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## Original Article

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
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# Shell growth and age determined from annual lines in the southern warm-water limpet *Patella depressa* at its poleward geographic boundaries

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## Abstract

In shell-secreting molluscs, age and growth rate of individuals and hence their performance can normally be measured using growth lines that are deposited in the shell throughout their lives. An annual periodicity of growth line formation of the warm-water limpet *Patella depressa* was established using marked and recaptured individuals from north Wales, UK. Length at age from suitably prepared shell sections was determined in limpets from non-range-edge populations and at two range edges, where different demographic attributes have been recorded. Individuals collected from their poleward range-edge in north Wales were older when compared with individuals at their range-edge in southern England. Shells collected from southern England were characterized by rapid growth with most individuals reaching >30 mm in maximum length by the fourth or fifth year, contrasting with those from north Wales, where most shells only reached this size at 7–10 years of age. Von Bertalanffy growth coefficients (*K*-values) were negatively related to *P. depressa* density, showing faster growth in lower total densities of both *P. depressa* and *Patella vulgata* combined. Higher intra-specific effects on *K*-values were found in *P. depressa* compared with its congener *P. vulgata*, with stronger effects in north Wales than in southern England. These results confirm differences in population patterns and individual traits between the two leading edges of *P. depressa*. Understanding annual growth in *P. depressa* over large scales could help to disentangle the processes determining differences in shell growth and age structure seen at the two range edges of this limpet species.

## Introduction

Growth rate can be considered as a metric of performance of an organism (Pörtner *et al.*, 2005), being influenced by both abiotic and biotic factors (Richardson, 2001). Invertebrate skeletons, particularly of molluscs, frequently provide a continuous record of ontogenetic growth and an archive of environmental variation during the lifespan of an individual (Rhoads & Lutz, 1980). In shelled molluscs, such as limpets, the shell is accreted incrementally as the animal grows (MacClintock, 1967). This incremental growth is separated by growth rings or growth lines (Richardson, 2001), which may be observed on both the surface (rings) and in cross-sections of the shell (lines; Rhoads & Lutz, 1980). As such, they can be used to estimate individual growth rates and hence population performance within the distribution of a species.

Growth line formation reflects responses to a variety of abiotic and biotic conditions (Richardson, 2001). Seasonal patterns of seawater temperature can have substantial impacts on growth in benthic molluscs (Surge *et al.*, 2013). Winter conditions inhibit normal limpet shell and somatic growth, resulting in slower growth rates (Lewis & Bowman, 1975), manifested in narrower annual and tidally related shell growth increments (Crisp *et al.*, 1990; Richardson & Liu, 1994; Lomovasky *et al.*, 2020). By contrast, rapid shell growth, as illustrated by wider increments, is driven by more favourable conditions. Fast growth associated with warmer seawater temperatures and greater food availability yields increments that are easily distinguishable when compared with finer increments accreted during colder, less favourable conditions (Picken, 1980; Surge *et al.*, 2013; Lomovasky *et al.*, 2020). For instance, seasonal variation in shell deposition in the bivalve *Arctica islandica* (Butler *et al.*, 2009) and in limpet species such as *Patella vulgata* (Surge *et al.*, 2013; Ambrose *et al.*, 2016; Gutiérrez-Zugasti *et al.*, 2017) and *Patella rustica* (Prusina *et al.*, 2015) results in annual growth line formation, but marine benthic molluscs often also show increments at daily (Bock & Miller, 1994; Schöne *et al.*, 2005) and tidal scales (Richardson *et al.*, 1979, 1980; Bock & Miller, 1994).

Growth line formation in molluscs depends on processes internal to the individual (e.g. reproduction). Reproduction requires a large, and often exclusive, energy investment

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(Blackmore, 1969), thereby reducing the budget for somatic and shell growth (Wright & Hartnoll, 1981; Sato, 1995; Pörtner *et al.*, 2005) and leads to slower growth during the reproductive season (e.g. in *P. vulgata*, Hawkins & Hartnoll, 1982). Thus, an annual line is commonly formed at the end of the gonadal development and spawning periods as, for example, in the top shell *Phorcus lineatus* (García-Escárzaga *et al.*, 2019). Cessation of shell growth produces a discernible break in the microstructure of the shell, which may be recognized as a thick line in both cross-section and external features of these gastropod (García-Escárzaga *et al.*, 2019, 2020) and bivalve shells (Sato, 1995). In the northern hemisphere, annual line formation in bivalve species usually occurs during winter in species growing at high latitudes (Surge & Schöne, 2014), although factors such as habitat depth and interaction with the dynamics of mixed seawater layers may play a role in changing the timing of increment growth in deeper-water species (Estrella-Martínez *et al.*, 2019). Hence, these seasonal influences and reproductive traits imprinted in shells can be recognizable and thus quantifiable as records of species' growth rates.

Patellid limpets play a major role in controlling and structuring intertidal communities by the consumption of microbial biofilms, which are composed of cyanobacteria, microalgae, propagules and juveniles of macroalgae, thereby regulating algal cover (Hawkins, 1981; Hawkins & Hartnoll, 1983; Hartnoll & Hawkins, 1985; Jenkins *et al.*, 2005; Coleman *et al.*, 2006). Due to their simple geometric morphology (Ekaratne & Crisp, 1983) as well as their sessile nature and homing behaviour (Santini *et al.*, 2014), limpets have been used as a tractable species to investigate population parameters such as growth rates (Jenkins & Hartnoll, 2001; Henriques *et al.*, 2012; Sousa *et al.*, 2017), age structure (Fenberg & Roy, 2012; Borges *et al.*, 2015, 2016; Martins *et al.*, 2017), sexual maturity (Guerra & Gaudencio, 1986; Ribeiro *et al.*, 2009), recruitment and mortality (Henriques *et al.*, 2012, 2017; Sousa *et al.*, 2017), including recent work testing hypotheses about the relative performances of overlapping species of cold- and warmer-water limpet species in various parts of their geographic distributions (e.g. Lima *et al.*, 2016; Aguilera *et al.*, 2018; Oróstica *et al.*, 2020). External limpet shell rings (annuli) are often preserved and quantifiable with the naked eye (Bretos, 1980; Picken, 1980) although distinguishing between disturbance rings and those of annual origin can be problematic. However, annual increments are frequently clearly visible in limpet shell cross-sections (e.g. Prusina *et al.*, 2015; Gutiérrez-Zugasti *et al.*, 2017; Prendergast & Schöne, 2017; García-Escárzaga *et al.*, 2020), enabling estimation of the lifespan and growth rates of *Patella* species to be determined at different locations within their geographic distribution.

