Epigenetic estimation of age in humpback whales

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Abstract

Age is a fundamental aspect of animal ecology, but is difficult to determine in many species. Humpback whales exemplify this as they have a lifespan comparable to humans, mature sexually as early as 4 years and have no reliable visual age indicators after their first year. Current methods for estimating humpback age cannot be applied to all individuals and populations. Assays for human age have recently been developed based on age-induced changes in DNA methylation of specific genes. We used information on age-associated DNA methylation in human and mouse genes to identify homologous gene regions in humpbacks. Humpback skin samples were obtained from individuals with a known year of birth and employed to calibrate relationships between cytosine methylation and age. Seven of 37 cytosines assayed for methylation level in humpback skin had significant age-related profiles. The three most age-informative cytosine markers were selected for a humpback epigenetic age assay. The assay has an $R^2$ of 0.787 ($P = 3.04e-16$) and predicts age from skin samples with a standard deviation of 2.991 years. The epigenetic method correctly determined which of parent–offspring pairs is the parent in more than 93% of cases. To demonstrate the potential of this technique, we constructed the first modern age profile of humpback whales off eastern Australia and compared the results to population structure 5 decades earlier. This is the first epigenetic age estimation method for a wild animal species and the approach we took for developing it can be applied to many other non-model organisms.

Keywords: age, cetacean, DNA methylation, epigenetics, population

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Introduction

Animal age is a determinant of many individual and population characteristics. Estimates of age can be used to understand traits in wild animals such as reproductive potential (Clapham 1992), developmental processes (Cook et al. 2006) and factors affecting survival and reproductive success (Chaloupka et al. 1999; Campana 2001). A common approach for age estimation in wild animals that have not previously been encountered is to count annually accrued features such as the growth rings found in fish otoliths that are widely used for producing population age estimates for informing fishery management (Campana 2001). Animals that cannot be lethally sampled provide a greater challenge as many have no outward features that change with age. In these species, individuals can be marked artificially, or records can be made of natural individual features in early life and the elapsed time determined upon re-identification. Whale age estimation was important when whales were the subject of commercial fisheries for monitoring population status (Chittleborough 1959, 1965). It continues to be important for monitoring the recovery of whale populations from the effects of past harvesting (Chaloupka et al. 1999; Baker & Clapham 2004).

Humpback whales (Megaptera novaeangliae, ‘humpbacks’) are one of the best studied cetacean species. However, they display no accurate visual markers for age after they are weaned at about 1 year (Chittleborough 1959) and lack of age information continues to limit our understanding of them. Sexual maturity is reached early in life and age at first parturition can be as young as 5 years (Chittleborough 1965; Clapham 1992; Barlow & Clapham 1997). The age of dead humpback whales can be estimated from ear plug growth layer groups (GLGs), a waxy structure within the ear (Chittleborough 1959, 1965; Lockyer 1984; Gabriele et al. 2010), and baleen plate thickness or ovary condition (Chittleborough 1959). Humpbacks have a maximum verified lifespan of 95 years (Chittleborough 1965), if single ear plug GLGs are considered to accrue annually (Gabriele et al. 2010). None of these age estimation methods can be applied to live whales.
Age estimation methods for live humpbacks are based either on repeat photographic identification or minimally invasive sampling of whale tissues. Humpbacks have natural individual markings on the underside of the tail flukes that identify them (Chaloupka et al. 1999). Repeat sightings of whales whose natural markings were photographed as calves allow age to be estimated (Gabriele et al. 2010). Humpback DNA is used for a wide range of genetic analyses including population size by mark-recapture (Palsbøll et al. 1997; Rew et al. 2011), population structure (Baker et al. 1993; Schmitt et al. 2013), population assignment of individuals (Pomilla & Rosenbaum 2005), effective population size (Roman & Palumbi 2003), kinship (Balsecchi et al. 2002) and sex determination (Morin et al. 2005). Attempts to use humpback whale DNA for age estimation have so far focused on telomere length assays (Dennis 2006; Olsen et al. 2012), but these suffer from numerous sources of measurement error (Olsen et al. 2012); factors other than age that cause telomere length changes (Dunsehra et al. 2011); and a wide range of inherited telomere lengths at birth (Kappei & Londonio-Vallejo 2008). In cetaceans, there is an additional problem in that ‘telomeric’ repeat sequences occur in nontelomeric regions (Dunsehra et al. 2011). The only molecular method currently available for age estimation of live humpback whales is lipid profile analysis (Herman et al. 2009).

