Research Paper

The telomere-mitochondrial axis of aging in newborns

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ABSTRACT

Aging starts at the beginning of life as evidenced by high variability in telomere length (TL) and mitochondrial DNA content (mtDNAc) at birth. Whether p53 and PGC-1 α are connected to these age-related markers in early life is unclear. In this study, we hypothesized that these hallmarks of aging are associated at birth.

In 613 newborns from the ENVIRONAGE birth cohort, p53 and PGC-1 α protein levels were measured in cord plasma, while TL and mtDNAc were measured in both cord blood and placental tissue. Cord blood methylation data of genes corresponding to the measured protein levels were available from the Human MethylationEPIC 850K BeadChip array. Pearson correlations and linear regression models were applied while accounting for selected covariates. In cord, a 10% increase in TL was associated with 5.22% (95% CI: 3.26 to 7.22; *p* < 0.0001) higher mtDNAc and -2.66% (95% CI: -5.04 to -0.23%; *p* = 0.032) lower p53 plasma level. In placenta, a 10% increase in TL was associated with 5.46% (95% CI: 3.82 to 7.13%; *p* < 0.0001) higher mtDNAc and -2.42% (95% CI: -4.29 to -0.52; *p* = 0.0098) lower p53 plasma level. Methylation level of TP53 was correlated with TL and mtDNAc in cord blood and with cord plasma p53 level.

Our study suggests that p53 may be an important factor both at the protein and methylation level for the telomere-mitochondrial axis of aging at birth.

INTRODUCTION

Aging is universal, unavoidable, and starts at the very beginning of life with an acceleration at middle-age. The general cause of aging is considered the timedependent accumulation of cellular damage [1-3]. The aging phenotype can be determined by cellular and molecular hallmarks that are generally considered to contribute to the aging process. Recently, primary hallmarks of aging were defined, which include genomic instability, telomere attrition, epigenetic alterations, loss of proteostasis, deregulated nutrient sensing, mitochondrial dysfunction, cellular senescence, stem cell exhaustion and altered intercellular communication [4]. These pathways have mainly been studied individually. A major challenge is to dissect the interconnection between the candidate hallmarks and their contribution to the aging-phenotype.

In an experimental study using mouse embryonic fibroblasts, Sahin et al. [5] revealed a direct connection between two primary hallmarks of aging i.e., dysfunctional telomeres which resulted in altered mitochondrial biogenesis and function via the tumor suppressor *TP53*, which in turn repressed peroxisome proliferator-activated receptor gamma, coactivator 1 alpha and beta (*PGC1A* and *PGC1B*), also known as master regulators of the mitochondria. Through observations from their experimental study, the "core axis of aging" was put forth, involving telomeres, mitochondria, p53 and PGC.

Until now, research on aging-mechanisms has mainly been limited to experimental research. Translation of these findings to population-based studies is scarce and focusses mostly on the older segment of the population [6]. However, human aging may start early and even from birth onward. It is therefore important to extent research on aging-mechanisms to the younger segment of the population. We evaluated the connection between TL and mtDNAc in 613 newborns from the ENVIRONAGE birth cohort and we evaluated whether p53 and PGC-1 α are on the path of this aging biomarker link as experimentally suggested.

METHODS

Study population

Mothers with a singleton full-term birth were selected from the ongoing population-based prospective ENVIRONAGE (ENVIRonmental influence ON early AGEing) birth cohort study, which is located in Limburg, Belgium. Detailed study procedures have been described previously [7]. Between 2010 and 2017, 1530 mother newborn pairs were recruited. In 691 samples, cord plasma protein levels (p53 and PGC-1 α) were measured. After removing 4 missing data, 18 outlying data (3 SDs from the mean) for protein levels and 10 outlying data for TL and mtDNAc, a total of 613 participants were used for statistical analysis. After delivery, mothers completed study questionnaires to provide detailed information on maternal age, paternal age, maternal education, smoking status, parity, and newborn's ethnicity. Maternal education was classified as "low" when mothers did not obtain any diploma, "middle" when they obtained a high school diploma, and "high" when they obtained a college or university diploma. Mothers were categorized as "never smoker", "former smoker" when they had quit smoking before pregnancy, and "smoker" if they had smoked at any time point during pregnancy. Parity was categorized in mothers having their first newborn, having their second newborn, or having their third or more newborn. Newborns were classified as either "European" when two or more grandparents were European, or as "non-European" when at least three grandparents were of non-European origin. Other maternal and perinatal parameters such as maternal pre-pregnancy Body Mass Index (BMI), newborns' sex, birth weight and birth date were collected from medical records after birth. Maternal pre-pregnancy BMI (kg/m²) was calculated based on data obtained during the first antenatal consultation. The date of conception was estimated by combining data on the first day of the mother's last menstrual period and the first ultrasonographic examination. The ENVIRONAGE study protocol has been conducted according to the Helsinki Declaration and was approved by the Ethical Committees of Hasselt University in Diepenbeek, Belgium (reference no. B371201216090 and B371201524537) and East-Limburg Hospital in Genk, Belgium. Written informed consent was provided by all participating mothers.

