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## Telomere-independent ageing in the longest-lived non-colonial animal, *Arctica islandica*

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### ABSTRACT

The shortening of telomeres as a causative factor in ageing is a widely discussed hypothesis in ageing research. The study of telomere length and its regenerating enzyme telomerase in the longest-lived non-colonial animal on earth, *Arctica islandica*, should inform whether the maintenance of telomere length plays a role in reaching the extreme maximum lifespan (MLSP) of >500 years in this species. Since longitudinal measurements on living animals cannot be achieved, a cross-sectional analysis of a short-lived (MLSP 40 years from the Baltic Sea) and a long-lived population (MLSP 226 years Northeast of Iceland) and in different tissues of young and old animals from the Irish Sea was performed. A high heterogeneity of telomere length was observed in investigated *A. islandica* over a wide age range (10–36 years for the Baltic Sea, 11–194 years for Irish Sea, 6–226 years for Iceland). Constant telomerase activity and telomere lengths were detected at any age and in different tissues; neither correlated with age or population habitat. Stable telomere maintenance might contribute to the long lifespan of *A. islandica*. Telomere dynamics are no explanation for the distinct MLSPs of the examined populations and thus the cause of it remains to be investigated.

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### 1. Introduction

The bivalve *Arctica islandica* is the longest-lived non-colonial animal known to science. *A. islandica* can be found in the northern hemisphere and lives on continental shelves across the North Atlantic down to Virginia at the western Atlantic coast and to the Barents Sea in the east (Begum et al., 2009), where it can tolerate a wide range of different environmental factors (salinity, temperature, oxygen availability) (Basova et al., 2012). Their distribution into the Baltic Sea (BS) and the White Sea seems to be limited by ecological attributes of brackish environment like salinity, temperature, and oxygen gradients (Philipp et al., 2012). The oldest individual ever observed in this species was found Northeast of Iceland (IC) with an age of more than 500 years (Butler et al., 2013). The Irish Sea (IS) population has an estimated maximum lifespan (MLSP) of 220 years (Ridgway et al., 2012) and individuals of the BS population reach an age of approximately 40 years (Begum et al., 2010). Thus MLSP of this species varies extremely between

populations and seems to depend on the respective environmental conditions (Basova et al., 2012). The extreme difference in MLSP in *A. islandica* populations offers a promising opportunity for ageing research, such as the examination of the telomere dynamics with age.

In many species a loss of proliferative capacity of somatic cells can be observed during cellular ageing. Shortening of telomeres is discussed as one responsible factor for the replicative senescence of somatic cells, and is proposed to be one of the reasons why animals age and show an age-related increase of mortality (Wright and Shay, 2005). The telomere structure, consisting of DNA repeat sequences (5'-TTAGGG-3'), is highly conserved among vertebrates (Meyne et al., 1989) as well as among invertebrates (Traut et al., 2007). Cells approach senescence if telomere length reaches a certain threshold, the so-called Hayflick limit (Bolzán and Bianchi, 2006; Hayflick, 1965). The activation of the enzyme telomerase (Aubert and Lansdorp, 2008) or alternative telomere lengthening pathways (Bryan et al., 1997) can maintain telomere lengths, leading to theoretically unlimited proliferative potential. Hence, telomerase is expressed primarily in germ cells (Zalenskaya and Zalensky, 2002), stem cells (Mason, 2003) and in actively proliferating transit cells (Buchkovich and Greider, 1996). In normal human tissue telomerase is activated in early embryogenesis and during the first trimester, thereafter it is repressed in adult tissues, supposedly resulting from the relationship of active telomerase linked to cancer (Forsyth et al., 2002; Shay and Wright, 2011).

Abbreviations: MLSP, maximum lifespan; BS, Baltic Sea; IC, Iceland; IS, Irish Sea; LSG, line of strongest growth; PSU, practical salinity unit.

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Progressive telomere shortening is very often linked to tissue and organismal ageing (Campisi, 1996; Proctor and Kirkwood, 2002) or to a stressful environment (Metcalf and Monaghan, 2003); and telomere maintenance has been shown to play a role in organismal longevity (Haussmann et al., 2005; Joeng et al., 2004). In contrast, an increase of telomere length with age has been observed in the extreme long-lived bird, the Leach's storm petrel, which reaches a MLSP of 36 years (Haussmann et al., 2003b). Similar results with such a positive relationship of telomere length and/or telomerase activity with age are observed in long-lived trees (*Pinus longaeva*, (Flanary and Kletetschka, 2005)), the sand lizard (*Lacerta agilis*, (Olsson et al., 2010)) and the water python (*Lias fuscus*, (Ujvari and Madsen, 2009)). More intriguing examples are asexual and sexual planarian flatworms or colonial ascidians showing significantly different patterns in telomere maintenance and telomerase activities (Sköld et al., 2011; Tan et al., 2012). Other studies e.g. in humans, mice and birds found that the rate of telomere shortening depends on different life stages. Higher rates of telomere shortening were observed in earlier life stages or during development, whereas during adult life the rate stays nearly constant (Forsyth et al., 2002; Hall et al., 2004; Pauliny et al., 2006). Another interesting aspect regarding telomere dynamics and longevity is a high telomerase activity combined with low telomere attrition examined in tissues or organs of animals which grow throughout their entire life (indeterminate growth) even during adulthood. This applies to most fish, lobsters, molluscs, and other long-lived marine invertebrates (e.g. Estabrooks, 2007; Klapper et al., 1998a; Klapper et al., 1998b; Owen et al., 2007; Plohl et al., 2002; Sköld et al., 2011).