Biological ageing is a combination of programmed processes (Berdasco & Esteller 2012; Horvath 2013) and accumulated changes caused by unrepaired environmental damage (Kujoth et al. 2005). Recent evidence suggests that epigenetic changes are both directing the process of ageing and being caused by it (Maegawa et al. 2010; Koch et al. 2011; Winnefeld & Lyko 2012; Hannum et al. 2013; Horvath 2013). The best studied class of epigenetic change in vertebrates is methyl group presence or absence at the C5 position of Cytosine residues that are adjacent to Guanidine residues (‘CpG sites’). Clusters of CpG sites are common in the 5’ regulatory regions of vertebrate genes (Hannum et al. 2013). CpG methylation levels play an important role in control of gene expression, where higher methylation levels (‘hypermethylation’) generally reduce gene transcription rate. Methylation changes at specific CpGs have been linked to age in mice (Maegawa et al. 2010) and humans (Christensen et al. 2009; Grønniger et al. 2010; Bocklandt et al. 2011; Koch & Wagner 2011; Hannum et al. 2013). Epidermal methylation changes in some genes that relate to age have been shown to affect a set of CpG sites distinct from the sites that change in relation to environmental impacts such as sun exposure (Grønniger et al. 2010). This suggests that some CpG sites are linked to genetically programmed ageing and are less influenced by environmental variables. Identification of epigenetic changes in tissues that can be sampled relatively noninvasively such as blood (Koch et al. 2011), skin (Grønniger et al. 2010) and buccal cells (Bocklandt et al. 2011) provides opportunities for development of epigenetic age assays for live animals. Several recent studies have used epigenetic assays to estimate the age of humans or human tissues (Bocklandt et al. 2011; Koch & Wagner 2011; Garagnani et al. 2012; Hannum et al. 2013; Horvath 2013).

We developed an epigenetic method for estimating humpback whale age with DNA purified from skin biopsy samples, hereafter referred to as the HEAA (Humpback Epigenetic Age Assay). We identified three CpG sites in the 5’ regulatory regions of three humpback whale genes with cytosine methylation levels that have a strong age relationship. We measured methylation levels in 45 whales of known ages ranging from <1 to 30 years to characterize the statistical properties of the HEAA including its precision and accuracy. We also specifically explored the ability of the HEAA to determine which of a pair of humpback whales is older and which is younger (ordinal age estimation) because this has a specific application in close-kin population size estimation. The HEAA was then used to estimate the age profile for a sample of 63 humpback whales of unknown age of Australia. This age profile was compared to profiles from the same population 47–57 years earlier as an example of one application of this technique. Finally, we applied the methylation assays used in the HEAA to sperm whale DNA to explore cross-species application of age assays based on DNA methylation.

Materials and methods

Sample collection and processing

Skin samples were collected from humpback whales by punch biopsy darts fired from crossbows or modified rifles. Samples were preserved in 75% ethanol or frozen and later transferred to ‘RNA later ICE’ (Qiagen). DNA was purified by CTAB extraction (Stewart & Via 1993). Three populations of humpback whales were sampled at three sites: the Gulf of Maine off eastern North America (43° N, 68° W) from May to August in 2007–2011; Evans Head in eastern Australia (29° S, 153° E) from June to July 2009; and Exmouth, Western Australia (22° S, 114° E) in September 2009. The sex of the whale associated with each sample was determined with a qPCR assay (Morin et al. 2005).

Three sample sets were assembled. The ‘calibration’ samples were chosen to represent an even distribution of ages within the available range from photo-identification studies (a few weeks to 30 years) and to evenly represent both sexes. They consisted of 45 DNA samples purified from skin biopsies, 40 from the Gulf of Maine, one...
known-age adult from Evans Head and three calves from Exmouth. There were 21 samples from females and 24 from males. The 24 ‘mother–calf pair’ samples consisted of six pairs from Exmouth, two from Evans Head and four from the Gulf of Maine. The ‘test’ samples were purified from 63 skin biopsies collected off Evans Head, 50 of which were males and 13 female.

Age determination for ‘calibration’ population whales

The age of 40 humpbacks from the Gulf of Maine was determined by resighting of individuals first seen as dependent calves. Biopsies taken in the Gulf of Maine were linked in the field to fluke photographs. Identity and year of birth was confirmed against a photographic identification catalogue curated by the Centre for Coastal Studies. The age of one humpback sampled near Evans Head, Australia was also known as it has been repeatedly resighted since it was a calf (Polanowski et al. 2011). Samples of four calves from near Exmouth, west Australia were estimated to be 4–6 weeks old based on size.