Sample collection and preparation

Procedures for umbilical cord blood and placental tissue collection for TL and mtDNAc assessment have been described in detail previously [7]. BD Vacutainer[®] Plus plastic whole-blood tubes with spray-coated K2EDTA (BD, Franklin Lakes, NJ, USA) were used to collect umbilical cord blood samples immediately after delivery. To obtain buffy coat and retrieve blood plasma, samples were centrifuged at 3,200 rpm for 15 min and stored separately. Within 10 minutes after delivery, placentas were collected and stored at -20° C. Four different placental biopsies (approximately 1- 2 cm^3) were taken at the fetal side at 4 cm from the cord umbilical and directly underneath the chorioamniotic membrane. Contamination by the chorioamniotic membrane was avoided by dissection followed by visual examination. All samples were stored at -80°C until DNA extraction. DNA was extracted using the QIAamp DNA Mini Kit (Qiagen Inc., Venlo, the Netherlands), according to the manufacturer's instructions. The quantity and purity of the sample was measured with the Nanodrop spectrophotometer (ND-1000; Isogen Life Science, the Netherlands). DNA samples were normalized to ensure a uniform DNA input of 5 ng for each qPCR reaction, and this was checked using the Quant-iT[™] PicoGreen[®] dsDNA Assay Kit (Life Technologies, Europe). Extracted DNA was stored at -80°C until further use.

Average relative TL and mtDNAc measurement

TL and mtDNAc were measured in cord blood buffy coat and placental tissue using a previously described modified quantitative real-time polymerase chain reaction (qPCR) protocol [8–10]. Details are provided in supplement (Supplementary Material). On each run, a 6-point serial dilution of pooled buffy coat or placental DNA was used to assess PCR efficiency as well as eight inter-run calibrators (IRCs) to account for inter-run variability. Non-template controls were also used in each run. All samples were measured in triplicate on a 7900HT Fast Real-Time PCR System (Applied Biosystems) in a 384-well format. qPCR curves for each sample were visually inspected and when technical problems were detected or triplicates showed too high variability, samples were removed for further analysis. Using gBase plus software (Biogazelle, Zwijnaarde, Belgium), all measurements were processed and normalized to a reference gene, taking into account run-to-run

differences. Average relative TL was calculated by determining the ratio of one telomere gene copy number (T) to one reference gene (36B4) (S). Average mtDNAc was calculated by determining the average of the ratio of two mitochondrial gene copy numbers (MTF3212/R3319 and MT-ND1) (M) to two nuclear reference genes (36B4 and β -Actin) (S). Telomere assay-reliability was assessed using an intra class correlation coefficient (ICC). The inter-assay ICC was 0.936 (95% CI: 0.808 to 0.969) and the intra-assay ICC was 0.952 (95% CI: 0.947 to 0.956).

p53, PGC-1a and SIRT-1 protein measurement

After thawing cord blood plasma, 100 µl of plasma was quantified for p53 protein levels (U/mL) using a Human p53 ELISA Kit according to the manufacturer's instructions (ref. ab46067, Abcam, Cambridge, United Kingdom). For PGC-1a (ng/mL), 100 µl of 1:250 diluted plasma was quantified according to the manufacturer's instructions (ref. E-EL-H1359, Elabscience, Texas, USA). Two commercially available Human SIRT1 ELISA Kits were tested for the detection of cord plasma SIRT-1 protein levels, but for none of them the limit of detection was reached (ref. ab171573, Abcam, Cambridge, United Kingdom and ref. E-EL-H1546, Elabscience, Texas, USA). All protein levels were measured in duplicate using a FLUOstar Omega microplate reader (BMG Labtech, Ortenberg, Germany). To minimize variability, all samples were randomized across 96 well plates and all ELISAs for one protein were measured in one day, while using the same serial dilution. Five IRCs per plate were taken into account to control for potential variability between plates. The intra-assay coefficient of variation was 5% for p53, and 10% for PGC-1a while the inter-assay coefficient of variation reached 14.5% for p53 and 25% for PGC-1a.