In *A. islandica*, the relationship of MLSP to growth rate or development is comparable, i.e. late onset of maturation, slow growth rate, with other well established long-lived model systems for ageing (e.g. primates, naked mole-rat) (Ridgway et al., 2011). Further, the species demonstrates indeterminate growth with very low growth rates compared to other bivalves (Begum et al., 2010; Strahl et al., 2007; Witbaard, 1996) and a constant rate of cell proliferation in many different tissues (Strahl and Abele, 2010). *A. islandica* is found in habitats with extremely different environmental conditions and populations exhibit large variations in maximum lifespans (Basova et al., 2012). Recent studies in marine bivalve molluscs have revealed an association between longevity and resistance to multiplex stressors (oxidative, genotoxic stress) as well as between longevity and proteostasis in several species including *A. islandica* (Treaster et al., 2013; Ungvari et al., 2013a; Ungvari et al., 2011; Ungvari et al., 2013b). Remarkably no enhanced antioxidant capacity or protein recycling was detected in *A. islandica* compared to shorter-lived species (Ungvari et al., 2011) but a higher ability to withstand stress factors in general was observed leading to the conclusion that retarded ageing may be the cause for stable antioxidants and not vice versa (Ridgway, 2012). A deeper understanding of telomere dynamics with respect to longevity and environmental influences in organisms may be gained by the investigation of the exceptionally long-lived bivalve, the ocean quahog. The BS population endures great fluctuations in temperature, salinity and oxygen availability (Begum et al., 2010), whereas abiotic conditions in the Northeast of Iceland are much more constant and stable (Basova et al., 2012). It is hypothesized this may greatly influence the animals' energy allocation to tissue maintenance, growth or reproduction but also the physiological response to different environmental stressors as recently reported for different *A. islandica* populations (Abele and Philipp, 2012; Basova et al., 2012; Philipp et al., 2012). The vast difference in MLSP and the environmental variables makes the two populations from the BS and IC with shortest (BS, MLSP 40 years) and longest (IC, MLSP >500 years) MLSP reported so far interesting objects for ageing studies, especially for the examination of telomere dynamics in *A. islandica*.

We therefore investigate two questions: (i) Is there a relationship between telomere length and/or telomerase activity with age in *A. islandica*? (ii) Is the large variation in maximum lifespan between populations a consequence of a significantly faster telomere shortening

that may result from higher oxidative stress levels in less favourable environments, e.g. in the BS, leading to faster telomere shortening and subsequently a reduced lifespan?

Hence, telomere dynamics among *A. islandica* individuals of differing ages and tissues of the IC, BS and the IS population are examined. In order to analyze the relationship between telomere dynamics and age, cross-sectional-analyses are necessary because longitudinal observations in *A. islandica* are, due to its long lifespan and generation cycle, not possible.

## 2. Material and methods

### 2.1. Sampling of *Arctica islandica* specimen

*A. islandica* of the BS were sampled at the station "Süderfahrt" (54°32,6 N 10°42,1E) in February 2010. In March 2010 IC animals were collected Northeast off Iceland (66°01,5 N 14°50,9 W) and in April 2010 IS animals were collected from the mouth of Belfast Lough (54°42.10 N, 5°35.25 W). All populations were fished from 20 to 25 m water depth with a hydraulic rigid-toothed dredge and kept under stable aquarium conditions with flow through water from respective environments (BS: approx. 5 °C and 20–25 PSU at the Geomar Helmholtz Centre for Ocean Research (Kiel, Germany), IC: approx. 5 °C and 35 PSU at the Suðurnes University Research Centre (Sandgerði, Iceland), IS: approx. 8 °C and 34 PSU at the School of Ocean Sciences (Bangor, Wales)). After 5–7 days of recovery from sampling stress the animals were dissected. Gill, mantle, adductor muscle for all populations and foot, the heart, digestive gland for BS and IC only were sampled, shock-frozen in liquid nitrogen and stored at –80 °C until further analysis.

### 2.2. Age determination of animals

*A. islandica* lay down annual growth rings in their shell (Schöne et al., 2005; Witbaard et al., 1994) from which the animals' individual ages can be deduced. The age determination was performed as described by Strahl et al. (2007) and Begum (2009). The right half of all shells was freed from organic remains with 13% NaOCl (AppliChem, Germany) and the line of strongest growth (LSG) embedded in two component liquid metal glue (Toolkraft, Germany). A 1 cm wedge comprising the LSG was cut out with a diamond blade using an FKS/E bench saw (Proxxon, Germany). The wedge was then cut along the LSG with an IsoMet low-speed saw (Buehler, USA). The LSG surface was sanded (with P400, P800, P1200; higher grits P2500, P4000 whenever necessary) and subsequently polished using 1 µm MicroPolish II (Buehler-met, Germany). The age was determined by age ring counts directly under a microscope (Zeiss Imager Z.1) with a 2.5 magnification lens or on the PC monitor using an AxioCam HRc camera with the AxioVision software.