Identification of age-related epigenetic markers in humpbacks

Genes with age-related epigenetic changes in humans and mice were identified through literature searches. The genes tested and the studies demonstrating their age-related methylation are given in Table 1. Candidate 5’ regulatory region sequences were taken from GenBank and used as queries for BLASTN (Altschul et al. 1990) searches of cetacean sequences in GenBank and BLAT searches of the dolphin (*Tursiops truncatus*) genome (Vollmer & Rosel 2012). Where candidate genes had a clearly orthologous 5’ regulatory regions in the *T. truncatus* genome, primers for amplification of humpback sequences were designed by eye based on all available homologous cetacean sequences. Humpback gene regions were amplified in 10 μL PCR reactions containing 5 μL 2× Phusion HF (NEB) master mix, 1 μM of each amplification primer, 10 ng of humpback whale DNA and milli-Q H₂O with thermal cycling conditions appropriate to each primer set and predicted amplicon. The fragments were purified with Ampure magnetic beads (Agencourt) and bidirectionally sequenced by dye terminator v 3.1 chemistry on an ABI 3100 Sanger sequencer (Applied Biosystems).

Measurement of cytosine methylation levels

Cytosine methylation levels were measured with Qiagen PyroMark assays. The pyrosequencing assays were designed using Pyromark Assay Design Software (Version 2.0.1, Qiagen). Humpback DNA was converted using the Epitect Bisulphite Conversion Kit (Qiagen). The assay regions were PCR amplified using a biotin-labelled, HPLC-purified primer and standard sequencing grade primer (Table S1, Supporting information). Amplification reactions consisted of 12.5 μL PYROMARK mastermix, 2.5 μL Coral Load, 1 μL each of 5 μM forward and 5 μM reverse primers, 2 μL of bisulphite converted template DNA and 6 μL of water. Thermocycling conditions were 15 min at 95 °C followed by 45 cycles of 30 s at 95 °C, 30 s at 56 °C and 30 s at 72 °C and a final extension step of 10 min at 72 °C. Pyrosequencing was performed on a PyroMark 24 Pyrosequencing System (Qiagen). The PyroMark Q24 software gave percentage methylation values for each CpG site.

Selection of sites for the HEAA

Cytosine methylation percentages for 37 CpG sites were compared to the ages of the 45 whales in the ‘calibration’ sample set. Linear regression was used to show how much of the variation in CpG methylation was explained by age differences. To correct P values for multiple age-methylation comparisons, a Bonferroni–Holm correction procedure was applied (Holland & Copenhaver 1987). The CpG sites that had significant relationships with age were considered for incorporation into the HEAA. Twenty different combinations of either two or three CpG sites that were found in separate gene regions were combined into multiple linear regression models. The combination of sites that produced a multiple regression with maximum predictive power was selected by scores for the Akaike Information Criterion (AIC) (Table S2, Supporting information).

Measurement of HEAA accuracy and precision

The accuracy of the HEAA was assessed with multiple linear regression. The overall precision of the HEAA was assessed with a Leave One Out Cross Validation (LOOCV) (Picard & Cook 1984). For each of the 45 samples in the ‘calibration’ set, age was estimated with the model using the other 44 samples to calculate the multiple linear regression. The difference between the known and estimated age value was recorded for each sample. The distribution of these residuals was assessed for normality and an assessment of leverage of individual points made.

An estimate of the proportion of assay error that could be attributed to the PyroMark assay was made by repeated measurement of CpG methylation levels in the same DNA samples. Four samples with known ages of 0.4, 5.5, 7.5 and 22.3 years were assayed six times each. The mean standard deviation for all 24 measurements...
CpG sites screened for age-related methylation in *Megaptera novaengliae*