DNA methylation measurement

Cord blood DNA was used to determine the epigenomewide DNA methylation levels. DNA samples were bisulfite-converted, amplified and then hybridized to the Illumina Infinium Human MethylationEPIC 850K BeadChip array (Illumina, San Diego, CA, USA) at the GenomeScan lab (Leiden, The Netherlands). The array measurements were scanned using an Illumina iScan and the data quality was assessed using the R script MethylAid. DNA methylation data preprocessing, quality control, outlier detection, batch effect removal and probe filtering were described previously [11]. In total, 57 CpG loci with their UCSC reference gene name referring to TP53, PGC1A or SIRT1 were selected for the present study. Methylation levels of TP53, PGC1A and SIRT1 were available for 205 participants and used for further analysis. Details are provided in Supplementary Table 1.

Statistical analysis

All statistical analyses were performed using R studio version 3.6.2 (R Core Team, Vienna, Austria). Shapiro-Wilk's test was used to check the normality of the distributions. Average relative TL, mtDNAc and cord plasma protein levels were log₁₀-transformed to better approximate a normal distribution. For the descriptive statistics, continuous variables were presented as means \pm standard deviation (SD) and categorical variables as numbers (frequency in percentage). Pearson correlation was used to systematically evaluate following correlations: (1) TL in cord blood and placenta and mtDNAc in the respective tissue, (2) TL in cord blood and placenta and cord plasma protein levels, (3) mtDNAc in cord blood and placenta and cord plasma protein levels and (4) cord plasma protein levels. Multiple linear regression was applied to further confirm the associations independent of potential confounding effects. We adjusted for a priori selected covariates based on known associations of these factors with TL, mtDNAc, and protein levels as shown previously in multiple studies [8, 12]: Technical covariates (sample storage and batch effects), newborn's sex, gestational age, maternal BMI, maternal and paternal age, ethnicity, parity, smoke status, maternal education and month of delivery. All model estimates were presented as percentage difference with 95% CI and expressed for a 10% increment in explanatory variable.

In a secondary explorative analysis, we evaluated in a subpopulation whether cord blood methylation levels of genes corresponding to our measured protein levels (p53, PGC-1a and SIRT1) were related to the studied age-related markers and protein levels, to further explore the interrelationship of the telomeremitochondrial aging axis. We restricted these DNA methylation association studies to age-related markers measured in cord blood, as it is known that DNA methylation levels are highly tissue- and cell typespecific [13–15]. Since multiple CpGs were measured for each gene, we first made a correlation matrix showing the Pearson correlations between the CpGs for each gene. Second, we performed Principal Component Analysis (PCA) to reduce the dimensionality of our dataset [16]. To determine the number of Principal components (PCs) to retain in our analysis, we created scree plots for each gene (Supplementary Figure 1). PCs on the left side of the "elbow" of the graph were retained in the analysis. For all genes, this resulted in 5 PCs to retain. Next, we correlated the PCs with the CpGs, to determine their loadings (Supplementary Table 2). CpG loci were selected as relevant to a factor if the absolute value of their factor loadings were larger than 0.45 [16].

Characteristic	Mean \pm SD or <i>n</i> (%) (<i>n</i> = 613)
Mothers	
Age, y	29.3 ± 4.6
Pre-pregnancy BMI, kg/m ²	24.6 ± 4.8
Educational level	
Low	79 (12.9%)
Middle	227 (37.0%)
High	307 (50.1%)
Smoking status	
Never smoker	391 (63.8%)
Former smoker	154 (25.1%)
Current smoker	68 (11.1%)
Parity	
1	337 (55.0%)
2	206 (33.6%)
≥3	70 (11.4%)
Newborns	
Sex	
Female	321 (52.4%)
Gestational age, wk	39.2 ± 1.7
Birth weight, g	3420 ± 496
Ethnicity	
European-Caucasian	533 (86.9%)
Season of birth	
Winter	150 (24.5%)
Spring	149 (24.3%)
Summer	151 (24.6%)
Autumn	163 (26.6%)

Table 1. Descriptive characteristics of mother-newborn pairs from a subset (n = 613) of the ENVIRONAGE birth cohort.

First, these first five PCs were used to assess the Pearson correlations between methylation data and both cord blood age-related markers and cord plasma protein levels. Second, multiple linear regression was applied to confirm these associations while adjusting for the aforementioned covariates.