### 2.3. Genomic analysis of DNA

Genomic DNA from several populations (Kattegat, White Sea, Norway and Iceland) and tissues (foot, adductor muscle) were analysed to verify the conserved telomeric repeat 5'-TTAGGG-3' in this species. Collection of animals that were included in the preparation of the molecular libraries is described in detail in Begum et al. (2010). The DNA was extracted with the Qiagen DNeasy Blood and tissue protocol (Qiagen, Germany) using foot tissue (from Norway and from Iceland) or adductor muscle (from Kattegatt and from the White Sea). DNA concentration was determined with a Nanodrop ND-100 (Thermo Scientific) and 1.25 µg of DNA from each specimen were pooled and sequenced on a Roche GS-FLX 454 sequencer (MacroGen, South Korea); for further details on preparation of the library see (Leese et al., 2012). The conserved telomere sequence 5'-TTAGGG-3' was blasted (blastn) (Altschul et al., 1990; Altschul et al., 1997) as repetitive sequence (30 repeats = 180 bp) against the genomic DNA sequence reads of *A. islandica* with a minimum eValue of 1e-003.

## 2.4. Measurement of telomere length

Mean terminal restriction fragment length (referred to as mean telomere length hereafter) was measured in gill tissue of 24 BS individuals and 38 IC animals across the respective total determined lifespan (10–36 years for BS, 6–226 years for IC animals) using the *TeloTAGGG* Telomere Length Assay (Roche, Germany) according to the protocol. DNA was extracted using DNeasy Blood&Tissue Kit (Qiagen, Germany) according to the manufacturer's instruction. The DNA yield was determined using a NanoDrop ND-1000 Spectrophotometer (Peqlab Biotechnology GmbH, Germany), 1 µg of DNA was digested by restriction enzymes (*HinfI* and *RsaI*) provided in the *TeloTAGGG* Telomere Length Assay over night at 37 °C in a heat block (Thermomixer compact, Eppendorf, Germany) and run on a 0.8% LE agarose (Biozym, Germany) gel at 30 V for 5 h in case of BS and IC animals. The gel was southern blotted onto a nylon Hybond-N+ membrane (Amersham, USA) over night and the DNA fixed by baking the membrane at 80 °C for 2 h. After hybridization and washing as described in the *TeloTAGGG* Telomere Length Assay manual, the southern blots were exposed to chemiluminescence for 20 min and analysed in a Molecular Imager ChemiDoc XRS Imaging System (BioRad, USA). Mean terminal restriction fragment lengths were detected as genomic DNA smears resolved in the gel and the area of highest density detected by the QuantityOne-4.6.8 software by BioRad was determined as the mean length of the telomeres (Fig. 1).

DNA extracted from gill, mantle and adductor muscle from 6 *A. islandica* individuals (3 young and 3 old) from the IS population was run on a 0.8% agarose gel in 0.5xTBE at 60 V for 12 h. Telomere lengths of this population were detected using a radioactive probe as described in Herbert et al. (2001).

## 2.5. Measurement of telomerase activity

Telomerase activity was measured using the Quantitative Telomerase Detection Kit (US Biomax Inc., USA) via realtime PCR. Enzyme activity was assessed in gill tissue of 10 BS and 15 IC animals previously used for telomere length determination. For 3 of these individuals per population, telomerase was additionally measured in mantle, digestive gland, and foot to investigate tissue specific enzyme activity in tissues

with different proliferation rates (Strahl and Abele, 2010). Protein extraction was performed according to the manufacturer's instructions, using 40 mg ground tissue and 100–200 µl provided lysis buffer including the protease inhibitor cocktail (product no. 11836170001, Roche, Germany). The protein concentration was determined using the DC Protein Assay (BioRad, USA) according to the protocol, and measured at 612 nm wavelength in a Tecan GENios Pro plate reader (Tecan, Germany). Protein extracts were shock frozen in liquid nitrogen and stored at –80 °C until further analysis. The Quantitative Telomerase Detection assay measures the telomerase activity from the extracted protein based on the enzyme's ability to add telomeric repeats on a substrate oligonucleotide. The elongated substrate is then amplified and quantified in a qRT-PCR. The initial concentration of telomerase can be inferred from the number of amplified qRT-PCR products using a standard curve generated with a provided standard template (TSR template). The volumes of the PCR-reaction were halved but to reduce pipetting errors, 1 µl sample was added and the water content adjusted. Prior to qRT-PCR, samples were incubated at 25 °C for 20 min to allow telomerase mediated extension of provided primers with telomeric repeat ends. 40 PCR cycles were run as recommended in the manual. All samples were run in duplicates. Prior to sample measurements, a dilution series ranging from 0.044 to 2.1 µg/µl of protein extract from a mixed sample were run to identify the optimal protein concentration of 175 ng/µl for sample analysis. For positive controls HELA cell-extracts and for negative controls heat inactivated *A. islandica* protein and heat inactivated HELA cell extracts were run to validate the method. For the calculation of telomerase activity in the samples, a TSR standard curve with the provided TSR control template was performed using 0.1 amol/µl (= 60000 molecules), 0.02 amol/µl (= 12000 molecules), 0.0008 amol/µl (= 480 molecules), 0.000032 amol/µl (= 20 molecules), and 0.0000064 amol/µl (= 4 molecules).