| Gene       | Evidence                     | CpG position | Age relationship             |
|------------|------------------------------|--------------|------------------------------|
| TET2       | Human hypermethylation       | −12          | Hypomethylation, $R^2 = 0.211$, $P = 0.000877$ |
| KF791963   | (Grönniger *et al.* 2010)    | +16          | Hypomethylation, $R^2 = 0.174$, $P = 0.00256$ |
|            |                              | +21          | Hypomethylation, $R^2 = 0.189$, $P = 0.00159$ |
|            |                              | +31          | Hypomethylation, $R^2 = 0.409$, $P = 1.75 \times 10^{-6}$ |
|            |                              | +58          | None, $P > 0.05$              |
| CDKN2A     | Human hypermethylation       | +297         | Hypermethylation, $R^2 = 0.409$, $P = 1.37 \times 10^{-6}$ |
| KF791964   | (Gonzalez-Zulueta *et al.* 1995; Krishnamurthy *et al.* 2006; Koch & Wagner 2011; Koch *et al.* 2011; Horvath 2013) | +303         | None, $P > 0.05$              |
|            |                              | +309         | Hypermethylation, $R^2 = 0.344$, $P = 1.39 \times 10^{-5}$ |
|            |                              | +327         | None, $P > 0.05$              |
| TRIM58     | Human hypermethylation       | +181         | None, $P > 0.05$              |
| KF791966   | (Koch *et al.* 2011)         | +190         | None, $P > 0.05$              |
|            |                              | +205         | None, $P > 0.05$              |
|            |                              | +222         | None, $P > 0.05$              |
|            |                              | +230         | None, $P > 0.05$              |
|            |                              | +257         | None, $P > 0.05$              |
|            |                              | +291         | None, $P > 0.05$              |
|            |                              | +295         | None, $P > 0.05$              |
|            |                              | +309         | None, $P > 0.05$              |
|            |                              | +312         | None, $P > 0.05$              |
|            |                              | +314         | None, $P > 0.05$              |
|            |                              | +323         | None, $P > 0.05$              |
|            |                              | +329         | None, $P > 0.05$              |
|            |                              | +332         | None, $P > 0.05$              |
|            |                              | +340         | None, $P > 0.05$              |
| HoxA9      | Human hypermethylation       | +252         | None, $P > 0.05$              |
| KF791967   | (Grönniger *et al.* 2010; Koch *et al.* 2011) | +255         | None, $P > 0.05$              |
|            |                              | +265         | None, $P > 0.05$              |
| DDAH2      | Human hypermethylation       | +31          | None, $P > 0.05$              |
| (Grönniger *et al.* 2010) | +52          | None, $P > 0.05$              |
|            |                              | +65          | None, $P > 0.05$              |
|            |                              | +85          | None, $P > 0.05$              |
| TOM1L1     | Human hypermethylation       | +539         | None, $P > 0.05$              |
| (Bocklandt *et al.* 2011) | +533         | None, $P > 0.05$              |
|            |                              | +517         | None, $P > 0.05$              |
| Edaradd    | Human hypermethylation       | −20          | None, $P > 0.05$              |
| KF791968   | (Bocklandt *et al.* 2011)    | −31          | None, $P > 0.05$              |

Regressions of CpG methylation with age for 37 CpG sites in eight *Megaptera novaengliae* genes. The name of the homologous gene in humans is given and the accession no. of the GenBank entry for the *M. novaengliae* sequence produced in this study. GenBank entries for the *M. novaengliae* DDAH2 and TOM1L1 sequences were not possible as these sequences were too short to be accepted by GenBank. The source and nature of the evidence for age-related CpG methylation in humans or mice is shown. The position of the 5' Cytosine of each CpG in each humpback gene is indicated relative to the gene’s start codon with negative values indicating distance in base pairs to the 5' of the start codon and positive values 3' of the start codon. The CpG and age regression $R^2$ values are shown. All regression $P$ values <0.05 were significant after Bonferroni–Holm correction.

was calculated from differences between known and estimated age.

**Estimate of whale ages in a test population**

The age distribution estimated by the HEAA for the 63 Evan’s Head whales (21% female, 79% male) was compared to age distributions determined by ear plug GLG counts and ovarian measurement for east coast Australian humpbacks in 1952–1962 (Chittleborough 1965). These age estimates were doubled to conform to more recent evidence that one ear plug GLG accrues annually (Gabriele *et al.* 2010). The numbers of animals with age estimates for 1952–1962 were, respectively 598,
Estimates of ordinal age differentiation performance

The performance of the HEAA in correctly determining the age order of whales was estimated for a range of age differences. The exact ages for the older and younger age were converted into estimates that the HEAA might produce by selecting a value at random from a normal distribution centred on the exact age with a standard deviation of 2.991 as estimated with LOOCV. This was done 10,000 times for all age differences from 0 to 100 (Fig. S1A, Supporting information).

To estimate how often age order in parents and offspring in a real population would be correct, population age distributions were simulated as negative exponential distributions with the \( \lambda \) parameter estimated from the mean age of the 2009 HEAA results for a growing population (\( \lambda = 1/10.01 = 0.0999 \)) and from mean age recorded in 1952 (Chittleborough 1965) for the same population (\( \lambda = 1/21.8 = 0.0459 \)) (Fig. S1B, Supporting information). Ten thousand parental ages were selected at random from the portion of these distributions greater than the minimum parturition age for humpbacks of 5 years. An offspring age was also selected at random with a maximum age limit of the parental age minus 5 years. This gave the distribution of different age intervals present in the population (Fig. S1C, Supporting information). These were multiplied by the proportion that would be correctly aged and the results integrated across all ages to give an overall proportion that would be correctly age ordered.