RESULTS

Study population characteristics

Demographic, pregnancy-related and perinatal characteristics of the mother-newborn pairs included in this study (n = 613) were summarized in Table 1.

Our study subset was representative for the original study population (Supplementary Table 3). Mothers were on average 29.3 (SD: 4.6) years old and had an average pre-pregnancy BMI of 24.6 (SD: 4.8) kg/m². Half of the participating women were highly educated (50%). The majority of the pregnant women never smoked cigarettes (63%), 25% stopped smoking before pregnancy, and 11% kept smoking on the average 3.4 cigarettes/day during pregnancy. Among the newborns, 52% were girls. Newborns had an average gestational age of 39.2 (SD: 1.67) weeks, an average birth weight of 3420 (SD: 496) grams, and most were of European origin (86%). Information about the age-related and protein markers is given in

Age-related marker	Mean ± SD	
Placenta		
TL (T/S ratio)	0.99 ± 0.26	
mtDNAc (M/S ratio)	1.05 ± 0.65	
Cord blood		
TL (T/S ratio)	0.99 ± 0.19	
mtDNAc (M/S ratio)	1.03 ± 0.57	
p53 plasma level (U/ml)	12.5 ± 9.72	
PGC-1α plasma level (μg/ml)	1145 ± 360	

Data presented as mean ± SD or number of participants (%). TL and mtDNAc are normalized separately in cord blood and placental tissue. Abbreviations: TL: telomere length; mtDNAc: Mitochondrial DNA content.

Table 2. Cord plasma SIRT-1 levels were excluded from the analysis since the measurements did not reach the limit of detection (LOD) of 0.31-0.63 ng/ml.

The telomere-mitochondrial axis of aging: links between TL, mtDNAc, p53 and PGC-1α

Unadjusted correlations

Cord blood and placental TL were positively correlated (r = 0.40, p < 0.0001, Supplementary Figure 2A), but no correlation between cord blood and placental mtDNAc was observed (Supplementary Figure 2B). The following correlations were systematically evaluated: (1) TL and mtDNAc, (2) TL and cord plasma protein levels, (3) mtDNAc and cord plasma protein levels and (4) between the cord plasma protein levels.

First, a positive correlation was found between TL and mtDNAc in both cord blood (r = 0.23, p < 0.0001, Supplementary Figure 3A) and placental tissue (r = 0.28, p < 0.0001, Supplementary Figure 3B).

Second, cord plasma p53 levels were negatively correlated with cord blood TL (r = -0.13, p = 0.0015, Supplementary Figure 4A) but not with placental TL (Supplementary Figure 4B). Cord plasma PGC-1 α levels were negatively correlated with both cord blood TL (r = -0.10, p = 0.013, Supplementary Figure 4C) and placental TL (r = -0.09, p = 0.029, Supplementary Figure 4D).

Third, no correlation was observed between cord plasma p53 levels and cord blood mtDNAc (Supplementary Figure 5A), but tended to be negatively correlated with mtDNAc in placenta (r = -0.07, p = 0.094, Supplementary Figure 5B). No correlation was observed between cord plasma PGC-1 α levels and cord blood mtDNAc (Supplementary Figure 5C), but a negative correlation was observed with placental mtDNAc (r = -0.11, p = 0.0082, Supplementary Figure 5D).

Fourth, cord plasma p53 and PGC-1 α levels were not correlated (Supplementary Figure 6). In Figure 1, a visual summary of all correlations is presented, which is based on the experimentally derived telomere-mitochondrial axis of aging hypothesis.

Covariate adjusted linear models

Using adjusted regression models, we further evaluated the associations between TL and (1) mtDNAc and (2) cord plasma protein levels (Table 3). We furthermore evaluated the associations between mtDNAc and the cord plasma protein levels, and also the association between the cord plasma protein levels (Supplementary Table 4). All associations were adjusted for technical covariates (sample storage and batch effects), newborn's sex, gestational age, maternal BMI, maternal and paternal age, ethnicity, parity, smoke status, maternal education and month of delivery. First, an increase in cord blood and placental TL was associated with higher mtDNAc in the respective tissue. A 10% increase in cord blood TL was associated with 5.22% (95% CI: 3.26 to 7.22; p <0.0001) higher cord blood mtDNAc and a 10% increase in placental TL was associated with 5.46% (95% CI: 3.82 to 7.13%; p < 0.0001) higher placental mtDNAc. Second, both cord blood and placental TL were associated with lower p53 levels, which is largely in line with the observed unadjusted correlations. A 10% increase in cord blood TL was associated with -2.66% (95% CI: -5.04 to -0.23%; p = 0.032) lower p53 plasma levels, while a 10% increase in placental TL was associated with -2.42% (95% CI: -4.29 to -0.52; p = 0.0098) lower p53 plasma levels. The observed correlations between TL and cord blood PGC-1a levels could not be confirmed in adjusted models. Furthermore, the correlations observed

