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The Bottlenose Dolphin Epigenetic Aging Tool (BEAT): A Molecular Age **Estimation Tool for Small Cetaceans**

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Age constitutes a critical parameter for the study of animal populations, providing information about development, environmental effects, survival, and reproduction. Unfortunately, age estimation is not only challenging in large, mobile and legally protected species, but often involves invasive sampling methods. The present work investigates the association between epigenetic modifications and chronological age in small cetaceans. For that purpose, DNA methylation at age-linked genes was characterized in an extensively studied, long-term resident common bottlenose dolphin (Tursiops truncatus) community from Sarasota Bay (FL, United States) for which sampled individuals have a known age. Results led to the identification of several CpG sites that are significantly correlated to chronological age in this species with the potential for sex to play a role in the modulation of this correlation. These findings have allowed for the development and validation of the "Bottlenose dolphin Epigenetic Age estimation Tool" (BEAT), improving minimally-invasive age estimation in free-ranging small cetaceans. Overall, the BEAT proved to be accurate in estimating age in these organisms. Given its minimally-invasive nature and potential large-scale implementation using skin biopsy samples, this tool can be used to generate age data from free-ranging small cetacean populations.

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INTRODUCTION

Dolphins and porpoises are small cetaceans that occur in all marine habitats around the globe, and are threatened by a wide range of human activities (Pompa et al., 2011; Davidson et al., 2012). Common bottlenose dolphins (Tursiops truncatus), the model organism used in this study, are wide-ranging animals (Dans et al., 2010; Wells and Scott, 2018) occurring in estuarine, coastal, shelf, and oceanic habitats in temperate, subtropical, and tropical waters around the globe. Unfortunately, because small cetaceans often associate in coastal or near coastal areas, they often are exposed to multiple concurrent anthropogenic stressors (e.g., pollution, boat traffic, bycatch in fishing gears, etc.) with potentially significant impacts on populations. These species are often difficult to study, and access to carcasses to collect biological data is relatively limited, particularly for wild migratory

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populations. Such problem has been alleviated by the progressive implementation of remote collection biopsies over the past two decades, improving our understanding of the population structure, feeding ecology, and ecotoxicology of free-ranging small cetacean populations from around the world (Fossi et al., 2000; Torres et al., 2003; Kiszka et al., 2011). Yet, the accurate estimation of chronological age, a critical population parameter, remains elusive from biopsy samples in these organisms.

Age constitutes a critically important parameter for population studies. Within a given population, several aspects of the biology of species can be investigated, including reproductive success, number of individuals of reproductive age, and the age composition of the population (Campana, 2001; Westgate and Read, 2007). In addition, age can be correlated to a variety of parameters, including pollutant loads, foraging tactics, and social structure. However, it is notoriously difficult to determine the age of bottlenose dolphins (as well as many other marine organisms), partly because they do not have conclusive external phenotypic traits that can be visually associated with age (Wilson et al., 1997; Miller et al., 2010; Hartel et al., 2015).

Current methods to estimate age in dolphins and other marine mammals include the use of body size, although this generally provides a rough estimation of the age group in which the animal may be included [e.g., adolescent versus adult (Baker et al., 2018)]. Another very accurate method includes removal of a tooth by a veterinarian, under local anesthesia, longitudinal sectioning, and quantification of growth layer groups in the tooth (International Whaling Commission, 1980; Hohn et al., 1989). This approach requires capture and handling of animals, which may not be feasible or desirable in some cases such as with endangered populations. Cost-effective, efficient, less-invasive alternative methods for obtaining accurate age estimates for large numbers of animals, without the need for capture, would be very useful. Biopsy dart sampling is a well-established technique for obtaining small tissue samples from cetaceans (Barrett-Lennard et al., 1996; Krutzen et al., 2002), and efforts have been made to use such samples for age estimation. Initial work has included the analysis of chromosomal telomere length and lipid accumulation in skin samples of cetaceans (Herman et al., 2009; Olsen et al., 2012). These techniques are limited by the strong influence of environmental conditions and for displaying a very wide range of values across individuals of similar chronological age. More specifically, telomere analyses can also be biased by the high variability in telomere lengths displayed by cetaceans (Olsen et al., 2012), as well as by its correlation with paternal age (Kappei and Arturo Londoño-Vallejo, 2008; Broer et al., 2013).