The HEAA’s ordinal age estimation performance was tested empirically on 12 samples from mothers and calves. The pairs were initially identified visually as mothers with dependent young and their relationship confirmed by microsatellite genotyping following previously published methods (Schmitt et al. 2013). The four mother–calf pairs collected from the Gulf of Maine (real ages 17.4 ± 0.4, 10.3 ± 2.5, 25.3 ± 3.6, 25.8 ± 2.2) had their age estimated with the HEAA calibrated with the other 44 (total 11) samples in the calibration sample set as for the LOOCV analysis, where all other pairs used the full HEAA calibration.

Test of HEAA assays on sperm whale DNA

Samples of sperm whale (Physeter catodon) skin and teeth were collected from whales that died in mass strandings at Perkins Island off northern Tasmania, Australia (42°2 S, 145°14 E) in December 2004 and January 2009. Ages were estimated at 0 years for a foetus, 1 year for a calf and 15, 20, 28 and 38 years for four individuals by standard tooth growth ring counting methods (Evans & Hindell 2004). The HEAA was applied to these samples as for humpbacks.

Results

Development of the HEAA

Humpback whale’s regulatory regions were isolated for eight genes that had evidence of CpG methylation changes in other mammals. Thirty-seven CpGs in these genes were assayed for correspondence between CpG methylation levels and age (Table 1). CpG sites considered for the HEAA were selected from the seven CpG sites that had a significant regression relationship with age. Multiple linear regression models were made for all 20 combinations of CpG sites that were not from the same gene region. The models were ranked by Akaike Information Criterion (AIC) score to identify which combination of two or three sites had the best ability to predict age, as shown in Table S3 (Supporting information). The model with the best AIC score had the sites TET2_CpG+31, CDKN2A_CpG+297 and GRIA2_CpG+202. These three sites were also those with the strongest regression relationship with age (Table 1). The regressions of age and CpG methylation levels in the three sites selected for the assay are shown in Fig. 1. There was no significant difference in the regressions for female and male CpG levels and age (analysis of covariance \( P > 0.05 \) in all cases). This was also true for the other four sites that had significant relationships between age and CpG methylation.

The main reason for selecting only one CpG site from each gene was that sites within the same region that have been methylated or demethylated as part of the same process would not provide independent biological age estimates. Concerted methylation changes are often found among CpG sites in the same human gene regions (Grönniger et al. 2010; Koch & Wagner 2011) and we found this in all sites within the same regions of humpback TET2 and CDKN2A (Fig. S2 and Table S2, Supporting information). Sites from the same region would also be subject to similar experimental error when assayed in the same PCR amplifications and PyroMark assays. This strategy follows those taken for developing human epigenetic age assays (Bocklandt et al. 2011; Koch & Wagner 2011).

Characteristics of the HEAA

The accuracy of the HEAA can be assessed from the multiple linear regression for its three CpG markers and known age shown in Fig. 2A. The regression \( R^2 \) of 0.787
indicates that although most of the response can be attributed to age, there are other factors affecting CpG levels at the HEAA sites. The significant y intercept of 2.395 means that young whales will have their age slightly overestimated. The gradient of the HEAA regression shows that the age of older whales will be slightly underestimated.

The precision of the HEAA as assessed by Leave One Out Cross Validation (LOOCV) is shown in Fig. 2B. The overall precision of the HEAA was estimated as the standard deviation of the mean difference between known and estimated ages, which was 2.991 years. The distribution of the differences in known and estimated age was approximately normally distributed (Fig. S3A, Supporting information). A Shapiro–Wilk normality test demonstrates that the difference from normality is not significant (W = 0.984, P = 0.785) and this can be seen in a quantile–quantile plot (Fig. S3B, Supporting information). The differences in known and estimated age show little heteroscedasticity, meaning that variance in the differences is similar throughout the range of ages assayed (Fig. S3C, Supporting information) and there is an even dispersion of these values around the mean (Fig. S3D, Supporting information). The leverage effect of outlying points on the multiple regression was only minor, with all values for Cook’s Distance < 0.5 (Fig. S3E, Supporting information). The residuals of the 45 estimates had a mean of 3.575 years and the 95% confidence interval for age estimates was 8.947 years.

The precision of the PyroMark system alone was estimated from the standard deviation of six repeated measurements of the same four DNA samples to be 2.205 years. This indicates that a reasonably high proportion of the precision error is attributable to error in
measurement of methylation levels. This is likely to be because the percentage methylation differences measured in the CDKN2A_CpG+297 and GRIA2_CpG+202 assays are small (Fig. 1B and C).

