and
  heat transfer of a fiddler crab (*Leptuca pugilator*). Journal of Experimental Marine Biology and
  Ecology, 530, p.151428.

Magnúsdóttir, H., Pálsson, S., Westfall, K.M., Jónsson, Z.O., Örnólfsdóttir, E.B., 2019.
Morphological variation in genetically divergent populations of the common whelk, *Buccinum undatum* (Gastropoda: Buccinidae), across the North Atlantic. Biological Journal of the Linnean
Society, 128(1), pp.93-106.

547 Marine Management Organisation., 2020. UK Sea Fisheries Statistics 2020. Office for National
548 Statistics, London.

Martell, K.A., Tunnicliffe, V., Macdonald, I.R., 2002. Biological features of a buccinid whelk
(Gastropoda, Neogastropoda) at the Endeavour ventfields of Juan de Fuca Ridge, Northeast
Pacific. Journal of Molluscan Studies, 68(1), pp.45-53.

552 McIntyre, R., Lawler, A., Masefield, R., 2015. Size of maturity of the common whelk, *Buccinum* 

*undatum*: is the minimum landing size in England too low? Fisheries Research, 162, pp.53-57.

Melatunan, S., Calosi, P., Rundle, S.D., Widdicombe, S., Moody, A.J., 2013. Effects of ocean
acidification and elevated temperature on shell plasticity and its energetic basis in an intertidal
gastropod. Marine Ecology Progress Series, 472, pp.155-168.

Melzner, F., Stange, P., Trübenbach, K., Thomsen, J., Casties, I., Panknin, U., Gorb, S.N.,
Gutowska, M.A., 2011. Food supply and seawater pCO<sub>2</sub> impact calcification and internal shell
dissolution in the Blue Mussel *Mytilus edulis*. PloS One, 6(9), e24223.

561 Shell damage and mortality in the common whelk *Buccinum undatum* caused by beam trawl562 fishery. Journal of Sea Research, 43(1), pp. 53-64.

563 Ohnishi, S., Akamine, T., 2006. Extension of von Bertalanffy growth model incorporating growth

- 564 patterns of soft and hard tissues in bivalve molluscs. Fisheries Science, 72(4), pp.787-795.
- Page, H.M., Hubbard, D.M., 1987. Temporal and spatial patterns of growth in mussels *Mytilus edulis* on an offshore platform: relationships to water temperature and food availability. Journal
  of Experimental Marine Biology and Ecology, 111(2), pp. 159-179.
- 568 Palmer, A.R., 1983. Relative cost of producing skeletal organic matrix versus calcification:
  569 Evidence from marine gastropods. Marine Biology, 75(2-3), pp. 287-292.
- 570 Palmer, A.R., 1990. Effect of crab effluent and scent of damaged conspecifics on feeding,
  571 growth, and shell morphology of the Atlantic dogwhelk *Nucella lapillus* (L.). Hydrobiologia,
  572 193(1), pp. 155-182.
- 573 Palmer, A.R., 1992. Calcification in marine molluscs: how costly is it?. Proceedings of the
  574 National Academy of Sciences, 89(4), pp.1379-1382.
- 575 Pechenik, J.A., Chaparro, O.R., Franklin, A., Mardones, M.L., Montory, J.A., 2019. Thermal
  576 tolerance of intertidal and subtidal adults and embryos of the marine gastropod *Crepipatella*577 *peruviana*. Marine Ecology Progress Series, 616, pp.67-81.
- 578 Pepin, P., 1991. Effect of temperature and size on development, mortality, and survival rates
  579 of the pelagic early life history stages of marine fish. Canadian Journal of Fisheries and Aquatic
  580 Sciences, 48(3), pp.503-518.

- 581 Pons, G., Evangelisti, V., Caprì, F., Mozzone, S., Viarengo, A., 2002. Cytochemical localization
- 582 and quantification of plasma membrane Ca2 -ATPase activity in mollusc digestive gland cells.
- 583 European Journal of Histochemistry, 46(1), p. 31-40.
- 584 Preston, S.J., Roberts, D., 2007. Variation in shell morphology of *Calliostoma zizyphinum*585 (Gastropoda: Trochidae). Journal of Molluscan Studies, 73(1), pp.101-104.
- 586 Prosser, C.L., Graham, G., Galton, V., 1991. Hormonal regulation of temperature acclimation in
- 587 catfish hepatocytes. Journal of Comparative Physiology B, 161(1), pp.117-124.
- 588 Ramsay, K., Richardson, C.A., Kaiser, M.J., 2001. Causes of shell scarring in dog cockles
- 589 *Glycymeris glycymeris* L. Journal of Sea Research, 45(2), pp. 131-139.
- 590 Richardson, C.A., 2001. Molluscs as archives of environmental change. Oceanography and
  591 Marine Biology: an Annual Review, 39, pp. 103-164.
- S92 Ruppert, E.E., Fox, R.S, Barnes, R.D., 2004. Invertebrate Zoology. Delhi: Cengage Learning,
  S93 Chapter 12: Mollusca, pp. 269-301.
- Sebens, K.P., 1987. The ecology of indeterminate growth in animals. Annual review of ecologyand systematics, 18(1), pp.371-407.
- Seed, R., Hughes, R.N., 1995. Criteria for prey size-selection in molluscivorous crabs with
  contrasting claw morphologies. Journal of Experimental Marine Biology and Ecology, 193(1-2),
  pp. 177-195.
- Sepúlveda, R.D., Jara, C.G., Gallardo, C.S., 2012. Morphological analysis of two sympatric
  ecotypes and predator-induced phenotypic plasticity in *Acanthina monodon* (Gastropoda:
  Muricidae). Journal of Molluscan Studies, 78(2), pp.173-178.