*Patella depressa*, a warm water limpet species, is distributed from North Africa to the British Isles (Figure 1A; Orton & Southward, 1961; Guerra & Gaudencio, 1986; Southward *et al.*, 1995; Ribeiro *et al.*, 2009). It has two separate leading edges as it has spread northwards after the last Ice Age (Figure 1; Southward *et al.*, 1995), in north Wales (N Wales; Crisp & Knight-Jones, 1954) and in south and south-east England (S/SE England; Crisp & Southward, 1958). In S/SE England, the English Channel has been described as analogous to a poleward gradient (Herbert *et al.*, 2009) for multiple intertidal species; the eastern Channel and southern North Sea are colder in winter than the western side due to continental influences from central Europe (for details see Crisp & Southward, 1958; Lewis, 1964). Different patterns of abundance (Kendall *et al.*, 2004; Oróstica, 2018), individual growth and population mortality rates have recently been reported at the two range limits of *P. depressa* (Oróstica *et al.*, 2020). The last northerly and easterly breeding populations of *P. depressa* occur at Abersoch in N Wales and

Southsea in S/SE England (Figure 1B; S. J. Hawkins & M. H. Oróstica pers. obs.).

Our goal was to investigate the age and growth rate of individual *P. depressa*, through validation of the putative annual growth lines seen in shell sections of *in situ* marked and re-measured limpets in the field. We compared the age and growth of *P. depressa* in populations at its two poleward range edges with non-range-edge populations in south-west England (SW England; Figure 1B). Here, *P. depressa* populations resemble those further south in Europe (France, Spain or Portugal) making up over 50% of the total limpet population on the mid-shore, with occasional patches where up to 100% can be found in a 50 × 50 cm quadrat (see Hawkins *et al.*, 2008; S. J. Hawkins unpubl. data). The number of growth lines were counted to estimate their longevity and mean length at age measured; von Bertalanffy growth curves were fitted to these data to estimate shell growth rate. We expected that at both range edges, individuals of *P. depressa* would grow more slowly. In addition, since unsuitable thermal conditions will curtail species reproduction, we expect limited or variable recruitment at species' range edges (Helmuth *et al.*, 2006), and hence a greater proportion of older individuals representing sporadic year class success. Additionally, we examined biotic control of growth. Density-dependent processes such as competition can influence limpet growth patterns (Thompson *et al.*, 2000; Boaventura *et al.*, 2002, 2003). Thus, we also tested the influence of population density on growth performance of *P. depressa* at the scale of the habitat-patch.

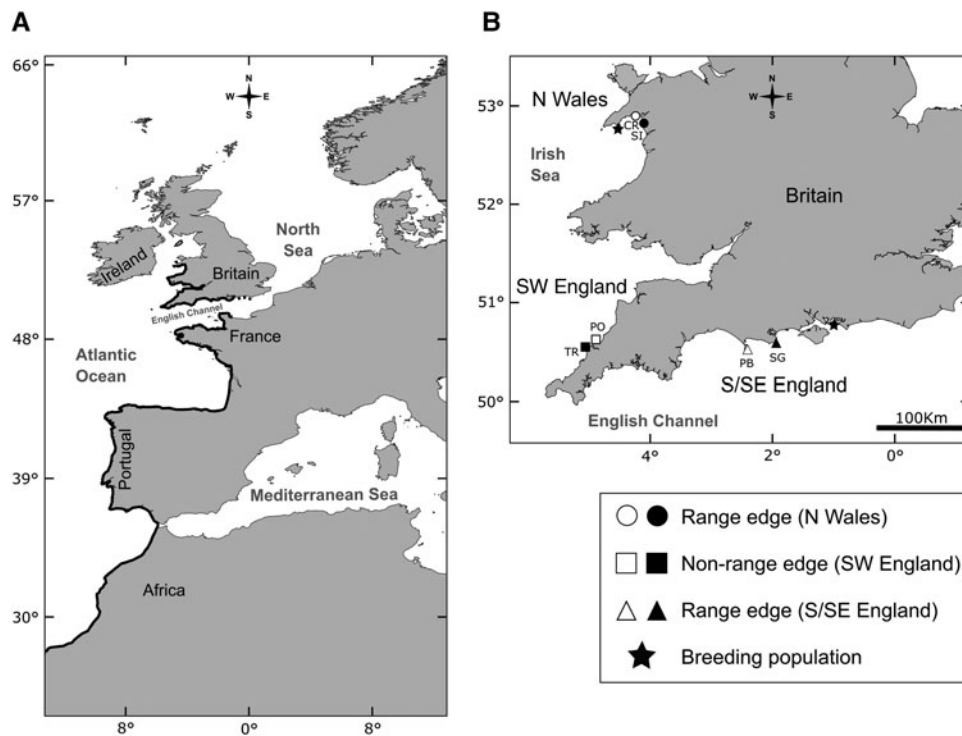
## Materials and methods

### Morphological traits of *P. depressa*

The shell of *P. depressa* is usually flatter than *P. vulgata* and *Patella ulysipponensis* with distinctive orange-brown marginal rays on the inner surface (Evans, 1947; Bowman, 1981). The apex (Ap) is located towards the anterior (Ant) end of the central axis of the shell (Figure 2A). Shells have fine radiating ribs and a markedly oval or triangular shape at their posterior (Post) end (Figure 2B; Bowman, 1981). The maximum length (ML) of *P. depressa* is usually between 30–35 mm (Figure 2A; Bowman, 1981) and although larger individuals have been found (Orton & Southward, 1961; Borges *et al.*, 2015), it never grows as large as *P. vulgata* in Britain (Evans, 1947; Borges *et al.*, 2015).