Complementary to traditional molecular analyses, the study of epigenetics [i.e., the study of phenomena and mechanisms that cause chromosome-bound, heritable (mitotically and/or meiotically) changes to gene expression that are not dependent on changes to DNA sequence (Deans and Maggert, 2015)] is emerging as a very promising framework to estimate chronological and biological age from DNA samples. Epigenetic age estimation (hereinafter referred to as aging) is based on the analysis of DNA methylation (addition of a methyl group to a Cytosine in the DNA sequence to form 5-methylcytosine). This modification is found at 60–80% of CpG sites (where a cytosine,

C, is next to a guanine, G, in the DNA sequence) across vertebrate genomes (Jones, 2001, 2012) both at gene bodies and promoters (Deaton and Bird, 2011), participating in the regulation of gene expression in response to developmental and environmental signals (Eirin-Lopez and Putnam, 2018). Most importantly, it has been shown that the %DNA methylation at specific positions of certain genes correlates linearly with chronological age, making age prediction reliable as demonstrated in the case of mammals including humans (Grönniger et al., 2010; Polanowski et al., 2014), as well as in other vertebrates (Paoli-Iseppi et al., 2019). This property has boosted the development of epigenetic aging tools that are now being successfully used in human forensics (Horvath, 2013; Shabani et al., 2018) as well as in marine organisms (Polanowski et al., 2014; Eirin-Lopez and Putnam, 2018).

In the present study we investigate the relationship between epigenetic modifications in age-responsive genes and chronological age in mammals, and their potential to develop an age evaluation tool for bottlenose dolphins and potentially other small cetacean species. For that purpose, genes displaying age-influenced changes in DNA methylation (*GRIA2*, *TET2*, and *CDKN2A*) were examined, in agreement with previous reports on humpback whales (Polanowski et al., 2014). In addition, we explored the potential for sex-specific differences in aging.

MATERIALS AND METHODS

Sample Collection

Skin samples from 39 bottlenose dolphins collected from June 2004 through May 2009 by the Sarasota Dolphin Research Program (Sarasota, FL, United States), as part of capturerelease health assessments of the long-term resident Sarasota Bay dolphin community (Wells et al., 2004, 2005) were used. This community has been studied since 1970, through at least six generations, and at any given time includes up to five concurrent generations within a single maternal lineage, and individuals up to 67 years of age (Irvine and Wells, 1972; Wells, 2014). As of 2015, the community was composed of 157.53 ± 43.05 SE dolphins (Tyson and Wells, 2016). In combination, health assessments, tagging and tracking, and monthly systematic photographic identification surveys have provided background information on the members of the resident community, including age, sex, genetic relationships, morphometrics, health, tissue concentrations of environmental contaminants, and reproductive histories (Wells, 2009). Skin samples were chosen as model biological material in the present study given the ability to perform samplings in large numbers of individuals in a minimally-invasive manner. This is further supported by previous reports describing age-related changes in DNA methylation in mammals including humans and humpback whales using skin tissue (Grönniger et al., 2010; Polanowski et al., 2014). Skin samples were obtained as subsamples of wedgeshaped surgical biopsies of blubber and skin collected under local anesthesia. The samples were frozen immediately in liquid nitrogen. Samples were stored at -80° C. Overall, the samples

analyzed were chosen to represent a group consisting of 15 females and 24 males with an even age distribution between sexes ranging from 2 to 36 years of age. However, for both males and females, fewer older animals were sampled than younger. Sex was determined by observation during capture-release efforts (see Figure 1 and Supplementary File S1).

DNA Extraction

DNA was extracted from skin samples using a phenol/chloroform extraction protocol, adapted from Sambrook et al. (1989). Accordingly, a small piece of skin (<2 mm) was cut into small pieces and placed in 500 µL of CTAB lysis buffer (100 mM Tris, 20 mM EDTA, 1.2M NaCl, 2% CTAB, pH 8.0) with 10 µL of proteinase K and 40 µL of Dithiothreitol (DTT). Samples were digested overnight at 50°C in a water bath, and subsequently vortexed and mixed with 600 µL of 4°C phenol/chloroform/isoamyl alcohol (25:24:1 mixture, Fisher BioReagents). Samples were vortexed again and centrifuged at 12,000 g, 4°C for 5 min. Approximately, 90% of the supernatant was removed and transferred to a new tube. Two additional cycles of extraction with phenol/chloroform/isoamyl alcohol were performed, placing the resulting supernatant fraction on a new tube and precipitating the DNA using 50 μL of ammonium acetate and 980 µL of 99% isopropyl alcohol at 4°C. After an overnight incubation on ice, samples were vortexed and centrifuged at 12,000 g, 4°C for 30 min to form a pellet. The isopropyl alcohol was subsequently removed and the pellet was subject to two consecutive washes using ice-cold 70% ethanol. The precipitate was completely dried using a vacuum centrifuge, DNA quality was assessed using agarose gel electrophoresis, and DNA concentration was calculated using Oubit fluorometric quantitation (Thermo Fisher Scientific).

