Physiological aging in the Icelandic population of the ocean quahog Arctica islandica

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ABSTRACT: The ocean quahog Arctica islandica is one of the longest-living and slowest growing marine bivalves. The oldest specimens obtained for the present study approached 200 yr. To achieve such a long lifespan, accumulation of oxidative damage markers in tissues must ideally be maintained at low levels over time, because the accumulating debris disturbs cellular functions. We investigated shell growth and cellular aging in an Icelandic population of A. islandica. Specifically, we analyzed protein carbonyl concentration as a marker for the oxidative deterioration of tissue proteins, and the accumulation of the fluorescent age pigment lipofuscin over quahog lifetime in gill, mantle and adductor muscle. The very slow growth rates of A. islandica correlate with very efficient maintenance of body proteins compared to other, faster aging bivalves. Lipofuscin granules accumulated mainly in connective tissues of gill and mantle. Lowest lipofuscin accumulation was found in the adductor muscle, and there, only outside the myofibrils. Consistent with the pleiotropic theory of aging, A. islandica seems to trade slow growth and late onset of reproduction for a very efficient autophagic potential that mitigates oxidative damage accumulation and supports long lifetime and presumably reproduction in very old ocean quahog.

KEY WORDS: Arctica islandica · Ocean quahog · Growth · Aging · Lipofuscin · Protein oxidation

INTRODUCTION

Marine bivalves, especially from cold temperate and polar seas, can reach outstandingly long life spans (>300 yr) among aquatic ectotherms (Brey 1999). Finch (1990) put such species in a ‘negligible aging’ senescence category, indicating that proliferation continues slowly and persistently in most tissues until very old age. Variable environmental stress levels (temperature, light, food availability) affecting populations from different geographical locations, or differing levels of stress that individuals in these populations experience through predator attack or competition, can stochastically modify the process of aging in bivalves and populations, and differentially shorten or lengthen individual life expectancy. Thus, populations within a stressful environmental setting can exhibit a life expectancy distinctly lower than mean longevity in the species. In fact, once a species’ longevity under optimal conditions is known, mean life expectancy of individuals in that population can serve as an indicator of the level of stress being experienced. This, and the fact that shells of long-lived cold-temperate bivalves are increasingly used as environmental archives (Richardson et al. 2001, Schöne et al. 2005), have raised interest in bivalve life strategies, as well as in the physiological changes in cells and tissues of aging bivalves. Recent work from our laboratory (Philipp et al. 2005a,b, Philipp et al. 2006) indicates that physiological aging in marine mud-dwelling bivalves is in line with the ‘Free Radical - Rate of Living’ theory established by Pearl (1928) and Harman (1956). This theory links the process of cellular aging to the rate of metabolic oxygen consumption that, in turn, determines the rate of mitochondrial release of hazardous reactive oxygen species (ROS) and shortens bivalve life expectancy. As in mammals (Terman & Brunk 2005), accumulation of free radical damage markers is viewed as the major cause of aging in bivalves. In contrast, higher levels of enzymatic antioxidants and free radical scavengers, such as glutathione, support longer species life expectancy (Philipp et al. 2005a).
We investigated physiological aging of the mud clam *Arctica islandica* (ocean quahog), one of the longest-living and slowest growing of the marine bivalves. Individuals over 100 yr old are commonly found, and an outstanding maximum life span of >300 yr has been reported (Schöne 2003, Schöne et al. 2004). The extraordinary longevity and wide geographical distribution of *A. islandica* render it an interesting species for ecophysiological studies into the relative importance of extrinsic (environmental, e.g. temperature) and intrinsic (e.g. genetic or behavioural) factors in modifying animal physiology and altering the velocity of aging in ectotherms. In this paper, we present the first set of data from a study of physiological aging in *A. islandica*. Age-dependent oxidative damage of proteins (protein oxidation) and accumulation of lipofuscin (end product of lipid peroxidation, also called 'fluorescent age pigment') in gill, mantle and adductor muscle of young (5 to 11 yr) and old (110 to 192 yr) *A. islandica* individuals from a cold-boreal North Icelandic population were measured. Both parameters are indicative of cellular maintenance through autophagic activity (i.e. autodigestion processes), and lipofuscin accumulation in connective tissues has previously been shown to correlate with age in long-lived bivalves (Lomovasky et al. 2002). Counting the annual internal growth bands in the shell of each individual, we were able to distinguish different ages among similarly sized individuals (Thompson et al. 1980a). We were interested to know if keeping the levels of oxidative damage products low forms part of the strategy employed by the ocean quahog to enable such extreme longevity, and whether or not there are tissue-specific differences in the aging process.