### Field validation of growth line formation

To validate the periodicity of growth lines seen in the shells of *P. depressa*, 80 specimens between 19–25 mm in ML were selected on rock boulders at mid tide level (MTL) on the exposed rocky shore at Shell Island in N Wales (Figure 1B), between the 22–29 June 2015. Each limpet was dried *in situ* with absorbent paper and labelled using a small (5 × 5 mm) waterproof numerical label (Brady®, TMM-0-49-PK model, <https://www.bradyid.com>) affixed to their shell with superglue. The ML of each tagged limpet was measured initially and again on three different occasions: 3 August 2015, 23 November 2015 and 21 March 2016. All field measurements were made with callipers to a resolution of 0.1 mm. After a two-year period had elapsed (Date: 13 June 2017), 18 limpets with their labels still attached were located. ML was measured *in situ* and they were carefully removed from the rock surface. Their soft tissues were removed in the laboratory and shells rinsed clean with fresh water and air-dried at ambient temperature. A subgroup of 10 shells with the least epibionts or least damage to the growing edge was selected for embedding in resin and growth line validation analysis (Table S1).



**Fig. 1.** (A) Geographic range of *Patella depressa* (black line; from N Wales to S/SE England to Senegal [not shown on map], Africa). (B) Regions (N = 3) and locations (N = 6) selected to measure growth and age at three regions defined by two leading edges of the range of *P. depressa*: (1) northern (N Wales: ○ Criccieth [CR]; ● Shell Island [SI]), and (2) eastern (S/SE England: △ Portland Bill [PB]; ▲ Swanage [SG]), and by non-range-edge populations in (3) SW England (□ Polzeath [PO]; ■ Trevone [TR]). In (B), the 2 black stars indicate last breeding populations of *P. depressa* towards both range edges in Britain (S. J. Hawkins & M. H. Oróstica pers. obs.).

### Geographic variation in longevity and shell growth rate

Shells of *P. depressa* were studied from six locations in the British Isles (Figure 1B). They were located at both range edges in N Wales (Criccieth and Shell Island) and S/SE England (Portland Bill and Swanage); and from non-range-edge populations in SW England (Polzeath and Trevone; see Figure 1B).

The abundance of *P. depressa* and *P. vulgata* was estimated between June and July 2016 from approximately mean high water neap (HWN) to mean tide level (MTL), where *P. depressa* reaches its maximum abundances on both semi-exposed and exposed shores (Orton & Southward, 1961; Oróstica *et al.*, 2020). At each location, the total numbers of *Patella* species were counted in each of ten  $0.5 \times 0.5$  m quadrats along a transect parallel to the coastline, ~1 m apart. From each quadrat, the largest *P. depressa* with the best-preserved shell was collected. The ML was measured, and the quadrat number of each shell recorded in the field. In the laboratory, the soft tissues were removed, the shells rinsed clean with fresh water and air-dried at ambient temperature before embedding in resin. A subgroup of five shells per location was selected for further age and growth analyses (total shells = 30; Table S2).

### Shell embedding and growth line analysis

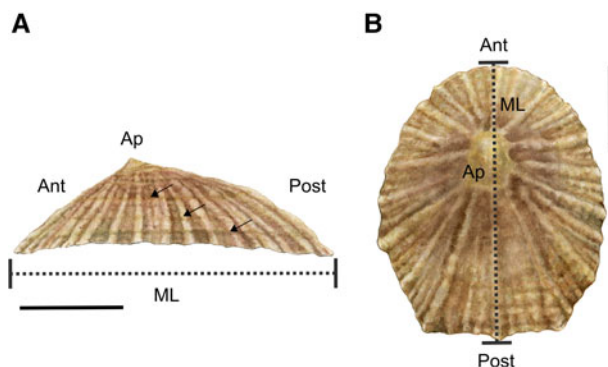
Shells collected were heavily eroded and thus many samples were excluded from the approach because of the difficulty of counting increments near the shell apex. Thus, 10 shells were used for annual line validation (see Table S1), and 30 shells for age and growth analysis (see Table S2). Each shell was embedded in epoxy resin (Kleer-Set Type FF, Polyester Casting Resin, MetPrep Ltd, UK). Embedded shells were processed using a standard procedure described by Ekaratne & Crisp (1982, 1984). The shells were sectioned using a Buehler ISOMET 5000

precision saw (cut rate  $14 \text{ mm min}^{-1}$  at 5000 rpm) along their maximum growth axis from the anterior to posterior side of the shell (see Figure 2B). One half of each resin block was polished using progressively finer grades of abrasive papers (P120, P400, P1200 and P1200/4000; MetPrep Ltd, UK) and finally polished using  $2 \mu\text{m}$  diamond paste (Maiapul Polishing Cloth Diamond, Spectrographic Ltd, UK). Polished shell sections were thoroughly cleaned using detergent and rinsed in tap water, etched for 30 s by immersion in 5% HCl, rinsed with distilled water and air-dried at ambient temperature for 24 h in a fume hood. Dry shell section surfaces were flooded with ethyl acetate and a  $0.35 \mu\text{m}$  thick sheet of acetate film (Replication Material G255, Agar Scientific Ltd, UK) applied to the etched shell surface, then air-dried at ambient temperature for 45 min (see also Richardson *et al.*, 1979). Dry acetate peels were gently removed from the polished shell section, trimmed, and mounted between microscope slides for visual analysis of shell increments under transmitted light microscopy.

High-resolution photographs ( $5\times$  magnification) of the entire shell section seen in reflected light or acetate peels viewed in transmitted light were taken using a Lumenera Infinity 3 camera (Infinity3-3URC 00199474, Canada) attached to a Meiji Techno Co. MT8100, Ltd, Japan microscope. ImagePro Premier® 9.1.4 (Built 5368, Media Cybernetics®) was used to generate photomontages and to catalogue each sample. Each peel (i.e. a replicate of each shell) was stored and referenced for further measurements.