Quantification of %DNA Methylation at CpG Sites

Freshly extracted DNA was divided into 500 ng aliquots (20 ng/ μ L) and subsequently subject to bisulfite transformation followed by pyrosequencing (EpigenDx, Hopkinton, MA, United States) to identify the %DNA methylation at individual

CpG sites, based on multiple DNA copies. Pyrosequencing yields approximately 20X coverage of each target sequence. Age-responsive candidate genes (*TET2*, *GRIA2*, and *CDKN2A*) were targeted. The selection of these genes was informed by previous reports identifying age-responsive genes in humans and particularly in a large cetacean species, the humpback whale (Polanowski et al., 2014).

Dolphin ortholog sequences for genes TET2, GRIA2, and CDKN2A were retrieved from the dolphin genome database (McGowen et al., 2012; Martinez-Viaud et al., 2019) and used to design primers using PyroMark software (Qiagen), specifically targeting most CpG sites present within the promoter sequences for these target genes (*TET2*, n = 4; *GRIA2*, n = 7; and *CDKN2A*, n = 6). Since *CDKN2A* is not annotated in the bottlenose dolphin genome, the killer whale (Orcinus orca) genome was used to verify this ortholog using the target sequence from the HEAA (Polanowski et al., 2014). The target sequences used and the CpG site location in relation to the start codon are detailed in Table 1 along with an explanation of CpG site labeling for this study. The %DNA methylation of gene promoters was calculated based upon the amount of fluorescence produced by PCR amplifications of pyrosequencing products at the targeted CpG sites. As a result, these analyses provide a final %DNA methylation for each of the samples at each of the CpG sites targeted for the three genes.

Statistical Analyses of BEAT Accuracy and Precision

Simple linear regression analyses were performed for each CpG site of each age-responsive gene using chronological age as the independent known variable and %DNA methylation for each CpG site at the different age-responsive genes as the dependent variable using R software (R Core Team, 2013, Version 3.5.2; R codes are included in **Supplementary File S2**). For each linear regression analysis, the coefficient of determination (R^2) was calculated, representing the percentage of the variation in the dependent variable (%DNA methylation) that is explained by the independent variables. After simple linear regression was performed for all CpG sites, multiple regression analyses were

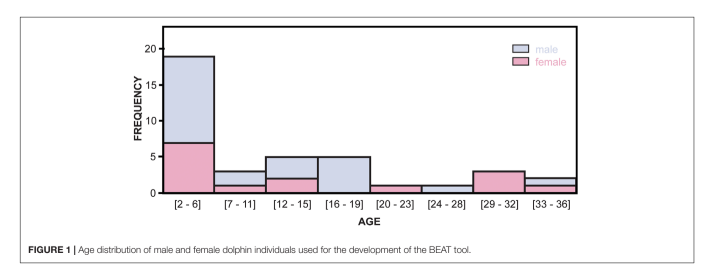


TABLE 1 | Location of the CpG sites identified by the present work at *TET2*, *GRIA2*, and *CDKN2A* genes.

Gene	CpG promoter location
TET2	GGTGGGC (TET2_1, -88384)CGGGG (TET2_2, -88379)CGGGGAGAAG (TET2_3, -88369)CGGGCCTGGGTCAAATTCCTAATTTGT (TET2_4, -88326)CGAGTCTTTAAAACTA
GRIA2	CCAGTCTC (GRIA2_1, -221)CGGACTT (GRIA2_2, -213)CG (GRIA2_3, -211)CGAG (GRIA2_4, -207)CGGGGAC (GRIA2_5, -200)CGGG (GRIA2_6, -196)CGCAGGG (GRIA2_7, -189)CGGCAGCCACCCGCAGGACCTTGGAAA
CDKN2A	(CDKN2A_1, -341)CGGAGGT (CDKN2A_2, -334)CGGAGT (CDKN2A_3, -328)CGGAGACCTCCTCT (CDKN2A_4, -314)CGG (CDKN2A_5, -311)CGA (CDKN2A_6, -308)CGCCTAGGGGGCTCAGGAAGCCACCGGAGGACTGAAAAG

CpG sites are indicated in boldface with their label and location (position of first nucleotide) in parentheses at the promoter region in respect to the transcription initiation site (negative numbers).