**HEAA estimation of age in east coast Australian humpbacks**

The ages of 63 humpbacks sampled near Evans Head on their northbound migration along the east coast of Australia were measured with the HEAA. The results of these estimates are summarized in Fig. 3A. The estimates had a mean age of 10.01 and a range of 0–52. The decreasing number of animals in each age class approximated a negative exponential decrease (two sample Kolmogorov–Smirnov $D = 0.141, P = 0.245$). An equivalent negative exponential distribution of ages with a $\lambda$ rate parameter calculated by treating the observed mean as the expected value of the distribution ($\lambda = 1/10.01 = 0.0999$) is shown in Fig. 3B. This indicates that the HEAA produces age distributions that are close to the distribution expected for noncalf animals in a population where interannual fecundity is reasonably constant and there is little difference in mortality rate among adult age classes (Beverton, Holt, 1956). The proportion of the test population that was within the age range of the calibration samples (0–30 years) was calculated from this distribution to be 95.6%.

Ages estimated for east coast Australian humpbacks sampled in 2009 were compared to ages for humpbacks recorded during the operation of the Byron Bay whaling station from 1952 to 1962 and estimated by measurement of ear plug GLGs and ovaries (Chittleborough 1965) as shown in Fig. 4. The age structure from 1952 to 1960 represents an almost unexploited age structure for migratory whales found in this region (Chittleborough 1965), which is the same region where the Evans Head samples were collected. The humpback fishery in this region began to collapse in 1961 and 1962 was the final year of operations.

**Parent–offspring ordinal age prediction**

The ability of the HEAA to correctly order the age of a parent and its progeny was tested by simulation. A negative exponential population age distribution was simulated for the 2009 east coast Australian humpback whales based on HEAA data (Fig. 3B) and a distribution also simulated from the 1952 data to provide an example distribution from an unexploited population. Parent–offspring age intervals with a minimum of 5 years were
estimated from these distributions and the age estimate errors that would be produced by the HEAA simulated from its standard deviation. In 10 000 estimates of parent–offspring age order, the proportion in which the age order was correctly determined was 93.7% and 99.1% of cases for the 2009 and 1952 age profiles respectively (Fig. S3, Supporting information).

An empirical test of HEAA ordinal age estimation was made with eight mother–calf pairs from the ‘test’ sample set. The ‘calibration’ sample set also contained four mother–offspring pairs, which had age order estimated with the LOOCV approach. In all 12 cases, the correct mother–offspring age order was found.

Cross-species testing of the HEAA

We successfully assayed the three CpG sites in HEAA in six samples of Sperm Whale (Physeter catodon) DNA with ages ranging from 0 to 38 years. Of the three HEAA sites, the P. catodon orthologue of TET2_CpG+31 was the only one with a significant age-related DNA methylation regression ($R^2 = 0.927$, $P = 0.0013$). Increased age was associated with hypomethylation, as found in humpbacks.

Discussion

Molecular methods for estimating animal age are improving as understanding of the processes of biological ageing deepens. Telomere length assays are the most extensively studied molecular ageing method for animals, but with a few exceptions, these have not proven useful for population studies (Dunshea et al. 2011). In many animals, average telomere length varies extensively at birth, so although changes in individuals can be tracked over time, they are not useful for cross-sectional studies of population age (Kappei & Londono-Vallejo 2008; Karlsson et al. 2008). Other recent molecular age estimation methods that have performed well are ‘T Cell Receptor Excision Circle’ (TREC) qPCR for humans (Zubakov et al. 2010); changes in levels of specific mRNAs in mosquitoes (Cook et al. 2006); assays for specific CpG site methylation levels in humans (Bocklandt et al. 2011; Koch & Wagner 2011); and accumulation of specific lipids in whale blubber (Herman et al. 2009). All of these methods estimate a ‘biological’ age as the features they measure do not change with an annual trigger, but change with constant biological processes such as immunological insults in TREC qPCR (Zubakov et al. 2010); mRNA expression relating to developmental stage (Cook et al. 2006); telomere length changes resulting from number of mitotic cycles (Karlsson et al. 2008) or accumulation of dietary lipids in adipose (Herman et al. 2009). There is almost always population-wide variability in correlation between ‘biological’ age estimates and ‘chronological’ age because individuals within a population have different genotypes and experience different life histories. Even an extremely thorough age estimation assay such as genome-wide measurement of methylation at 70 387 age-related CpG sites (Hannum et al. 2013) results in an assay that has a standard deviation in age prediction of approximately 5% of the lifespan of the animal (humans) which is similar to the HEAA’s standard deviation = 3.0/lifespan of $\sim 95 \approx 3.1\%$. This indicates that the underlying relationship between chronological age and proxy markers for age is the limiting factor for achieving precision. Adding extra CpG sites to the HEAA would therefore not necessarily improve its predictive ability. In fact, two of the human epigenetic age assays used combinations of three CpG sites with the best relationships with age selected from a large number of possible age-related sites (Bocklandt et al. 2011; Koch & Wagner 2011). The differences in percentage methylation found in CDKN2A_CpG+297 and GRIA2_CpG+202 were quite small and repeated measurements of the same samples demonstrated that some of the variation in age-related methylation change can be attributed to the measurement error of the PyroMark system. Technologies for measuring DNA methylation are constantly improving and this suggests that measurement of these sites with deeper sequencing or other more accurate methodologies could further improve the precision of the HEAA.