**MATERIALS AND METHODS**

**Quahog collection and maintenance.** Ocean quahog *Arctica islandica* were collected in August 2004 and in May 2005 northeast of Iceland (66° 01′ 54″ N, 14° 50.98′ W) between 14 and 22 m water depth using a hydraulic dredge. Water temperature on the sampling date was 4°C in May 2005 and 9°C in August 2004. The quahog were transported in cooling boxes to the Sandgerdi Marine Centre, Iceland, where they were kept at constant temperature (9°C) and salinity (34.9 psu) in 400 l tanks for 7 d. On Day 7, quahog were dissected and gill and mantle were snap frozen in liquid nitrogen for biochemical analysis. Protein oxidation and lipofuscin were analyzed only in quahog sampled in 2005. Tissues were frozen for determination of protein carbonyl content and samples of gill, mantle and adductor muscle were fixed in Bakers formalin for 24 h and stored in gum sucrose until further processing for histological determination of lipofuscin granule accumulation. Shells of all individuals from 2004 and 2005 were cleaned and numbered for later age determination and assignment to the samples for biochemical analysis. Samples were transported to the Alfred-Wegener-Institute in Bremerhaven, Germany, for analysis.

**Determination of individual age and growth.** Individual age was determined from internal shell growth bands in 179 sampled shells, collected in August 2004 and in May 2005, following the procedure of Mutvei et al. (1994) and Schöne et al. (2004). The right shell valve was embedded in epoxy resin and sectioned along the axis of maximum shell growth (height *H*) with a Buehler low-speed diamond saw. Cross-sections were ground on lapidary wheels using grits of P400, P1200, P2400 and P4000 grade and subsequently polished using a polycrystalline diamond suspension of 1 and 0.1 μm. Annual shell growth bands (see Schöne et al. 2004 for verification of annual band formation) were counted using a stereomicroscope at 10 to 80× magnification.

We fitted the general growth model of Schnute (1981) to the size-at-age data (*S* <sub>i</sub> versus age *t*) by means of the nonlinear iterative Newton algorithm:

\[
S_t = \left( Y_2^B + (Y_1^B - Y_2^B) \times \left[ \frac{1 - e^{-A \times (t-t_1)}}{1 - e^{-A \times (t-t_2)}} \right] \right)^{1/B}
\]

The model has 2 constants: A (time<sup>−1</sup>), and B (dimensionless); and the size parameters *Y*<sub>1</sub> (mm) and *Y*<sub>2</sub> (mm). The 2 age values, *t*<sub>1</sub>, and *t*<sub>2</sub>, are chosen by the user, e.g. youngest and oldest individual in sample. The starting values of *Y*<sub>1</sub> and *Y*<sub>2</sub> are set accordingly. Special cases of the Schnute model resemble historical growth models such as the von Bertalanffy, Richards, or Gompertz models. For our study, the Schnute model indicated that the von Bertalanffy growth model fitted the data best:

\[
H_t = H_\infty \times (1 - e^{-K \times (t-t_0)})
\]

where *H*<sub>\infty</sub> = asymptotic shell height, *K* = growth constant, *t* = age, and *t*<sub>0</sub> = age at which height is zero.

**Determination of protein carbonylation.** Protein carbonylation in 16 young (5 to 7 yr) and 17 old (110 to 192 yr) individuals was measured after Levine et al. (1990). For this analysis, 150 to 300 mg tissue were homogenised in 1 ml of 50 mM HEPE homogenization buffer, containing 125 mM KCl, 1.25 mM EDTA, 0.6 mM MgSO<sub>4</sub> and protease inhibitors (0.5 μg ml<sup>−1</sup> leupeptin, 0.7 μg ml<sup>−1</sup> pepstatine, 40 μg ml<sup>−1</sup> phenylmethylsulfonyl fluoride, 0.5 μg ml<sup>−1</sup> aprotinin, 0.44 mg ml<sup>−1</sup> EDTA, 160 μg ml<sup>−1</sup>MgSO<sub>4</sub> × 7H<sub>2</sub>O, pH 7.4), using a glass homogenizer (IKA Eurostar) at 300 rpm. Subsequently, 4 ml homogenization buffer was added and the homogenate was centrifuged at 31,000 × *g* (Ultra-centrifuÅget, Beckmann L7-80) for 22 min at 4°C. Two replicates of 0.4 ml each from the supernatant
were incubated with 1.4 ml of acidic DNP (10 mM 2,4-dinitrophenylhydrazine in 2 M HCl) at room temperature for 1 h to let DNP bind to the carbonyl groups. Blanks were run with HCl only. Following the incubation, protein was precipitated with 0.2 ml 100% trichloroacetic acid (TCA) and centrifuged for 5 min at 11 000 × g. The protein pellet was washed 3 times with ethanol:ethylacetate (1:1) and the final pellet left to dry out completely before being re-dissolved in 0.6 ml of guanidine hydrochloride (6 M in 20 mM potassium phosphate buffer at pH = 2.5), the carbonyl content in each sample and blank was measured spectrophotometrically at 360 nm (molar extinction coefficient ε = 22 000 M⁻¹ cm⁻¹). To relate carbonyl content to the amount of extracted proteins in each sample, the protein content was subsequently measured using Bradford’s (1976) method.