conducted. All possible combinations of significantly correlated CpG sites were tested. To avoid the effect of concerted DNA methylation changes among CpG sites on the same gene regions (Grönniger et al., 2010) only one CpG site from each gene was used in the combinations. This approach follows up on the strategy currently used for developing epigenetic age assays in other mammalian species (Koch and Wagner, 2011; Polanowski et al., 2014). Best models were defined based on Akaike information criterion (AIC) and Bayesian information criterion (BIC). The contribution of sex-specific traits to the epigenetic estimation of chronological age was also studied by discriminating between male and female samples in the analysis of %DNA methylation as a function of age. Thus, simple linear regression analyses for male and female datasets were additionally performed (see Supplementary Files S2, S3). The differences in the best epigenetic aging models obtained for male and female only analyses were evaluated using an analysis of covariance (ANCOVA) using R software ($\alpha = 0.05$). Lastly, the overall epigenetic aging model (including males + females) was validated through multiple regression analysis using the Leave-One-Out Cross Validation (LOOCV) method (Picard and Dennis Cook, 1984). This approach recurrently trains the model using all but one sample each time, until all samples had been left out once thereby predicting a value for each datapoint without using it to train the model.

RESULTS

CpG Methylation at Age-Responsive Genes in Bottlenose Dolphins

Overall, a total of 17 CpG sites were obtained across the three candidate genes targeted (TET2, n=4; GRIA2, n=7; CDKN2A, n=6) using pyrosequencing. The total %DNA methylation for all CpG sites at each of the genes analyzed showed a relatively low %DNA methylation, with gene average methylation falling between 1.6 and 12.2, with the highest percentage at a given site being 24.9 for an individual CpG

site at the promoter of CDKN2A_4. This was for a 2 year old female. The lowest amount of methylation was the absence of DNA methylation observed at several different sites for some individuals (Supplementary File S1). In addition, average %DNA methylation levels were calculated for each individual across all CpG sites for each gene. The individual with the highest average methylation for each gene was as follows: GRIA2 = female 189 age 32 (10.85 \pm 4.40); CDKN2A = female 189 age 32 (10.30 ± 2.80) ; *TET2* = female 217 age 2 (12.25 ± 5.20) . Similarly, the lowest averages were: GRIA2 = male 256 age 2 (1.60 \pm 1.60); $CDKN2A = \text{male } 252 \text{ age } 3 \text{ } (7.00 \pm 3.30); TET2 = \text{female } 193$ age 21 (4.30 \pm 3.00). The average %DNA methylation was also calculated for each gene, using average values across individuals for each gene. In this case, CDKN2A displayed the highest overall average (8.45 \pm 3.48; minimum: 4.35, maximum: 13.32), followed by TET2 (8.10 \pm 3.32; minimum: 3.86, maximum: 11.23) and *GRIA2* (3.96 \pm 1.78; minimum: 2.12, maximum: 6.91; Supplementary File S1). Globally, bottlenose dolphins displayed %DNA methylation levels at CpG sites in GRIA2 and CDKN2A genes similar to those displayed by humpback whales (Polanowski et al., 2014). Oppositely, %DNA methylation at TET2 CpG sites were lower by almost an order of magnitude in bottlenose dolphins (8.03 \pm 1.850, compared with humpback whales 14.630 ± 4.070 ; **Table 2**).

Correlation Between DNA Methylation and Chronological Age

Simple regression analyses of CpG methylation as a function of chronological age revealed that, out of the 17 CpG sites identified, only 4 of them (*TET2_4*, *CDKN2A_2*, *CDKN2A_4*, *CDKN2A_6*) were not significantly correlated with age, and therefore excluded from subsequent analyses (see **Table 3** for all *R*² values, see **Figure 2** for top correlated CpG site graphs). Multiple regression analyses were then conducted for all possible combinations of CpG sites displaying significant correlation with age, using one CpG site from each of the genes to ensure that independent age estimates were being made. This approach was implemented to account for the observations made in other mammals that CpG sites within a gene tend to be strongly correlated to one another, more often than CpG sites from different genes possibly because of similar processes affecting multiple CpG sites across a single gene (Koch and Wagner, 2011; Polanowski et al., 2014).

Following multiple linear regression analysis, the top model (based on the lowest AIC and BIC and highest R^2 values, $R^2 = 0.779$, residual error = 4.83, p < 0.001; **Table 4**) included two genes, GRIA2 CpG site 5 ($GRIA2_5$) and TET2 CpG site 2 ($TET2_2$), and is hereafter referred to as the BEAT (Bottlenose dolphin Epigenetic Aging Tool). These two CpG sites displayed the strongest correlation with chronological age among all CpG sites studied in these two genes (**Figure 2**). Full details on the other top models from multiple regression analysis (along with AIC and BIC scores), are shown in **Table 4**. The validation of the best model using the LOOCV method yielded predicted points at $R^2 = 0.740$ (root mean square error = 5.14) which were then plotted against the known age values, and had a strong correlation ($R^2 = 0.780$; **Figure 3**).