## 2.6. Statistics

Data evaluation and further statistical analyses such as t-tests, Mann–Whitney test and calculation of nonparametric Spearman correlation were performed using GraphPad Prism 5 Version 5.04. Determination of outliers was undertaken using Grubb's test of GraphPad

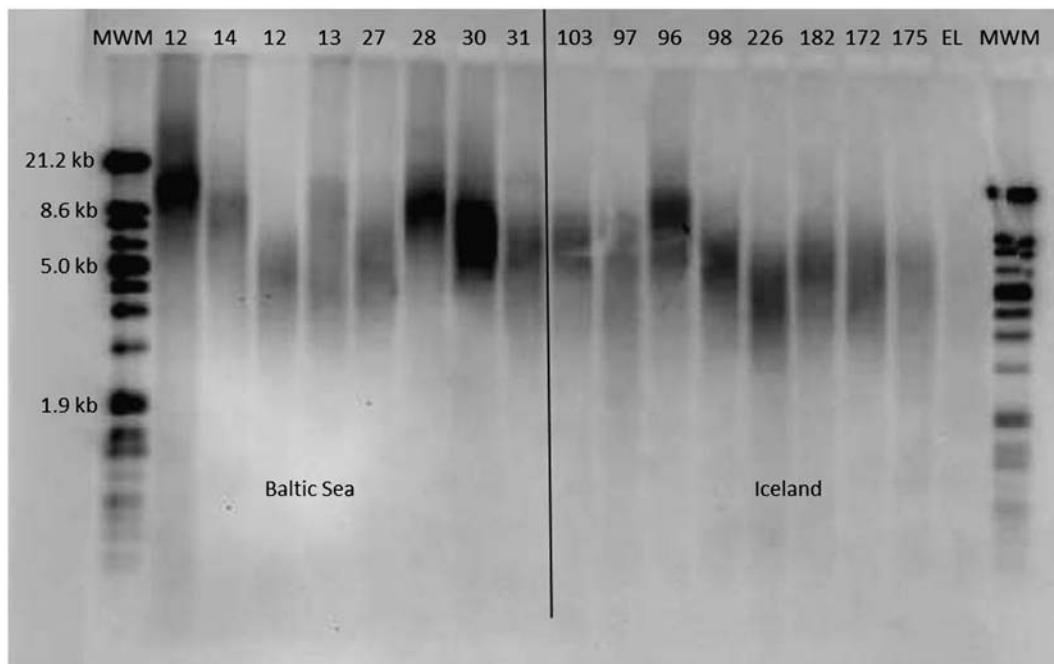


Fig. 1. Example of a Southern blot for the detection of telomere lengths. Mean terminal restriction fragment lengths are detected as smears in gill tissue of BS and IC animals. Column numbers are ages in years of investigated animals. MWM = molecular weight marker, EL = empty lane.

**Table 1**

Genomic *Arctica islandica* reads (Hit ID) that contain the telomeric repetitive sequence 5'-TTAGGG-3'. Perfection, Overlap, eValue and Score as well as matching area of query and hit are shown.

Hit ID	Perfection	Overlap [bp]	eValue	Score	qStart	qEnd	hStart	hEnd
GJ39D1101DR0VF	93%	123	1.00E-45	180	58	180	425	303
GJ39D1101CMCPB	92%	77	6.00E-23	105	103	179	315	239
GJ39D1101BDSS3	89%	64	2.00E-10	63.9	1	64	220	282
GJ39D1101ENZV3	89%	64	2.00E-10	63.9	37	100	220	282
GJ39D1101DMC7E	87%	64	5.00E-08	56.0	114	177	215	153
GJ39D1101B8FSK	88%	59	2.00E-07	54.0	119	177	288	231
GJ39D1101BMK7T	88%	53	1.00E-08	58.0	18	70	11	63
GJ39D1101C39UO	90%	51	2.00E-07	54.0	50	100	529	578
GJ39D1101BZGHQ	100%	36	9.00E-13	71.9	129	164	36	1
GJ39D1101ENDY3	100%	26	8.00E-07	52.0	17	42	56	81

Prism Software. GraphPad Prism 5 designates statistical significances as follows:  $p \leq 0.05$  (\*),  $p < 0.01$  (\*\*),  $p < 0.001$  (\*\*\*),  $p < 0.0001$  (\*\*\*\*). In column charts, data are expressed as means  $\pm$  SEM, unless otherwise indicated.

### 3. Results

#### 3.1. Genomic analysis of DNA sequences

The BLAST search of the well conserved telomeric repeat sequence, 5'-TTAGGG-3', in genomic readout data of *A. islandica* from different populations and tissues revealed ten different reads (Table 1) containing the repetitive telomere sequence. An overlap of 26–123 bp with a perfection between 88 and 100% was detected.