- Sibly, R. M., Baker, J., Grady, J. M., Luna, S. M., Kodric-Brown, A., Venditti, C., Brown, J. H., 2015.
  Fundamental insights into ontogenetic growth from theory and fish. Proceedings of the
  National Academy of Sciences 112, 13934-13939.
- Sokolova, I.M., Pörtner, H.O., 2003. Metabolic plasticity and critical temperatures for aerobic
  scope in a eurythermal marine invertebrate (*Littorina saxatilis*, Gastropoda: Littorinidae) from
  different latitudes. Journal of Experimental Biology, 206(1), pp.195-207.
- Sokolova, I.M., Bock, C., Pörtner, H.O., 2000. Resistance to freshwater exposure in White Sea *Littorina* spp. I: Anaerobic metabolism and energetics. Journal of Comparative Physiology B,
  170(2), pp.91-103.
- Solas, M.R., Hughes, R.N., Márquez, F., Brante, A., 2015. Early plastic responses in the shell
  morphology of *Acanthina monodon* (Mollusca, Gastropoda) under predation risk and water
  turbulence. Marine Ecology Progress Series, 527, pp.133-142.
- 614 Stillman, J.H., 2003. Acclimation capacity underlies susceptibility to climate change. Science615 (New York), 301(5629), p. 65.
- 616 Tanabe, K., 1988. Age and growth rate determinations of an intertidal bivalve, *Phacosoma*617 *japonicum*, using internal shell increments. Lethaia, 21(3), pp. 231-241.
- Thomsen, J., Casties, I., Pansch, C., Körtzinger, A., Melzner, F., 2013. Food availability outweighs
  ocean acidification effects in juvenile *Mytilus edulis*: Laboratory and field experiments. Global
  Change Biology, 19(4), pp. 1017-1027.
- 621 Thomsen, J., Gutowska, M., Saphörster, J., Heinemann, A., Trübenbach, K., Fietzke, J.,
  622 Hiebenthal, C., Eisenhauer, A., Körtzinger, A., Wahl, M., Melzner, F., 2010. Calcifying

- 623 invertebrates succeed in a naturally CO<sub>2</sub> enriched coastal habitat but are threatened by high
- 624 levels of future acidification. Biogeosciences, 7(11), pp. 3879-3891.
- Tomanek, L., Somero, G.N., 1999. Evolutionary and acclimation-induced variation in the heatshock responses of congeneric marine snails (genus *Tegula*) from different thermal habitats:
  implications for limits of thermotolerance and biogeography. The Journal of Experimental
  Biology, 202(21), pp. 2925-2936.
- Valentinsson, D., Sjödin, F., Jonsson, P.R., Nilsson, P., Wheatley, C., 1999. Appraisal of the
  potential for a future fishery on whelks (*Buccinum undatum*) in Swedish waters: CPUE and
  biological aspects. Fisheries research, 42(3), pp.215-227.
- 632 Von Bertalanffy, L., 1938. A quantitative theory of organic growth (inquiries on growth laws. II).
  633 Human Biology, 10(2), pp. 181-213.
- West, G. B., Brown, J. H., Brian J. Enquist, B. J., 2012. A general model for ontogenetic growth.
  Nature 413, 628-631.
- Zdelar, M., Mullin, F., Cheung, C., Yousif, M., Baltaretu, B., Stone, J.R., 2018. Pollution-,
  temperature-and predator-induced responses in phenotypically plastic gastropod shell
  traits. Molluscan Research, 38(1), pp.34-40.
- 639 Zuur, A. F., Ieno, E. N., Walker, N., Saveliev, A. A., Smith, G. M., 2009. Mixed effects models and
  640 extensions in ecology with R. New York, USA: Springer.
- 641