TABLE 2 | Comparison between %DNA methylation levels at studied CpG sites in the BEAT and the HEAA models.

	BEAT			HEAA		
	Average %DNA methylation ¹	Minimum	Maximum	Average %DNA methylation ¹	Minimum	Maximum
TET2	8.03	4.70	13.1	14.63	7.51	25.7
GRIA2	5.75	1.60	19.3	1.91	1.07	3.06
CDKN2A	4.41	0.00	6.7	2.39	1.03	4.83

¹ Averages were calculated by combining %DNA methylation at each individual for the top CpG site correlated to age and then dividing by the total number of individuals.

TABLE 3 | CpG sites identified at *TET2*, *GRIA2*, and *CDKN2A* genes, along with R^2 and p-values obtained in simple linear regression analyses.

CpG site	R ²	p-value	
TET2_1	0.0878	0.0669	
TET2_2	0.2962	0.0003	
TET2_3	0.3791	3.02×10^{-5}	
TET2_4	0.0642	0.1196	
GRIA2_1	0.7310	4.26×10^{-12}	
GRIA2_2	0.6082	4.87×10^{-9}	
GRIA2_3	0.6082	4.87×10^{-9}	
GRIA2_4	0.5933	9.83×10^{-9}	
GRIA2_5	0.7556	7.10×10^{-13}	
GRIA2_6	0.4806	1.00×10^{-6}	
GRIA2_7	0.6887	6.54×10^{-11}	
CDKN2A_1	0.3392	0.0001	
CDKN2A_2	0.0005	0.8870	
CDKN2A_3	0.4428	3.81×10^{-6}	
CDKN2A_4	0.0076	0.5956	
CDKN2A_5	0.3834	2.64×10^{-5}	
CDKN2A_6	0.0001	0.9621	

The following equation created from the multiple regression analysis is as follows: Age estimate = 8.602 + 4.392 (%DNA GRIA 5) – 1.149 (%DNA *TET2 2*).

The contribution of the sex of dolphin individuals to the epigenetic estimation of chronological age was evaluated using analysis of covariance (ANCOVA). While the results revealed a lack of significant differences between male and female changes in DNA methylation with age for the BEAT (p=0.760), simple linear regression analyses performed using only female and male datasets revealed differences for the two CpG sites used in the BEAT model (**Figure 4** and **Supplementary File S3** for all other sites). Most importantly, for site $GRIA2_5$, the slopes for males

TABLE 4 | Ranked list of the models best describing the relationship between %DNA methylation at CpG sites studied for *TET2*, *GRIA2*, and *CDKN2A* genes as a function of chronological age in bottlenose dolphins (male and female combined)

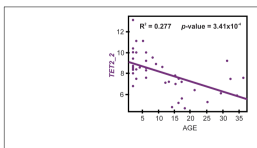
Ranking	Multiple regression	Adjusted R ²	p-value	AIC ¹	BIC ¹
1 (the BEAT)	TET2_2, GRIA2_5	0.7789	6.05×10^{-13}	238.32	244.98
2	TET2_2, GRIA2_5, CDKN2A_1	0.7779	3.78×10^{-12}	239.40	247.72
3	TET2_2, GRIA2_5, CDKN2A_5	0.7742	5.03×10^{-12}	240.04	248.36
4	TET2_2, GRIA2_5, CDKN2A_3	0.7729	5.55×10^{-12}	240.26	248.58
5	TET2_3, GRIA2_5, CDKN2A_1	0.7655	9.66×10^{-12}	241.51	249.82

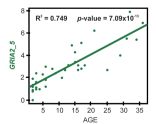
¹AIC, Akaike information criterion; BIC, Bayesian information criterion. The top model is referred as "the BEAT."

and females deviate from one another at older ages. For site $TET2_2$ we see that both male and female slopes are similar but males tend to have much less methylation than females. ANCOVA analyses for simple linear regressions did not find significant differences except for $TET2_2$ (p = 0.006), however the methylation of CpG sites at this gene did not display interaction with the sex of the individuals analyzed (p = 0.713).

DISCUSSION

Dolphins, as well as many other marine mammals, show relatively little to no visible signs of aging, therefore requiring the use of alternative aging methods. Unfortunately, the current methods used to estimate age in dolphins are limited due to logistical constraints including the need to capture and handle individuals (International Whaling Commission, 1980). Such limitation hampers monitoring, assessing, and studying age-dependent





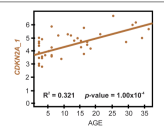


FIGURE 2 | CpG sites displaying the strongest correlation with chronological age across dolphin individuals in each of the three genes studied.