### Identification, determination of periodicity and timing of growth line formation

Shell sections were observed in reflected light whilst acetate peels were viewed in transmitted light to highlight the structural features of the shells. Using both approaches, prominent growth lines were observed in the apex region of the shell and these



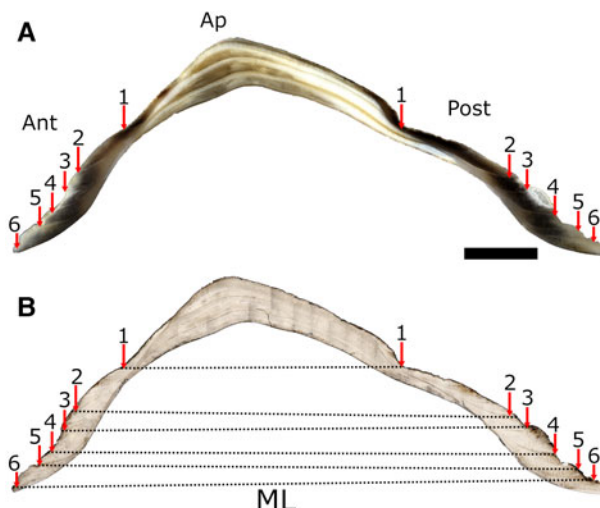
**Fig. 2.** Morphology of *Patella depressa*: (A) lateral and (B) dorsal views of the shell. Ant = anterior, Post = posterior and Ap = apex. Dashed black lines indicate maximum length (ML) of the shell. In (A) black arrows indicate potential annual growth rings or annuli. Scale bars: A–B, 10 mm.

lines were followed through into the anterior and posterior sides of the shell and the positions of the lines noted and labelled (i.e. line 1, line 2, line 3, etc.; see Figure 3). Only those lines that could be observed both in shell section and acetate peel and traced into both the anterior and posterior sides of the shell were considered to be ‘true’ lines (see Figure 3A, B). Similar growth lines in the apex of the shell of the related species *P. vulgata* have been validated as forming annually (e.g. Ambrose *et al.*, 2016).

To establish the periodicity of growth line formation in *P. depressa*, the position on the shell where they were initially measured at Shell Island (i.e. their initial ML in June 2015, see Table S1) was first identified in the acetate peels of the shell sections of the marked, measured and recovered limpets (N = 10). When the initial ML recorded at Shell Island was transposed onto the matching shell section–acetate peel (see Figure 3) it conveniently corresponded to a distinct line deposited when the shell was disturbed during labelling and initial measurement. The number of distinct growth lines deposited in the anterior and posterior sides of each shell was counted. Without exception, two lines corresponding to 2015–16 and 2016–17 were observed in all 10 shells, demonstrating unequivocally that the lines observed in the shell were formed annually. Timing of growth line deposition was established by transposing the ML measurements taken on the three other occasions in the field (see above), i.e. in August and November 2015, and in March 2016 onto the relevant shell section–acetate peel (Table S1). In all shells, the growth line in 2015–16 was deposited between the November 2015 and March 2016 measurements, indicating winter line deposition.

#### Growth rate estimations

Five *P. depressa* shells from each location were selected to study their shell growth (see above; Table S2). Annual lines in these shells were identified in both the anterior and posterior growing edges, counted to determine age and their positions marked on a photomicrograph image of the shell section–acetate peel. To check that the growth lines had been correctly identified in both growing edges, the cumulative distance between the shell apex and each annual line was measured in both growing edges of shells, from each population–region, and compared using a paired samples *t*-test (Gotelli & Ellison, 2013). No significant difference between the annual shell growth of the anterior and posterior growing shell-edges was observed (N Wales: *t*-value = -1.99, *df* = 143.85, *P* = 0.053; SW England: *t*-value = -1.87, *df* = 148.89; *P* = 0.062; S/SE England: *t*-value = -1.97, *df* = 143.88, *P* = 0.052), demonstrating that the positions of the annual lines had been identified consistently and correctly in both growing shell-edges.



**Fig. 3.** Photomicrograph of a cross-section of a six-year-old *Patella depressa* shell showing prominent (annual) lines (arrows) identified in both: (A) reflected light on the resin-embedded shell and (B) transmitted light through an acetate peel. From the apex (Ap) to the shell margin, increment number 6 indicates the last major growth line observed in (A) and (B). Growth rates were calculated by measuring the maximum length (ML, dotted line) at each annual line. Scale bar: A & B, 4 mm.

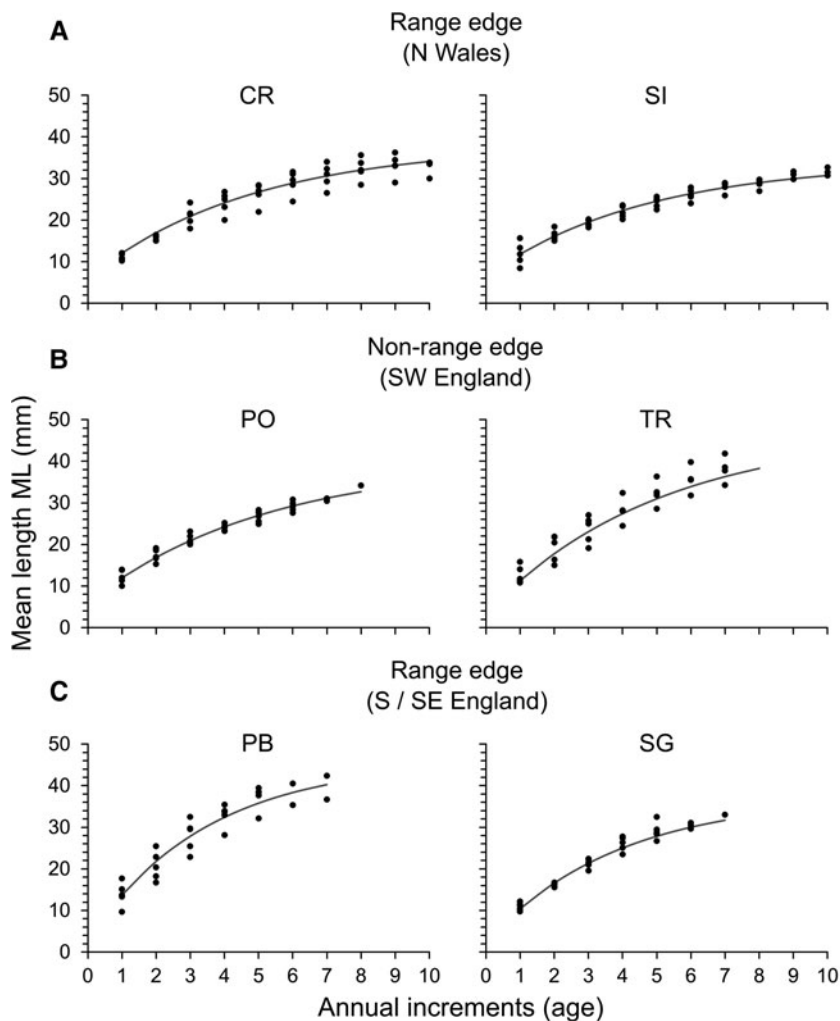
To evaluate if the number of annual lines (i.e. limpet age) varied between the two range-edges and non-range-edge populations of *P. depressa*, analysis of variance with two factors was performed (ANOVA; Gotelli & Ellison, 2013). The two factors included were: (1) Region as a fixed factor, with three levels: N Wales, SW England and S/SE England; and (2) Location as a random factor, nested within Region. Subsequently maximum length (ML) between the anterior and posterior growing edges at each growth line was measured directly from the shell section–acetate peels and/or photomicrographs (Figure 3). Von Bertalanffy growth (VBG) curves were fitted to the mean ML at age data for each limpet population at each location. The VBG parameters were estimated following the equation defined by:

$$L_t = L_\infty(1 - e^{-K(t-t_0)})$$

where  $L_t$  is the length at time  $t$ ,  $L_\infty$  is the theoretical ML that species would reach, the  $K$  parameter is a growth coefficient estimating how fast the individual approaches  $L_\infty$  and  $t_0$  is the theoretical age at zero length. Ford–Walford plots were used to estimate  $L_\infty$  and  $K$  (King, 2007). A Ford–Walford plot shows the linear relationship between  $L_t$  against the length at  $t + 1$  ( $L_{t+1}$ ). From this relationship,  $L_\infty$  and  $K$  can be calculated from the straight-line equation where  $L_\infty = y - \text{intercept}/(1 - \text{slope})$  and  $K = -\ln(\text{slope})$  (King, 2007). The remaining parameter in the von Bertalanffy growth equation,  $t_0$ , can be estimated if length at a particular annual line is known (King, 2007). Therefore, from the von Bertalanffy equation,  $t_0$  may be calculated as follows:

$$t_0 = t + (1/K) \times (\ln [(L_\infty - L_t)/L_\infty])$$

In addition, comparisons based on these parameters between locations and regions were made. Furthermore, as in fishes and invertebrates, whose growth can also be described by the von Bertalanffy function, comparisons were also made through the overall growth performance index, i.e.  $\Phi' = \log K + 2 \log L_\infty$  (see Clarke *et al.*, 2004; Pörtner *et al.*, 2005 for details). According to Pauly (1979),  $\Phi'$  describes the growth rate at the point of inflection of the von Bertalanffy growth curve (i.e. maximum growth rate; Heilmayer *et al.*, 2004). The TropFishR package was used to build Ford–Walford plots and estimate von Bertalanffy



**Fig. 4.** Von Bertalanffy growth (VBG) curves fitted to maximum length (ML) at age-data for shells of *Patella depressa* from two range edges populations: (A) N Wales (CR = Criccieth and SI = Shell Island) and (C) S/SE England (PB = Portland Bill and SG = Swanage); and (B) from non-range-edge populations in SW England (PO = Polzeath and TR = Trevone). See Table 1 for parameters of each VBG curve.

parameters in the CRAN R project environment (R Core Team, 2019, v3.5.3; Mildenerger *et al.*, 2017).

#### Effect of limpet density on shell growth

Variation in the growth coefficient (VBG-K) among regions, was determined using analysis of covariance (ANCOVA; Gotelli & Ellison, 2013), with two factors: Region (three levels: N Wales, SW England and S/SE England) and Location (two levels) nested in Region, and a covariate of total limpet density, i.e. total limpet number recorded in the individual quadrats where shells were collected for age and growth analysis (June–July 2016; see above). Density-dependent competition, both inter-specific (based on the density of *P. vulgata*) and intra-specific effects (based on the density of *P. depressa*) was examined by determining the relationship of density with the VBG-K of each individual ( $N = 5$ ) collected at each location. In addition,  $R^2$  was calculated for each relationship between individual  $K$  values of *P. depressa* and the total density of limpet species (i.e. total limpets, see above) and the density of *P. vulgata* and *P. depressa* when considered singly (Gotelli & Ellison, 2013). Bartlett's tests were used to check the normality of the residual variances before using both ANOVA and ANCOVA (Gotelli & Ellison, 2013). Tukey's post hoc tests were carried out for pairwise comparisons.

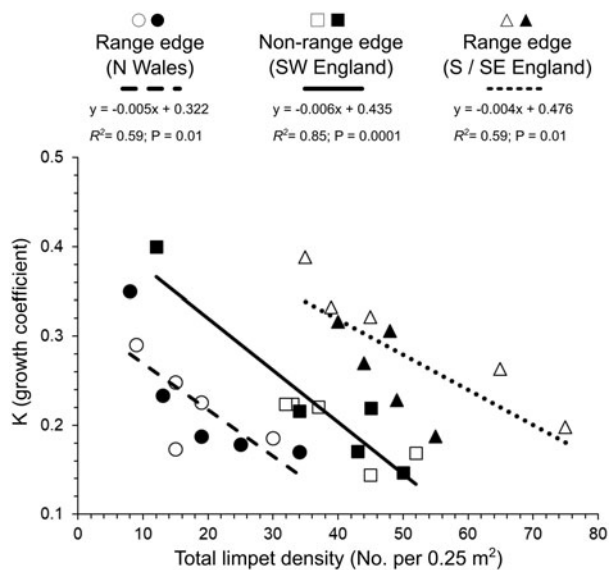
#### Results

Under low power magnification all the sectioned limpet shells revealed pronounced annual growth lines in both reflected light

on the shell sections and in transmitted light through the acetate peels (Figure 3A, B). Viewed at high magnifications many fine bands were observed in the acetate peels between the obvious strongly defined annual lines (Figure 3B). In some limpet species these fine bands are deposited tidally (see Ekaratne & Crisp, 1984; Crisp *et al.*, 1990; Richardson & Liu, 1994). In *P. depressa* shell sections, the low-contrast, weakly defined bands radiate from the apex towards the edges of the shells (Figure 3A). The separation between these bands becomes reduced as they gradually merge to form a single high-contrast growth line. From the seasonal shell samples it was possible to constrain the timing of line formation to the winter period between November 2015 and March 2016 when shell growth ceased. This growth cessation is characterized by a change in the orientation of the micro-growth bands towards the shell surface (Figure 3B). A similar pattern has been observed in *P. depressa*, and in the top shell *P. lineatus* (see García-Escárcaga *et al.*, 2019, 2020).