**Histological investigation of lipofuscin accumulation.** Accumulation of lipofuscin was assessed histologically in 8 young (5 to 11 yr) and 8 old (110 to 192 yr) individuals. Samples of 5 × 5 × 5 mm of gill, mantle and adductor muscle were transferred to histocettes and fixed in Bakers formalin (100 ml formaldehyde 40% and 20 g Ca(CH₃COO)₂ in 1 l distilled water) for 24 h. Subsequently, samples were transferred to a gum-sucrose solution (300 g sucrose and 10 g gum Arabic in 1 l distilled water) and transported to the Alfred-Wegener-Institute in Bremerhaven, where samples were maintained at 4°C until further processing.

For embedding, samples were washed twice with 0.2 M phosphate buffer (pH 7.4) for 30 min. Samples were dehydrated by 2 × 15 min in 70% and 2 × 15 min in 100% acetone immersions. Each sample was then transferred into 5 ml monomer solution (80 ml 2-hydroxyethyl-methacrylic acid, 12 ml 2-butoxyethanol, 0.27 g benzoylperoxide) and left for 12 h. Subsequently, samples were transferred to an embedding form and 0.1 ml of polymerization activator solution (polyethylene glycol 200 and N,N-dimethyamine, 10:1) were added to each 5 ml of monomer solution. Samples were left to polymerise at 4°C for 12 h and then dried 24 h in the fume hood before being cut into 2 μm slices with a rotation microtome (Leica, RM 2145). Lipofuscin was detected by Schmorl-staining modified after Pearse (1960). Sections of different tissues were washed for 10 min in distilled water and stained for 10 min in a mix of 100 ml of solution A (500 mg iron-(III)-chloride in 50 ml distilled water) and 100 ml of solution B (500 mg potassium-iron-cyanide in 50 ml distilled water). Then, sections were placed for 2 min in 1% acetic acid, washed for 10 min under running water, and then washed 3 times in distilled water. Subsequently, sections were dried, fixed in glycerine gelatine (Merck, Microscopy Kaiser glycerine gelatine) and analyzed by a Zeiss Axioscope light microscope equipped with a KS300 Zeiss image analysis software. The percentage of tissue area occupied by lipofuscin granules was assessed.

**Statistics.** Protein oxidation data were tested for normality (Kolmogorov-Smirnov test) and homogeneity of variances (Bartlett’s test). The effects of tissue type and age on the lipofuscin content were tested by full interaction analysis of covariance (ANCOVA) using the statistical programme package JMP 5.0.1 (SAS), i.e. ln(lipofuscin content) versus tissue type (gill, mantle, muscle) and covariate ln(age).

**RESULTS**

**Individual growth**

Individual growth is best described by the von Bertalanffy growth model (Fig. 1):

\[
H_t = 86 (1 - e^{-0.057(t + 0.399)}), \quad N = 179, \quad r^2 = 0.933
\]

Yearly growth increments were >1 mm yr⁻¹ in young quahog (<12 yr), but <0.5 mm yr⁻¹ at the age of 40 yr.