#### 3.2. Telomere length

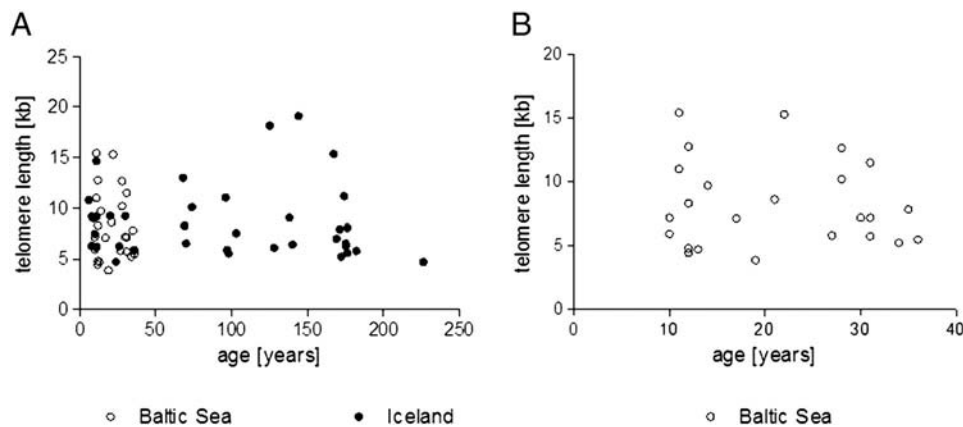
Mean telomere lengths were determined in gill tissue of 38 IC animals (age range 6–226 years), 24 BS animals (age range 10–36 years) and different tissues (gill, mantle, adductor muscle) of 6 IS animals (three young, 11 years and three old,  $2 \times 135$  and 194 years) using the TeloTAGGG Telomere Length Assay (Roche, Germany). The Southern blot in Fig. 1 reveals age-independent mean telomere length, e.g. similar telomere lengths of 12 and 27 year old BS (sample lane 3 and 5) and 98 and 182 year old IC individuals (sample lane 12 and 14). Mean telomere length in gill tissue of the IC population ranged from 4.72 to 19.11 kb, from 3.88 to 15.44 kb (Fig. 2) in the BS population and in investigated tissues of the IS population from 5.63 to 14.30 kb (Fig. 3). This indicates a high heterogeneity of mean telomere length in measured populations independent of individual age and tissues. Neither age (Figs. 2 and 3) nor size (data not shown) correlated with mean telomere length (nonparametric Spearman  $r_{SIC} = -0.22$ ,  $p$ -value =

0.18;  $r_{SBS} = -0.08$ ,  $p$ -value = 0.72) in the IC or BS populations and ANOVA of IS animals revealed no significant differences between the age groups. In young as well as old animals short and long telomeres were detected (Figs. 2 and 3). Further, no significant differences in mean telomere length between individuals of IC, BS and IS of similar age or between different tissues within the IS population could be observed (mean of IC = 8.55 kb, mean of BS = 8.26 kb, mean of IS = 7.73 kb).

#### 3.3. Telomerase activity

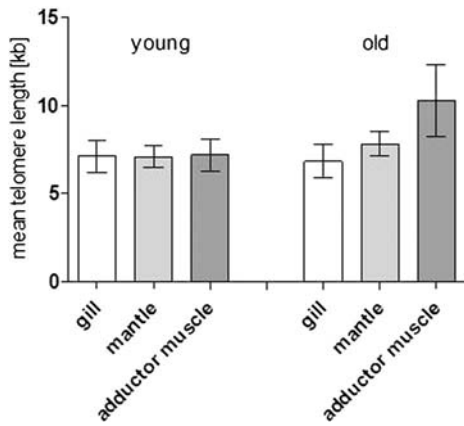
Telomerase activity was measured in gill tissue of the BS and IC populations across a wide age range (BS = 10–36 years, IC = 8–226 years). Telomerase activity was detected in individuals of all ages in both populations with high variability (Fig. 4), showing that the enzyme stays active throughout the complete lifespan of the animals. Statistical analysis showed no correlation of age with telomerase activity (nonparametric Spearman  $r_{SIC} = 0.14$ ,  $p$ -value = 0.64;  $r_{SBS} = 0.22$ ,  $p$ -value = 0.54). The enzyme activity of both populations at the same age was comparable. Since the same individuals were used for telomerase activity and telomere length analysis, results could be correlated but revealed no direct coherence between mean telomere length and telomerase activity (Fig. 5) (nonparametric Spearman  $r_{SIC} = 0.15$ ,  $p$ -value = 0.60;  $r_{SBS} = -0.35$ ,  $p$ -value = 0.33).

Additionally, telomerase activity was tested in gill, mantle, digestive gland, and foot tissue which are known to have different proliferation rates (Strahl and Abele, 2010). Enzyme activity was measured in BS individuals aged 21, 22 and 36 years and IC animals aged 68, 69 and 98 years. In 83% of investigated animals, telomerase was most active in foot and in 80% the enzyme was least active in mantle tissue (Fig. 6). In both populations the enzyme shows the same activity trend in the different tissues.



**Fig. 2.** Mean telomere length in kilobases (kb) in gill tissue of (A) 38 IC (closed circles) and 24 BS animals (open circles) measured across total determined lifespan of both *A. islandica* populations. (B) BS samples are separately shown for better overview.