#### Age patterns in *P. depressa*

Shells displayed significant geographic differences in the number of annual lines among the two range-edges and non-range-edge populations (two-way ANOVA:  $F_{(2,24)} = 13.8$ ,  $P < 0.05$ ). Limpets collected from the poleward edge in N Wales had a significantly greater number of annual lines (between nine and ten lines; see also growth curves in Figure 4), when compared with individuals at non-range-edge populations in SW England (six and eight lines per shell) and those at the range-edge populations in S/SE England (between six and seven lines per shell; Tukey's post hoc test,  $P < 0.05$ ).



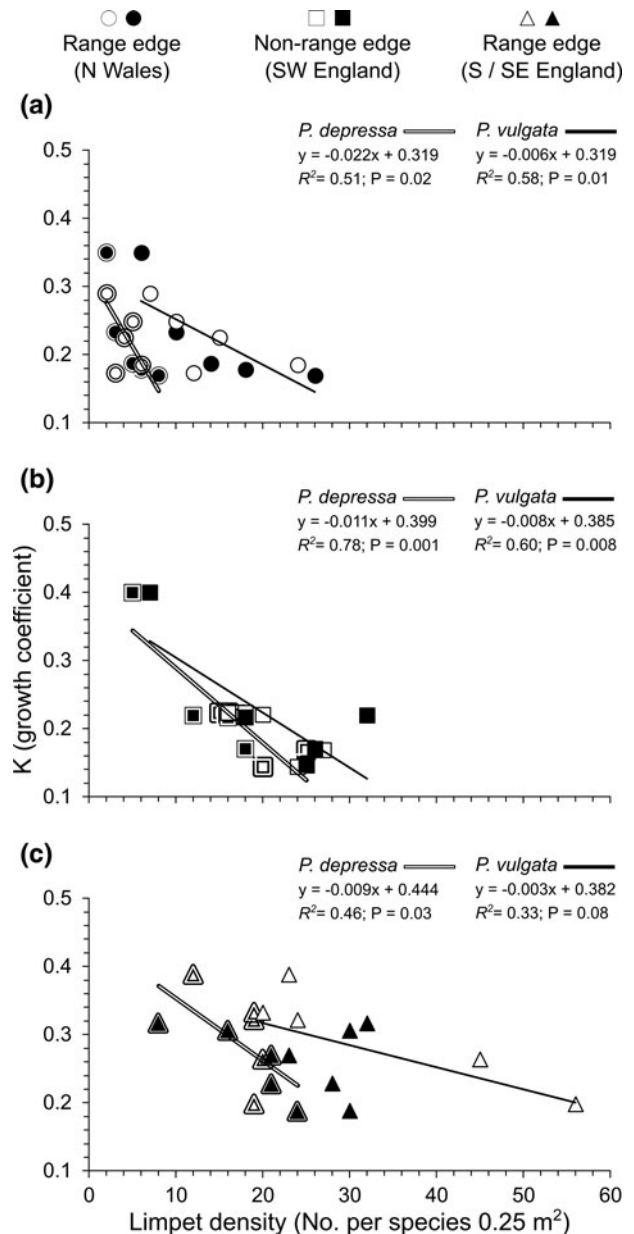
**Fig. 5.** Relationship between individual *Patella depressa* growth coefficients ( $K$ ) and total limpet density (combined number of *P. depressa* and *Patella vulgata* in  $0.25 \text{ m}^2$ ) measured in June–July 2016 at each region, defined by two range edges of *P. depressa*: N Wales (Criccieth =  $\circ$ ; Shell Island =  $\bullet$ ) and S/SE England (Portland Bill =  $\triangle$ ; Swanage =  $\blacktriangle$ ); and by non-range-edge populations in SW England (Polzeath =  $\square$ ; Trevone =  $\blacksquare$ ). Linear regression lines fitted to the data and  $R^2$  are indicated for each region.

### Growth rate in *P. depressa*

The 18 tagged and measured *P. depressa* from Shell Island (N Wales) displayed an average increase in ML of  $\sim 6$  mm over the 2-year period in the field (2015–17). The Von Bertalanffy growth (VBG) plots of limpets showed different growth patterns between the two range-edge populations (Table 1; Figure 4A–C). *Patella depressa* from S/SE England (Figure 4C) had higher  $K$  values and higher growth performance indices ( $\mathcal{O}$ ) compared with shells from N Wales (Table 1, Figure 4A). Rapid growth was identified in shells collected from range-edge populations in S/SE England, where most individuals reached over 30 mm in ML by the fourth or fifth annual line at Portland Bill and by the sixth line at Swanage (Figure 4C). By contrast, shells collected from the poleward populations in N Wales showed slower shell growth to reach their asymptotic maximum (i.e. flatter growth curves) with most shells only reaching a ML  $> 30$  mm between the seventh and tenth annual line (Figure 4A). Individuals collected from the non-range-edge populations (Figure 4B), showed both faster (i.e. Trevone) and slower (i.e. Polzeath) annual shell growth than individuals from both range edges, where individuals reached over 30 mm ML in 5 or 6 years with the greatest  $L_\infty$  in shells from Trevone (Table 1).