Humpback whales are an excellent example of a species where age estimation is difficult. The prevalent method for age estimation in this species during the commercial whaling era was counting ear plug GLGs (Chittleborough 1959). However, there is measurement error in this method (Chittleborough 1959) and historical lack of consensus on how many growth layers accrue annually (Lockyer 1984). It was initially thought that two GLGs were produced each year (Chittleborough 1959) but more recent evidence suggests only one group (Gabriele et al. 2010), which is also consistent with the accrual rate in other baleen whales (Lockyer 1984; Gabrielle et al. 2010) and odd-numbered ear plug GLG counts (Chittleborough 1965; Gabriele et al. 2010). The most reliable method currently used to determine humpback age is visual re-identification of individuals that were first seen as calves. This approach clearly requires substantial effort and cannot be applied to populations without extensive records of past sightings. The only available method for estimating the age of live, previously unencountered humpback whales is analysis of blubber lipid profiles (Herman et al. 2009). This method has a similar estimated precision (standard deviation = 3.1–5.3 in different populations) to our HEAA (standard deviation = 3.0). A limitation of this approach is that it
requires calibration for each population as many of the accumulated lipids are derived from dietary items that are not uniformly consumed in different places. The HEAA does not require population-specific calibration and may therefore be particularly useful for comparing age data among populations.

Methods for measuring the age of long lived animals such as humpback whales are difficult to calibrate for the older portion of their range because there are few data available for validation. In this study, the oldest sample in the calibration set was from a 30-year old, which approaches the span of photo-identification research but is only about a third of the potential lifespan of the species. This relationship should be further evaluated for older whales as validating data become available, but all of the age-methylation relationships identified in other mammals so far are approximately linear (e.g. Maegawa et al. 2010; Bocklandt et al. 2011; Koch & Wagner 2011) and the relationships shown in Fig. 1 are also, which suggests that age estimates outside the calibration range will be correct relative to estimates within the calibration range. Even with currently available samples for calibration, the age range over which the HEAA was calibrated was estimated to cover more than 95% of humpbacks that will be encountered in the wild because older whales are expected to be rare (see Fig. 3).

The CpG sites that we use in the HEAA are in the first exons of humpback whale genes homologous to genes with known functions in other mammals. TET2 (ten eleven translocation 2) is a member of a multigene family that encodes DNA-binding proteins. It is involved in regulation of cytosine methylation of other genes and is a proto-oncogene (Branco et al. 2012). In humans, TET2 becomes hypermethylated with age (Grönninger et al. 2010), so it was interesting to observe significant, concerted age-related hypomethylation in humpback whales in four of the five CpG sites that we assayed and hypomethylation in the single sperm whale TET2 site assayed. CDKN2A (cyclin dependent kinase inhibitor 2A) is part of a gene complex with various names (CDKN2A/CDKN2B, p16INK4a/p16INK4b, ARF) for which there is also widespread evidence of age-related methylation changes (Gonzalez-Zulueta et al. 1995; Maegawa et al. 2010; Koch et al. 2011; Horvath 2013). CDKN2A mRNA expression levels have been proposed as a biomarker for human age, which is likely to be at least partly regulated by CpG methylation (Krishnamurthy et al. 2006). Glutamate receptors such as GRIA2 (glutamate receptor Ia2/AMP2A) are the predominant receptors in the mammalian brain and have a role in neuronal death associated with ageing (Chakrabarti et al. 2001). GRIA2 shows strong age-associated hypermethylation in humans (Koch & Wagner 2011).