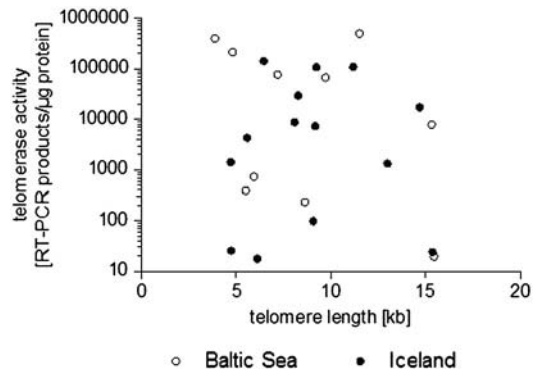


**Fig. 3.** Mean telomere length  $\pm$  SEM in kilobases (kb) in gill (open bars), mantle (light grey bars) and adductor muscle (dark grey bars) of three young (all 11 years) and three old ( $2 \times 135$  and 194 years) *A. islandica* from the IS.

#### 4. Discussion

The well conserved telomeric repeat sequence 5'-TTAGGG-3' that can also be found on human chromosome ends, has previously been detected in marine bivalves such as the mussel (*Mytilus galloprovincialis*, (Pohl et al., 2002)), the bay scallop (*Argopecten irradians*, (Estabrooks, 2007)), the Pacific oyster (*Crassostrea gigas*, (Guo and Allen, 1997)) and other marine invertebrate species e.g. the sea urchin (*Strongylocentrotus purpuratus*, (Lejnine et al., 1995)) (Traut et al., 2007). However, as a first study we report this sequence in genomic DNA of several populations of the longest-lived non-colonial animal *A. islandica*. It is therefore eligible to measure telomere length using the TeloTAGGG Telomere Length Assay by Roche relying on this repeat sequence.

Individuals of the long-lived bivalve *A. islandica*, sampled at a wide range of ages, demonstrate the potential to maintain telomere length throughout their life. Mean telomere length was measured in gill tissue across all ages in the short-lived (investigated ages: 10–36 years) BS population, in the long-lived (investigated ages: 6–226 years) IC population as well as in three different tissues of young (11 years) and old ( $2 \times 135$  and 194 years) *A. islandica* from the IS. Thus, 90% and 88% of the BS MLSP (40 years) and the IS MLSP (220 years) respectively were covered. The individual with the highest MLSP ever observed was recovered Northeast of IC from deeper grounds (>80 m depth, at 66°31,59'N 18°11,74 W (Butler et al., 2013)) whereas the population investigated in the presented study was recovered from ca. 20 m depth from another bay Northeast of IC (66°01,5 N 14°50,9 W). Populations may be less disturbed at deeper grounds so that older individuals

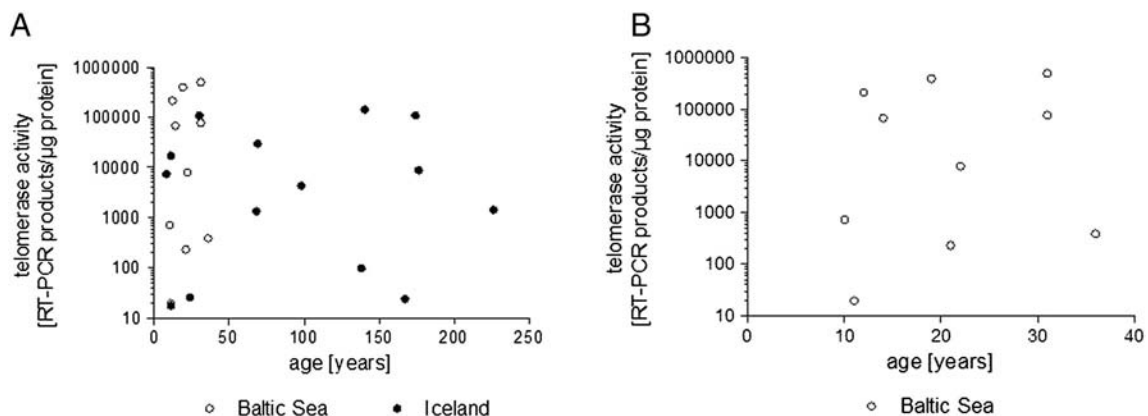


**Fig. 5.** Correlation of telomerase activity (in RT-PCR products/ $\mu$ g protein) with mean telomere length measured in gill tissue of the same individuals of IC (closed circles) and BS (open circles) population.

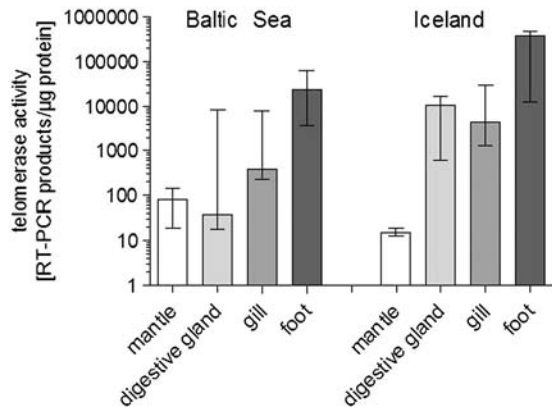
are more likely to be recovered from lower depths. We can assume that the studied population may reach a similar MLSP as reported for the population in Butler et al. (2013). However, the oldest individual in this study reached an age of 226 years, so that we consider this age as MLSP of this population. An overview of populations and studied ages are given in Table 2 demonstrating that telomere dynamics were studied in a high proportion of population-specific MLSPs.