### Density-dependent effects on limpet shell growth

At the quadrat scale ( $50 \times 50 \text{ cm}$ ), the variation in the growth coefficient ( $K$ ) was significantly negatively affected by the combined total density of *P. depressa* and *P. vulgata* present (ANCOVA:  $F_{(1, 18)} = 62.2$ ,  $P < 0.05$ ; Figure 5). *Patella depressa* with higher  $K$  values were found amongst lower limpet densities across range-edge and non-range-edge populations (Figure 5).  $K$  values also differed significantly among regions (ANCOVA:  $F_{(2, 18)} = 10.7$ ,  $P < 0.05$ ; Figure 5), but the interaction between limpet density and region was not significant ( $P > 0.05$ ). Furthermore, inter- and intra-specific relationships across sites indicated that  $K$  values of *P. depressa* shells were more negatively affected by *P. depressa* density than *P. vulgata* density (Figure 6). High intra-specific effects amongst *P. depressa* were observed across all the



**Fig. 6.** Negative relationships between individual *Patella depressa* growth coefficients ( $K$ ; calculated from five shells) and *P. depressa* density (double black outline on symbols,  $N = 10$ ) and *Patella vulgata* density (single black outline on symbols,  $N = 10$ ) at each location. Limpet density was measured in June–July 2016 at each region, defined by two range edges of *P. depressa*: (A) N Wales (Criccieth =  $\circ$ ; Shell Island =  $\bullet$ ) and (C) S / SE England (Portland Bill =  $\triangle$ ; Swanage =  $\blacktriangle$ ); and by (B) non-range-edge populations in SW England (Polzeath =  $\square$ ; Trevone =  $\blacksquare$ ). Linear regression lines fitted to the data and  $R^2$  are indicated.

selected regions, especially in the range-edge populations in N Wales ( $R^2 = 0.51$ ;  $P = 0.02$ ; Figure 6A) when compared with the range-edge in S/SE England ( $R^2 = 0.46$ ;  $P = 0.03$ ; Figure 6C). Although the relationship between  $K$  values of *P. depressa* with density of *P. vulgata* was not significant in range-edge populations in S/SE England ( $R^2 = 0.33$ ;  $P = 0.08$ ; Figure 6C), it was in N Wales ( $R^2 = 0.58$ ;  $P = 0.01$ ; Figure 6A), with both species showing strong and similar intra- and inter-specific relationship at non-range-edge populations in SW England ( $P < 0.05$ ; Figure 6B).

### Discussion

Our results showed that age, indicated by annual lines in shells, and growth performance, determined by their  $K$  values (growth

**Table 1.** Summary of calculated von Bertalanffy growth (VBG) parameters (i.e.  $K$ ,  $L_{\infty}$  and  $t_0$ ) using Ford–Walford plots for five shells selected in June/July 2016 at each location

Region	Location	$L$ max (mm)	$L_{\infty}$ (mm)	$K$ (years <sup>-1</sup> )	$t_0$	$\emptyset'$
Range edge (N Wales)	CR	36.4	38.1	0.207	-0.849	2.28
	SI	33.9	34.0	0.214	-0.994	2.29
Non-range edge (SW England)	PO	36.5	40.1	0.191	-0.866	2.49
	TR	43.1	47.3	0.198	-0.381	2.65
Range edge (S/SE England)	PB	41.5	45.9	0.299	-0.224	2.80
	SG	34.5	37.4	0.269	-0.247	2.58

There are three regions, defined by two range edges of *Patella depressa*: N Wales and S/SE England; and by non-range-edge populations in SW England.  $L_{\infty}$  (mm) is the theoretical maximum length that species can reach;  $K$  (year<sup>-1</sup>) is a measurement of the rate at which the maximum size can be reached; and  $t_0$  is the theoretical age at zero length. Maximum length ( $L$  max) and Growth Performance index, i.e.  $\emptyset' = \log K + 2 \log L_{\infty}$  are also indicated (see Clarke *et al.*, 2004 for details). The VBG curves are indicated in Figure 4.

coefficients) differed between the two range-edges of distribution of *P. depressa*. Our findings do not support the hypothesis that the two separate boundaries of *P. depressa* have older individuals with slower growth rates than non-range-edge populations. Individuals of *P. depressa* from the range-edge in S/SE England grew faster and had fewer annual growth lines than individuals from the poleward edge in N Wales. This suggests rapid annual growth rates in shells collected from the range-edge in S/SE England, indicating less long-lived individuals than individuals in N Wales. In addition, there was no clear variation in age and growth patterns between non-range-edge populations in SW England and those at either of the range edges. These observations suggest that both range-edge and non-range-edge populations of *P. depressa* could be characterized by different individual performance traits, which can be imprinted in shells as growth records (García-Escárczaga *et al.*, 2020). Furthermore, density-dependent processes negatively affected individual growth performance ( $K$  values) of *P. depressa* at the scale of the habitat-patch (i.e. individual quadrats).

#### Shell growth of *P. depressa* at poleward boundaries

Our study has shown that shell growth of *P. depressa* has different patterns at their two leading poleward edges of distribution that occur within Britain. This can be explained by regional differences in seawater temperature (Hiscock *et al.*, 2004) and inter-individual variations in the duration and intensity of both growth cessation and slow-down periods because of reproduction (e.g. see Gutiérrez-Zugasti *et al.*, 2017 for *P. vulgata*; García-Escárczaga *et al.*, 2020 for *P. depressa*). Regional differences in sea surface temperatures suggest that faster growth rates recorded in limpets at the range edge of *P. depressa* in S/SE England may be a consequence of warmer summer temperatures when compared with conditions in N Wales (Oróstica *et al.*, 2020), enabling early onset of the growth season and longer duration into the autumn. In addition, there is evidence of earlier onset of maturation in *P. depressa* during early spring and a prolonged spawning season into the autumn in recent years as a response to global warming (Orton & Southward, 1961; Moore *et al.*, 2011). The complex interaction between the energetic cost of gonad maturation (Blackmore, 1969; Wright, 1977; Wright & Hartnoll, 1981), seasonal foraging activity patterns and microbial food availability (Jenkins *et al.*, 2001; Thompson *et al.*, 2004) need further exploration throughout the range of limpets and other gastropod species.