**Age dependence of protein carbonylation (CO group formation)**

Young quahog (5 to 8 yr, mantle N = 10, gill N = 7) had (mean ± SD) 0.49 ± 0.19 nmol CO mg⁻¹ protein in mantle and 0.96 ± 0.49 nmol CO mg⁻¹ protein in gill tissue. Old individuals (110 to 192 yr, mantle and gill N = 8) had a mantle content of 0.58 ± 0.46 nmol CO mg⁻¹ protein and a gill content of 1.14 ± 0.54 nmol CO mg⁻¹ protein. CO content did not differ significantly between age groups or tissue types (p > 0.05).
Lipofuscin accumulation

Lipofuscin content was measured in gill, mantle and adductor muscle. Accumulation of lipofuscin in mantle (Fig. 2a,d) and gill (Fig. 2b,e) sections was not homogenous, but granules were concentrated in the connective tissue below the epidermis. The percentages (see Fig. 3) refer to only those areas of the tissues that contained granules (i.e. epidermis was not included). Adductor muscle was cut vertically to the fibres, and lipofuscin was encountered only in the sarcoplasmatic interstice (Fig. 2c,f) between myofibrils. We found that individual muscle fibres increased in size from 15 × 5 μm in young to 20 × 15 μm in older specimens. Histological evaluation of the density of lipofuscin granules in muscle was done for the interstitial compartment between the myofibrils which comprised <30% of the whole tissue in young and old quahog.

Lipofuscin content was significantly different in all 3 tissues (gill > mantle > muscle) and increased significantly over bivalve age (Fig. 3). The rate of increase with age did not differ between tissues (no significant interaction term, p = 0.117, Fig. 3): ln(lipo) = −3.070 + 0.657 × ln(age) + 1.134 × Dgill + 0.485 × Dmantle −

Fig. 2. Arctica islandica. Tissue slices from (a,d) mantle, (b,e) gill and (c,f) adductor muscle, embedded in gum sucrose and Schmorl-stained to visualize lipofuscin granules as irregular dark blue spots (= black spots in panels above). (a,b,c) Young quahogs (7 yr) vs. (d,e,f) old quahogs (122, 148 and 149 yr, respectively). Scale bars: 20 μm
1.919 × Dmuscle; N = 48, F = 79.44, all p < 0.001, where Dgill, Dmantle and Dmuscle represent ‘dummy’ variables (1 = this tissue, 0 = not this tissue).

Gills of old quahog (>100 yr) had the highest density of lipofuscin granules, followed by the mantle tissue, whereas lipofuscin accumulation in muscle was limited to the connective tissue between myofibrils. Even though only this part of the tissue was analyzed, the density of lipofuscin was low in muscle slices, but the rate of accumulation was the same as in mantle and gill.

DISCUSSION

In the present study, the oldest quahog individual was 192 yr old. Such extraordinary longevity requires fundamental behavioural or life history traits that support selection against aging. One important strategy would be to trade vigour, rapid growth, and reproductive output in young quahog for increased longevity. Fig. 4 depicts growth constant K of the von Bertalanffy growth function for 147 different bivalve populations (Brey 1999) and shows that _Arctica islandica_ is one of the slowest growing bivalves worldwide. Such slow growth may be the trade-off for slower aging and stable or increased reproductive output in later life, as documented for fish (e.g. Atlantic cod, Olsen et al. 2005; for a review see Pauly et al. 2002) and for long-lived turtles (Congdon et al. 2003). In both cases, older females increase reproductive output (egg production, clutch size or reproductive frequency) with age. Also, in long-lived iteroparous bivalves, reproductive output is known to increase in successive breeding seasons (Browne & Russell-Hunter 1978), the ‘trade-offs’ consisting of deferred maturation and general energy saving behaviour. Energy saving in this case means low metabolic rates involving periods of actual metabolic rate depression, as shown for _A. islandica_ from the Irish Sea (Taylor 1976). Gonadal maturation in the ocean quahog is slow and first spawning occurs between 10 and 14 yr of age (Thompson et al. 1980b). Furthermore, there is a general trend observable in aquatic ectotherms towards increased maximum life spans in the cold when comparing temperate and cold water species of similar lifestyle (Brey 1991, Brey et al. 1995, Ziuganov et al. 2000, Cailliet et al. 2001, La Mesa & Vacchi 2001), and deferred maturation and late reproductive output may form part of this life extension in cold water. This also implies a delay in physiological ageing in the cold. Lower metabolic rates at low temperatures reduce reactive oxygen species (ROS) propagation from mitochondrial respiration, thereby presumably supporting slower rates of physiological ageing (Abele & Puntarulo 2004). In the case of the ocean quahog, certainly a cold water species, it is likely that the energy saving, oxyconforming behaviour confers an adaptive value for life in the cold and, at the same time, increases longevity to support reproductive output in late age; it may also have a further slowing effect on physiological aging in addition to the effect of temperature.