The high heterogeneity of mean telomere length observed in *A. islandica* has also been reported in humans (Hewakapuge et al., 2008). The detected range of mean telomere length in *A. islandica* (3.88–19.11 kb) is comparable to those found in humans (3–17 kb) (Harley et al., 1990) and other species such as common terns (*Sterna hirundo*, (Hausmann et al., 2003a)) and the wedgeshell clam (*Donax trunculus*, (Pohl et al., 2002)). The maintenance of telomere length over age in *A. islandica* might be achieved by the activity of the enzyme telomerase found at all ages which did not change significantly over age. This could represent a prerequisite to prevent senescence and postpone mortality. Remarkably, the high variability in mean telomere length and telomerase activity at all ages between and within the BS and IC population could be observed leading to the conclusion that the high variation in telomere dynamics in the different age classes of *A. islandica* may be due either to external influences or individual properties.

Telomere dynamics have been widely studied as a proxy of cellular and organism senescence in relation to survival and lifespan (Hausmann and Marchetto, 2010; Horn et al., 2010). Telomere dynamics with age can show a large number of varieties: The classical pattern is that telomere length declines with age as observed in some animal species and also human (Aubert and Lansdorp, 2008; Harley et al., 1990); Zebra finch, (Hausmann and Vleck, 2002); and Great



**Fig. 4.** Telomerase activity (in RT-PCR products/ $\mu$ g protein) in (A) gill tissue of 15 IC (closed circles) and 10 BS (open circles) animals. No correlation of telomerase activity with age could be observed. (B) BS samples are separately shown for better overview.



**Fig. 6.** Mean telomerase activity  $\pm$  SEM in RT-PCR products/ $\mu$ g protein in mantle (open bars), digestive gland (light grey bars), gill (dark grey bars) and foot tissue (black bars) of the BS and IC population ( $n = 3$  individuals per tissue). In mantle tissue of the BS population telomerase activity of only two individuals was measured.

frigatebird, (Juola et al., 2006). However, shortening of telomeres does not occur at a constant rate with age (Horn et al., 2010). A second pattern, contradictory to the first, shows increasing telomere length with advancing age as found in the bristlecone pine, (*P. longaeva*, (Flanary and Kletetschka, 2005)), sand lizards (*L. agilis*, (Olsson et al., 2010)), and the water python (*L. fuscus*, (Ujvari and Madsen, 2009)). A third pattern describes no age-related telomere shortening in adulthood but a strong and often rapid shortening earlier in life, as observed in birds, the European shag and the wandering albatross (Hall et al., 2004; Pauliny et al., 2006). For *A. islandica* we document no change with age along the range of ages investigated (BS: 10–36 years, IS: 11–194 years, IC: 6–226 years) supporting several studies (Abele et al., 2008; Strahl et al., 2007; Ungvari et al., 2011) that found stable physical parameters in the ocean quahog over age which implies a lack of ageing in this species. Indeed the dynamics of both telomere length and telomerase activity in *A. islandica* seem to be stable and similar across all ages. Due to the chosen sampling pattern we have no information about telomere length during earlier growth stages of individuals younger than 6 years from the IC, 10 years from the BS and 11 years from the IS population. Telomerase expression of the sand scallop, *Euvola ziczac*, however has been detected in adult tissue (adductor muscle, gill, mantle, and male and female gonads) as well as during early developmental stages (Owen et al., 2007). The results of *A. islandica* are in line with other animals showing indeterminate growth, such as the rainbow trout, lobsters, molluscs, or other marine invertebrates, which express telomerase in all examined tissues during their complete investigated lifespan (Klapper et al., 1998b; Owen et al., 2007). As an emerging model system for ageing studies, bivalves, and especially the ocean quahog, can also compete with established long-lived model systems for ageing such as primates and the naked mole-rat (Austad, 2010; Fischer and Steven, 2011; Lewis et al., 2012) that show similar life history associations between MLSP and growth rate or development (i.e. late onset of maturation, slow growth rate) (Ridgway et al., 2011). These findings imply that *A. islandica* does not merely attain exceptional longevity through low metabolic rate but also possesses similar traits to long-lived animals of other taxa.