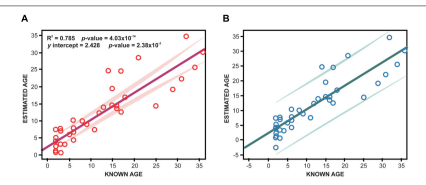
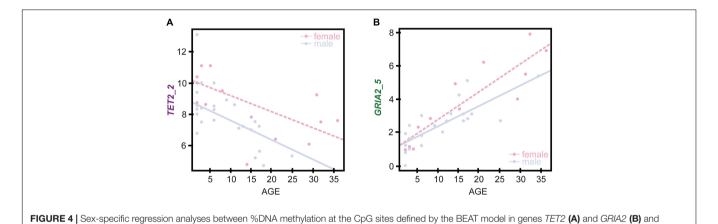


FIGURE 3 | Analysis of the precision of the BEAT model using **(A)** multiple regression analysis ($R^2 = 0.785$) and **(B)** its validation using LOOCV analysis (bottom graph, $R^2 = 0.740$). In **(A)**, the estimated age values were calculated for each individual using the equation generated by Multiple Regression analysis using all individuals to generate the model. In **(B)**, estimated age values were calculated for each individual using the equation generated by multiple regression analysis with the individual being estimated left out (see section "Materials and Methods").



effects of environmental stressors at the level of populations (Reeves, 2003; Westgate and Read, 2007; Agusa et al., 2008). Fortunately, studies in cetaceans can easily benefit from ongoing efforts creating more effective molecular aging tools in model systems such as humans or mice (Yi et al., 2014, 2015). In the present work, a similar experimental approach was used to investigate the relationship between epigenetic modifications in age-responsive genes and chronological age in mammals, and their potential to develop the BEAT (Bottlenose dolphin Epigenetic Aging Tool) to efficiently estimate chronological age using skin biopsy samples (Grönniger et al., 2010; Horvath, 2013; Polanowski et al., 2014). The BEAT is very similar to the HEAA [Humpback whale Epigenetic Aging Assay (Polanowski et al., 2014)] on its approach, but requiring only the analysis of CpG sites at two different genes, instead of the three genes required by the HEAA.

chronological age in females (open dots, dashed line) and males (solid dots and line).

Age-Responsive Genes Display Different Levels of CpG Methylation Across Dolphin Age Groups

The BEAT model defined in the present study is based on %DNA methylation information from two age-responsive genes

(GRIA2 and TET2) to determine chronological age. These genes were previously (and successfully) used to estimate age in other mammals, including North Atlantic humpback whales and humans (Koch and Wagner, 2011; Buscarlet et al., 2016). GRIA2 (glutamate receptor 2) belongs to a family of receptors that function in many neurophysiological processes (especially in the brain) and displays hypermethylation in older individuals (Koch and Wagner, 2011). TET2 (tet methylcytosine dioxygenase 2) encodes an epigenetic regulator mediating the transformation of 5-methylcytosine (5mC) into 5-hydroxymethylcytosine (5hmC) (Buscarlet et al., 2016), also functioning as a tumor suppressor that is heavily methylated in cancer tissues (Grönniger et al., 2010). Surprisingly, TET2 hypomethylation in older marine mammals contrasts with its hypermethylation in humans (Grönniger et al., 2010; Polanowski et al., 2014), underscoring the effects of the different constraints imposed by the regulatory role of DNA methylation on the function and subsequent longterm evolution of different genes across diverse ecosystems (Hernando-Herraez et al., 2015).

Overall, bottlenose dolphins displayed %DNA methylation levels at CpG sites in *GRIA2* and *CDKN2A* genes similar to those displayed by humpback whales (Polanowski et al., 2014). Oppositely, %DNA methylation at *TET2* CpG sites were

lower by almost an order of magnitude in bottlenose dolphins $(8.03 \pm 1.850$, compared with humpback whales 14.630 ± 4.070 ; **Table 2**). This difference underscores the importance of doing age assessment studies for different species and even developing species specific aging tools. Discriminating among the CpG sites used in the BEAT, the %DNA methylation at the GRIA2_5 site displayed a very high coefficient of determination for simple linear regression with age ($R^2 = 0.749$), which contrasts with the lower value found for TET2 2 ($R^2 = 0.277$), suggesting that other factors (besides age) may contribute to DNA methylation in these sites. Although the coefficient of determination obtained for the BEAT ($R^2 = 0.779$) is not as high as in similar tools developed for human forensics using high-resolution methods [e.g., $R^2 = 0.91$ (Yi et al., 2014)] this model can still be considered a massive improvement to previous methods used to estimate age in dolphins and small cetaceans, including additional room for improvement as the application of new methods becomes feasible (i.e., affordable) in non-model organisms.