As an example of the value of this technique, we estimated age structure for a suite of samples of unknown age obtained off the east coast of Australia and compared those results to historical whaling data. The population of humpbacks that migrate past the east coast of Australia was largely unexploited before 1952, so the age profiles for that time give an estimate of an unperturbed age structure (Chittleborough 1965). The fishery initially produced consistent annual yields and from 1952 to 1960 the average hunting time required to catch each whale was less than 2 h. In 1961 this increased to 4.5 h and in 1962 it had extended to 15 h as the fishery collapsed, ceasing to operate after 1962 (Chittleborough 1965) The population age structures for 1961 and 1962 reflect this depletion of adult whales. The 2009 population age structure estimated with the HEAA from this region is more similar to the age structures recorded in the final 2 years of the fishery than to those of the population encountered at the commencement of whaling. Our results suggest that 47 years after the cessation of whaling, the east coast Australian humpback population has not yet returned to a stable state. However, the modern sample size was small, the samples were not obtained for this purpose and the sampling did not necessarily replicate that of historical whaling operations. Biopsy samples were obtained during a portion of the northbound migration and humpback whales exhibit seasonal variation in migratory timing that varies with age, sex and reproductive status (Dawbin 1966). There may also be differences in selectivity between biopsy sampling and historical whaling operations. Despite these caveats, the strong bias towards young animals being represented and older ones being rare in our results is interesting and warrants further study. This analysis is included as an example of the sort of population comparisons that the HEAA allows, which will of course be more informative with larger sample sizes that are collected with the intent of assessing age structure. This population is growing rapidly (Chaloupka et al. 1999; Noad et al. 2011), so the 2009 age structure has probably resulted from high fecundity and survival, in contrast to the very similar age profiles for 1961 and 1962 that were caused by the size-dependent mortality imposed by the whale fishery. This illustrates a limitation of single ‘snapshot’ population age profiles in estimating underlying demographic parameters such as mortality and fecundity, as similar age profiles can result from changes in different parameters (De La Mare 1985).

The HEAA is more useful for improving demographic analysis methods based on mark–recapture or close-kin recognition. An assumption of many of the commonly used mark–recapture models is that all animals have an equal chance of recapture (Seber 1982). This assumption will be violated in many populations
where there is age-specific mortality. The HEAA enables estimation of mortality rates in different age classes. Ordinal age determination is especially useful for improving estimates of population size from close-kin recognition analyses (Skaug et al. 2010). Parent–offspring pairs can be identified by genotyping individuals with multiple markers such as microsatellites or SNPs, but it is often not possible to know which individual is the parent and which is the offspring. The power of close-kin population size estimation is diminished when the order of age is not known (Skaug et al. 2010). HEAA determination of age order from the same DNA sample that is used for genotyping will be particularly useful in this application.

Cytosine methylation is reasonably chemically stable and has been successfully assayed in ancient DNA purified from ~38 000-year-old Neanderthal bones (Briggs et al. 2010) and ~60 000-year-old bison bones (Llamas et al. 2012). This suggests potential for epigenetic age estimation from degraded DNA such as faecal samples, which would enable noninvasive sampling for age estimation material from whales or other vertebrates. Another clear application is age estimation from deceased animals. Age-specific population mortality rates could be derived from animals that die of natural causes (Sinclair 1977) as occurs in mass strandings of marine mammals (Barlow & Clapham 1997) and has been successfully assayed in ancient DNA purification material from whales or other vertebrates. Removal of deaminated cytosines and detection of in vivo methylation as highly conserved among mammal species as their associated gene and regulatory region DNA sequences and a plant (Yuan et al. 2014), so it is likely that specific CpG methylation in age-related sites can be identified for them as well. Epigenetic age estimation has great potential for expanding the scope of molecular studies of nonmodel organisms.

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A.P. performed experiments to develop the HEAA. J.R. collected samples for the calibration population. D.C. performed the PyroMark assays. A.P. and S.J. collected samples for the test population and jointly developed the concept for the HEAA. S.J. performed the data analyses. All authors contributed to writing the paper.

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Data Accessibility

All R scripts, DNA sequence data and CpG methylation data were deposited in the DRYAD database archive for this study (doi: 10.5061/dryad.h4b48). *Megaptera novaengliae* 5′ regulatory region sequences determined for six genes were deposited in GenBank and have accessions KF791963 – KF791968 (Table 1).

Supporting Information

Additional Supporting Information may be found in the online version of this article:

- **Fig. S1** Simulations for estimating performance of the HEAA in estimating age order.
- **Fig. S2** Co-plot of CpG methylation levels with age among seven sites considered for the HEAA.
- **Fig. S3** Assessment of normality of distribution of differences in known ages and ages predicted by our epigenetic age assay.
- **Table S1** Primers for the HEAA PyroMark assays.
- **Table S2** Akaike Information Criterion assessments of models with 25 different combinations of sites with CpG methylation levels relating to age in humpback whales.
- **Table S3** Correlation of age-related methylation in 18 pairwise combinations of CpG sites.