Surprisingly, *A. islandica* from IC showed no specific strategy according to the telomere dynamics as compared to the BS individuals which is remarkable in light of the 10-fold difference in maximum lifespan between the two populations (BS: MLSP 40 years, IC: MLSP >500 years). As mentioned above, however, no sign of ageing has been found in this species and the short lifespan of *A. islandica* in the BS may rather be attributed to poor shell formation in a low salinity environment (Begum et al., 2010; Hiebenthal et al., 2012). Moreover, compared to the highly stable marine IC habitat the brackish BS environment is highly fluctuating not only in respect to salinity but also in ionic compositions of the water body, water temperature, oxygen content and nutrient concentrations which implies a general more stressful habitat and an impact on telomere dynamics as hypothesized by others (Hall et al., 2004; Horn et al., 2008; Jennings et al., 2000). Philipp et al. (2012) for example could show that *A. islandica* from the Baltic Sea and Iceland were found to differ in transcriptional response towards hypoxia which was attributed to the adaptation to different environmental stability of the two populations. Stressful environments have been linked to higher levels of oxidative stress, associated to telomere shortening (Jennings et al., 2000; Von Zglinicki, 2002). However, no correlation between telomere length and telomerase activity with age in either population of *A. islandica* indicates that the maintenance of telomere length is independent of environmental conditions in this species. Several telomere-independent functions of telomerase e.g. protection of mitochondria, inhibition of apoptosis, enhancement of DNA repair, promotion of cell growth and stem cell proliferation (reviewed in Bollmann (2008)) might influence enzyme activity and are important for the individual's fitness in their respective environment. A sustained expression of telomerase observed in *A. islandica* could hence also contribute to telomere-independent functions of telomerase.

We detected telomerase throughout all ages but also in different tissues (gill, mantle, digestive gland, foot), with lowest activity in mantle and highest in foot in the BS and IC population (Fig. 6). Telomerase activity in diverse tissues of several marine organisms with indeterminate growth has already been reported previously (Klapper et al., 1998a; Klapper et al., 1998b; Owen et al., 2007). Belair et al. (1997) demonstrated telomerase activity serves as a marker for cell proliferation in human tumours and normal cells, thus high telomerase activity is expected in highly proliferative tissue. According to Strahl and Abele (2010) mantle and gill tissue of IC *A. islandica* show similar proliferation rates whereas in animals from the German Bight proliferation rates were higher in gills compared to mantle tissue. Proliferation conditions in the different tissues in *A. islandica* are therefore not resolved and it is unclear whether the higher telomerase activity in gill compared to mantle tissue in the present study reflects higher proliferation rates in gill tissue. Foot tissue was not measured by Strahl and Abele (2010), thus there is no information about proliferation activity which could be correlated with the higher telomerase activity found in this tissue compared to all other investigated tissues. Personal observations (E.E.R. Philipp), however, noted a reduction in foot size after 16 days of exposure of *A. islandica* to hypoxia. We hypothesize the foot may serve as glycogen storage tissue accessed during periods of anaerobiosis (Oeschger, 1990), and is diminished after debilitating conditions. As this species is regularly exposed to hypoxia/anoxia, self-induced or by the environment (Philipp et al., 2012), high proliferation rates in the foot might therefore be a prerequisite for fast utilization and recuperation of the glycogen storage.

**Table 2**

Sample attributes of investigated populations and studied ages in relation to population-specific MLSPs. N = sample size, TL = telomere length, TA = telomerase activity, NA = data not available.

Population	N TL/TA	Size-range [mm]	MLSP [years]	Investigated ages [years]	Investigated mean age for TL/TA [years]	Age-range of MLSP covered
Baltic Sea	24/10	33.16–60.6	40	10–36	17.7/20.7	65%
Irish Sea	6/NA	56.3–143.0	220	11,135,194	81.0/NA	83%
Iceland	38/15	22.11–105.02	226	6–226	83.6/101.5	97%

A possible selective mortality of individuals with shorter telomeres (Hausmann and Mauck, 2008) restricts cross-sectional-analyses in their conclusion about telomere shortening and telomerase activity over age and life stages. Another factor that cannot be examined in cross-sectional studies is environmental stressors which may have significant impacts on the rate of the telomere dynamics on both individual and population levels (Hall et al., 2004; Horn et al., 2008, 2010; Von Zglinicki, 2002).

#### 4.1. Summary and conclusion

As emerging model systems, bivalves exhibit similar life history and developmental traits (Ridgway et al., 2011) compared to other model systems such as primates and the naked mole-rat (Fischer and Steven, 2011; Lewis et al., 2012). Especially the longest-lived non-colonial animal known to science, the bivalve *A. islandica*, shows a general lack of signs of ageing (Abele et al., 2008; Strahl et al., 2007; Ungvari et al., 2011). Supporting these findings, our study is the first presenting a lack of ageing-associated telomere dynamics in the ocean quahog. We document the well conserved telomeric repeat sequence 5'-TTAG GG-3' in several populations of *A. islandica*. We further demonstrate highly variable mean telomere length and telomerase activity which is continually expressed in different tissues throughout the lifespan in the longest- and shortest-lived populations of *A. islandica*. The correlation of telomerase activity with age, size and tissue was investigated as well as the correlation of telomere length with telomerase activity. Apparently, *A. islandica* is extremely successful at maintaining cellular proliferation showing a lack of age-associated telomere shortening accompanied by a continuous telomerase activity throughout its lifespan. Equal telomere dynamics investigated in *A. islandica* populations with extremely different MLSPs indicate that MLSP is not determined by telomere length or telomerase activity so that the causative factor of this exceptional longevity remains to be investigated.

#### Conflict of interest

The authors have no conflicts of interests.

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