The BEAT Tool Provides Epigenetic Age Estimations Based on Two Age-Responsive Genes

In total, 17 CpG sites located at three age-responsive genes (GRIA2, TET2, and CDKN2A) were studied to develop the BEAT. The best model correlating %DNA methylation with dolphin chronological age used a single CpG site in TET2 (CpG site 2) and GRIA2 (CpG site 5) genes (multiple regression $R^2 = 0.779$; LOOCV validation $R^2 = 0.740$, Figure 3) approximately 78% of the variation observed in %DNA methylation among dolphins is explained by their differences in chronological age. The remainder of the variation (22%) could be potentially due to other environmental factors (e.g., stress exposure) and/or be determined by genetic variability. Contrary to the HEAA assay, the BEAT tool did not require incorporating CpG sites from the CDKN2A gene into the model, although this gene was used by the second best correlation model (Table 4). Combined, these results suggest that the BEAT model might be less influenced on multiple age proxy markers to produce age estimates, reducing its dependence on genotypic or environmental variation, a key feature supporting its accuracy (Polanowski et al., 2014). Indeed, the methylation at the CpG site used in GRIA2 confers by itself a very strong correlation to dolphin chronological age (Figure 2), which is further improved by the incorporation of an additional CpG site from TET2.

The BEAT model displays a high correlation between %DNA methylation and age for the studied CpG sites (Figure 3 and Table 4). Yet, the accuracy of the model decreases slightly as dolphins age, as suggested by the increasing variance of estimated ages for older individuals [the same observation was made by studies attempting to create an age estimation tool for humans (Xu et al., 2015; Yi et al., 2015)]. Nonetheless, this effect does not seem to hamper the ability of the model to produce accurate estimations for a wide range of ages as validated by the LOOCV analysis (Figure 3) and by the fact that the error of the BEAT model is never higher than 5 years meaning that age estimates by the model will be within 5 years

of the actual age. Obviously, as in every model, the error can be reduced by increasing the sample size (notably by adding older individuals). Nonetheless, its present form constitutes a very important advancement from traditional aging methods used on these organisms (Hohn et al., 1989; Olsen et al., 2012). Further validation of the model using unknown wild specimens was not conducted in this study (but are currently underway) due to the similarity of this model to the HEAA where this type of validation was conducted and was found to be successful (Polanowski et al., 2014). Two different and complementary explanations might account for the observed dispersion. First, in the present study the age distribution of the samples is skewed toward young individuals (2–4 years, n = 14), compared with older individuals (4–36 years, n = 25; Figure 1 and Supplementary File S1). Although such distribution was also a potential problem for the estimations made by the HEAA assay in humpback whales (Polanowski et al., 2014), it can be easily corrected by progressively incorporating older individuals in the dataset to better fit the model. In the present case, the oldest individual incorporated into the BEAT tool was 36 years old, making the model reliable for most dolphins within the boundaries of the lifespan of most individuals (Wells and Scott, 2018).

Sex-Specific Differences Do Not Affect the Age Estimations Made by the BEAT Tool

The second explanation for the wider spread of data points around the regression line for older individuals refers to the potential increasing contribution of sex-specific traits to DNA methylation after dolphins reach sexual maturity. Indeed, mammals display critical differences in gene regulation between females and males, including epigenetic mechanisms [e.g., X chromosome inactivation, imprinting, etc. (Bar et al., 2019; Mutzel et al., 2019)], and their role during aging (Kornack et al., 1991; Berchtold et al., 2008; Arslan-Ergul and Adams, 2014). In the present work, the analysis of younger individuals (either females or males) revealed a better correlation between %DNA methylation and age (Figures 2, 3) when compared with older, sexually mature dolphins (age > 15 years) (Wells and Scott, 2018).