Shells from non-range-edge populations in SW England showed both faster and slower annual growth patterns than individuals from both range-edges of *P. depressa*, depending on the specific location. In this context, sclerochronological approaches

in shells have shown that the patterns of annual lines can vary even within individuals of the same species (see Surge *et al.*, 2013 for *P. vulgata*; García-Escárczaga *et al.*, 2020 for *P. depressa*). For instance, Surge *et al.* (2013) found a mix between summer and winter annual lines in *P. vulgata* shells from the English Channel, where the biogeographic boundary between the cold- and warm-temperate regions can be found (Hiscock *et al.*, 2004). Conversely, in regions with cooler sea temperatures, shells have a slow growth rate and form a prominent annual line during winter (Fenger *et al.*, 2007; Surge *et al.*, 2013). A similar pattern has been found for *P. depressa* in northern Spain (nearer the centre of its geographic distribution), where individuals had growth cessations in winter, during the coldest months of the year, and occasionally also in summer, perhaps associated with strong upwelling (García-Escárczaga *et al.*, 2020). Our annual growth line validation using tagged individuals of *P. depressa* from N Wales agrees with these findings and multiple shell measurements at intervals throughout the year indicated that the annual line formed at the end of the year (i.e. winter, between November 2015 and March 2016, in N Wales, UK), when the seawater temperatures are at their coldest (see Oróstica *et al.*, 2020, their fig. 2). By contrast, individuals of *P. vulgata* inhabiting the north of Spain (toward the equatorward range-edge of its geographic distribution), form a noticeable annual line in mid-summer, presumably due to heat stress (Surge *et al.*, 2013). Further growth analysis using sclerochronological techniques on shells of *P. depressa* could elucidate annual growth line patterns across poleward edge, central and equatorward edge populations.

#### Age pattern of *P. depressa* at poleward boundaries

Populations of *P. depressa* were shown to have different age patterns at their two leading edges. Further south, in central populations of *P. depressa* (Silva *et al.*, 2003), growth rates of individuals were slightly higher compared with those individuals at higher latitudes (Oróstica *et al.*, 2020), which could suggest a shorter lifespan for those individuals in Portugal (Guerra & Gaudencio, 1986; Lewis, 1986). However, the values of growth parameters ( $K$  and  $\emptyset'$ ) we found were lower when compared with other patellean species (i.e. *Nacella*, *Scutellastra*, *Cymbula* and *Acmaea*; see Branch, 1981; Clarke *et al.*, 2004 for review). Northern populations of southern species are characterized by short reproductive periods in summer and frequent recruitment failures (Bates *et al.*, 2014). This can generate a latitudinal pattern, whereby marginal regions have an irregular limpet age structure with missing year classes and dominated by older and larger individuals (Lewis *et al.*, 1982; Lewis, 1986). Analysis of the age of *P. depressa* suggests that the patterns obtained here are not that far from those calculated for other limpet populations in Britain (Lewis &



Bowman, 1975; Wright, 1977), supporting a gradient of increasing age of *P. depressa* from Portugal (~3 years old; Guerra & Gaudencio, 1986; Lewis, 1986), to southern England (6–8 years old, this study) and N Wales (9–10 years old, this study). Studies on the northern cold-water species *P. vulgata* estimated a maximum lifespan in the Isle of Man (UK) of between 12 and 17 years (Wright, 1977), and between 15 and 17 years in north-east England (Lewis & Bowman, 1975). By contrast, further south at its equatorward limit in Portugal, size-frequency data suggest that *P. vulgata* has a lifespan of no more than 3 or 4 years (Guerra & Gaudencio, 1986), indicating that latitudinal effects can influence population age-structure in limpet species (Lewis & Bowman, 1975; Wright, 1977; Lewis et al., 1982; Guerra & Gaudencio, 1986; Lewis, 1986).

### Density-dependent effects on shell growth of *P. depressa*

In reconstructing shell growth measurements from the annual growth lines we have clearly shown that an increase in total limpet density negatively affected the growth of *P. depressa*. This pattern was supported by the negative relationship between growth performance (*K* values) of *P. depressa* and total limpet density (*P. depressa* plus *P. vulgata*). There are contrasting effects of limpet density on shell growth performance based on the scale used across range-edge and non-range-edge populations (see Oróstica et al., 2020 for details). This is the first time that this population process has been observed at such small spatial resolution in *P. depressa* from unmanipulated local areas (but see Boaventura et al., 2002, 2003 and Moore et al., 2007 for experimental studies). At a quadrat scale, we found localized and negative effects of limpet density on growth performance. In addition, greater intra-specific effects on growth performance between *P. depressa* individuals were detected, compared with the inter-specific effects with *P. vulgata* density. There was, however, a stronger inter-specific effect of *P. vulgata* on the growth of *P. depressa* in N Wales than at the eastern range-edge in S/SE England. Previous experimental studies show a similar pattern (see Boaventura et al., 2002, 2003 and Moore et al., 2007; Firth & Crowe, 2008, 2010). Lewis & Bowman (1975) for example suggested that intra-specific competition in *P. vulgata* arises in areas with densities above 300–450 limpets m<sup>-2</sup>, which are likely to reduce individual growth rates (Branch, 1981). These patterns suggest that growth rates in limpet species are highly variable, subject to abiotic and biotic factors (Branch, 1981) at various geographic and temporal scales as well as human impacts (Fenberg & Roy, 2012; Borges et al., 2016; Martins et al., 2017), with local processes being important to consider at species range edges (Helmuth et al., 2006).

### Conclusions

Our study has measured individual growth rates and determined age using validated annual line deposition in range-edge populations of *P. depressa*. Von Bertalanffy growth parameters indicated that range-edge individuals in S/SE England performed better (i.e. grew faster) than poleward individuals in N Wales. In addition, annual growth line analysis in *P. depressa* suggests a population structure consisting of mainly older individuals with slower growth rates in poleward N Wales populations than in range-edge populations in S/SE England. Limpet density at the quadrat scale influenced individual growth performance, with intra-specific effects more important than inter-specific effects. Our field validation of annual line formation in shells of *P. depressa* has suggested a winter season of formation (i.e. November to March) in populations at their northern boundaries (i.e. N Wales). However, a better understanding of the timing of annual growth

line formation in *P. depressa* over longer temporal as well as larger spatial scales could disentangle the mechanisms that underpin differences in shell growth and age structure seen in this study.

**Supplementary material.** The supplementary material for this article can be found at <https://doi.org/10.1017/S0025315421000539>

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