Taken together, these attributes of _Arctica islandica_ suggest it is a perfect example of antagonistic pleiotropy, a theory of longevity, established by Williams (1957) and later detailed by Kirkwood (2005), and originally applied to aging-studies models such as _Drosophila_ (for a review of evolutionary genetics of longevity see Zwaan 1999).
As mentioned above, one mechanistic basis underlying such slow aging involves mitigation of ROS generation during mitochondrial oxygen turnover and the accumulation of oxidative damage by cellular maintenance processes. The mean protein carbonyl concentrations in mantle tissue, ranged lower than in shorter living and faster growing soft shell clams like *Mya arenaria* and *Laternula elliptica*, or scallops like *Aequipecten opercularis* and *Adamussium colbecki* (see Table 1). Moreover, the age-dependent change in carbonyl concentration per mg protein in *A. islandica* did not reach significance in either gill or mantle tissue because degrading hydroxylase enzymes are confined to the cytosol outside becomes deprived of autophagic potential (see Terman & Brunk 2005 for a review). In adductor muscle of *A. islandica*, lipofuscin containing lysosomes were found only in the cells outside the myofibrils. In contrast, the myofibrils themselves were essentially free of any lipofuscin, which may be a prerequisite for the preservation of muscle contractile power. This is consistent with findings from senescent mammalian cardiac myocytes, which display apparently intact myofibrils in spite of abundant lipofuscin deposits and damaged mitochondria (Terman et al. 2003). In Fig. 2f it is easy to see that myofibril diameter increased in older quahog. However, the ratio of contractile elements to accompanying interstitial cells containing the lipofuscin remained the same in young and old quahog. This means the data can be extrapolated to the whole muscle and are thus comparable to lipofuscin accumulation in gill and mantle. The slope of accumulation in Fig. 3 indicates no difference in the accumulation rate between the 3 tissues. As yet, we have been unable to sample neuronal tissues from the ocean quahog, which would have been interesting to compare with respect to age pigment accumulation from the ocean quahog.

We have shown that slow growth rates, accompanied by sustained protection of body proteins from oxidative degradation even at old age, as well as effective disposal of oxidative damage waste products in the lysosomal bodies, are important traits of longevity in the bivalve *Arctica islandica*. Importantly, the lipofuscin granules are located in metabolically less active connective and interstitial tissues, where they are less likely to impair metabolic and diffusive processes of the organs. Good preservation of tissue protein could be due to low mitochondrial ROS output at low metabolic activity of the mud clam, and transient metabolic rate depres-

<table>
<thead>
<tr>
<th>Species</th>
<th>Location</th>
<th>MLSP (yr)</th>
<th>Carbonyls (mg⁻¹ tissue protein)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Mya arenaria</em></td>
<td>North Sea</td>
<td>13</td>
<td>1.154 ± 0.2992</td>
</tr>
<tr>
<td><em>Laternula elliptica</em></td>
<td>Antarctic</td>
<td>36</td>
<td>1.875 ± 0.5747</td>
</tr>
<tr>
<td><em>Arctica islandica</em></td>
<td>Iceland</td>
<td>&gt;200</td>
<td>0.58 ± 0.46</td>
</tr>
<tr>
<td><em>Aequipecten opercularis</em></td>
<td>Irish Sea</td>
<td>8 to 10</td>
<td>1.016 ± 0.07263</td>
</tr>
<tr>
<td><em>Adamussium colbecki</em></td>
<td>Antarctic</td>
<td>45</td>
<td>1.319 ± 0.1815</td>
</tr>
</tbody>
</table>
sion may be an important behavioural strategy here (see Taylor 1976). Elevated antioxidant protection and autophagic cell clean-up may further mitigate oxidative damage and support tissue maintenance over the long lifetime of *A. islandica*. Altogether, this seems to be part of the pleiotropic life strategy of the ocean quahog, favouring longevity and enhanced reproductive output at old age over rapid growth and activity in young quahog.

Acknowledgements. The authors thank Gudmundur Vidir Helgason and Halldor Palmar Halldorsson from Sandgerdi Marine Station, University of Iceland; Gudrun G. Thorarinsdottir from the Marine Research Institute in Reykjavik, Thorsteinn Thorbergsson and Noy for their support in sampling and maintenance of the experimental quahog during the field expeditions to Iceland. Nadja Neubert gave a helping hand on the second field trip to Iceland. Angela Koehler kindly provided support from her histological laboratory. Thanks to Stefanie Meyer, Sieglinde Bahns and Kerstin Beyer for valuable technical support.

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Editorial responsibility: Otto Kinne (Editor-in-Chief), Oldendorf/Luhe, Germany

Submitted: June 14, 2007; Accepted: September 11, 2007

Proofs received from author(s): September 21, 2007