There have been several published studies investigating differences between sexes in aging and many other life functions (Garm et al., 2013; Soares et al., 2014; Fischer and Riddle, 2018). Given the role of epigenetic mechanisms regulating gene expression, it is not surprising that a growing number of studies are reporting sex-specific changes in DNA methylation occurring with age (Horvath, 2013; Horvath et al., 2016), highlighting the importance of including sex in DNA methylation studies. Therefore, in order to ascertain if sex plays a significant role contributing to the BEAT model, additional analyses were performed, discriminating between females (n = 15) and males (n = 24), including similar numbers of 2–4 year old individuals (females n = 6, males n = 8), as well as additional individuals evenly representing all other age groups (see **Figure 1**). The results suggest that the addition of sex to the model does

TABLE 5 | Ranked models best describing the relationship between %DNA methylation at studied CpG sites as a function of chronological age discriminating between female and male bottlenose dolphins.

Ranking	Multiple regression	Adjusted <i>R</i> ²	p-value	AIC ¹	BIC ¹
Female 1	TET2_4, GRIA2_5	0.8877	7.96×10^{-7}	90.85	93.68
Female 2	TET2_4, GRIA2_5, CDKN2A_1	0.8852	4.85×10^{-6}	91.87	95.41
Male 1	TET2_2, GRIA2_1, CDKN2A_5	0.9006	8.27×10^{-11}	120.21	126.10
Male 2	TET2_2, GRIA2_1, CDKN2A_3	0.8861	3.22×10^{-10}	123.48	129.37

¹AIC, Akaike information criterion; BIC, Bayesian information criterion.

not improve the BEAT's ability to estimate age (ANCOVA, p=0.760, **Figure 4**). Interestingly, when simple and multiple regression analyses were performed separately between females and males, the results revealed the presence of sex-specific differences in the CpG sites displaying higher correlations with %DNA methylation (see **Supplementary File S3**). Similarly, sex-specific multiple regression analyses defined different best models for each sex. Accordingly, the best female-specific model used CpG sites at two different genes ($GRIA2_5$ and $TET2_4$), whereas the male-specific model used CpG sites from all three genes investigated ($GRIA2_1$, $CDKN2A_5$, and $TET2_2$, see **Table 5**).

Overall, although the incorporation of sex does not seem to significantly improve the BEAT model, it does not discard its role in the rate of DNA methylation change between males and females. One possibility is that the small number of reproductive age individuals in the dataset may be hindering such contribution. Indeed, the simple linear regression models for *GRIA2_5* and *TET2_2* discriminating between females and males (**Figure 4**), show that regression lines share similar slopes, with the deviation being more pronounced for older individuals for both CpG sites. As samples from more known-age individuals of greater age become available, sex differences may become more evident and could possibly improve the accuracy of age estimation in cetaceans.

CONCLUSION

The present study investigates the relationship between epigenetic modifications in age-responsive genes and chronological age in bottlenose dolphins, using that information to develop an epigenetic age determination tool, the BEAT. The model underlying the BEAT is based on the %DNA methylation displayed by CpG sites in genes TET2 (CpG site 2) and GRIA2 (CpG site 5), estimating chronological age within 4.8 years of accuracy and with a high coefficient of determination ($R^2 = 0.779$). Overall, this tool provides a much needed alternative for feasibly estimating ages for large numbers of individuals in cetacean populations of interest. Furthermore, the ability to train the model with additional samples, especially

older individuals with equal representation of males and females, will foster the further development of the BEAT tool providing a finer estimation of age depending on sex and age-groups, and environmental factors that might influence biological age.

DATA AVAILABILITY

Data corresponding to dolphin age, sex, and %DNA methylation for the different CpG sites studied, R scripts and correlation analyses associated to the BEAT on which the present research is based, are deposited in the repository of the CREST-CAChE NSF Center at Florida International University and publicly available at github.com/eelabfiu/beat.

ETHICS STATEMENT

The animal study was conducted under National Marine Fisheries Service Scientific Research Permits, and approvals under Mote Marine Laboratory's IACUC.

AUTHOR CONTRIBUTIONS

AB designed the study, developed the experiments, analyzed the data, and wrote the manuscript. JK designed the study and wrote the manuscript. RW designed the study, provided the samples, and wrote the manuscript. JE-L designed the study, provided the materials, analyzed the data, and wrote the manuscript.

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Center in the Institute of Water and Environment at Florida International University.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars. 2019.00561/full#supplementary-material

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FILE S1 | Bottlenose dolphin dataset used in the present work, including: age, sex, and %DNA methylation for the different CpG sites studied in genes *TET2*, *GRIA2*, and *CDKN2A* across all individuals analyzed.

- **FILE S2** | R codes used for the development of the BEAT and tests for normality including outlier analyses. Residual plots and Normality Q–Q plots are also included.
- **FILE S3** | Individual linear correlation analyses for CpG sites displaying a significant correlation between %DNA methylation and age, calculated from whole dataset and discriminating between females and males.
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- **Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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