	Cord TL (<i>n</i> = 603)		Placenta TL ($n = 558$)		
	% difference (95% CI)	<i>P</i> -value	% difference (95% CI)	<i>P</i> -value	
Cord mtDNAc	5.22 (3.26, 7.22)	< 0.0001	-0.45 (-1.96, 1.08)	0.56	
Placenta mtDNAc	1.02 (-1.06, 3.16)	0.34	5.46 (3.82, 7.13)	< 0.0001	
Cord p53	-2.66 (-5.04, -0.23)	0.032	-2.42 (-4.29, -0.52)	0.0098	
Cord PGC-1a	0.17 (-0.85, 1.21)	0.74	0.49 (-0.31, 1.29)	0.22	

Table 3. Association between TL in cord blood and placenta and (1) mtDNAc in the respective tissue and (2) cord plasma protein levels in the telomere-mitochondrial axis of aging.

Estimates are presented as percentage difference with 95% CI for a 10% change in explanatory variable. All models are adjusted for technical covariates (sample storage and batch effects), newborn's sex, gestational age, maternal BMI, maternal and paternal age, ethnicity, parity, smoke status, maternal education and month of delivery. Cord blood and placental Telomere Length represent the response variables, while the markers in the first column represent the exposure variables. Abbreviations: TL: telomere length; mtDNAc: mitochondrial DNA content.



Figure 1. Summary of the results found in this study integrated into the experimentally based telomere-mitochondrial axis of biological aging hypothesis. DNA damage and telomere shortening activate p53 leading to growth arrest, senescence or apoptosis. p53 might also impair mitochondrial function and mitochondrial DNA content indirectly through suppression of PGC-1 α – one of the master regulators of the mitochondria – leading to mitochondrial comprise and increased ROS levels, which leads to more DNA damage including telomere shortening. p53 and PGC-1 α could therefore be central players in the association between telomere length and mitochondrial DNA content and subsequently in the aging process. Solid lines represent significant associations between age-related or protein markers, while non-significant associations are represented by dotted lines. p53 and PGC-1 α levels were only measured in cord blood, while TL and mtDNAc were measured in both cord blood and placental tissue. Abbreviations: p53: tumor suppressor protein 53; PGC-1 α : peroxisome proliferator-activated receptor gamma co-activator 1 alpha protein. Figure based on the experimental work of Sahin et al. [5, 33]. between mtDNAc and the protein levels, nor between the protein levels themselves could not be confirmed in full adjusted regression models (Supplementary Table 4).

Cord blood TP53, PGC1A and SIRT1 methylation levels and the cord blood telomere-mitochondrial axis of aging

Unadjusted correlations

First, a Pearson correlation heatmap of the methylation levels of all CpGs for each gene (*TP53*, *PGC1A* and *SIRT1*, separately) is shown in Supplementary Figure 7. Second, the unadjusted Pearson correlation matrix between *TP53*, *PGC1A* and *SIRT1* methylation data (reflected by 5 PCs) and both cord blood age-related markers and protein levels in a subset of our study data (n = 205) is shown in Supplementary Figure 8. The first five PCs cumulatively explained 41%, 58% and 59% of the variation of *TP53*, *PGC1A* and *SIRT1*, respectively. In Supplementary Figures 9 and 10, significant correlations were additionally presented in correlations plots.

For TP53 methylation data, PC1 was negatively correlated with cord plasma p53 level (r = -0.17, p = 0.038), PC2 was positively correlated with cord blood TL (r = 0.20, p = 0.010) and PC5 was negatively correlated with cord blood mtDNAc (r = -0.16, p = 0.00073). The most relevant CpG to PC1 based on factor loadings (cg12041429, Supplementary Table 2) was located within the 5' untranslated region (5'UTR), meaning between the transcription start site (TSS) and the translation initiation codon (ATG) (Supplementary Table 1). The most relevant CpG for PC2 (cg18198734, Supplementary Table 2), was located in the gene body, meaning between the ATG and stop codon; irrespective of the presence of introns, exons, TSS or promoters (Supplementary Table 1). For PC5, absolute values of CpG factor loadings were not larger than 0.45, therefore, no CpG loci were selected as relevant to this factor.

For *PGC1A* methylation data, PC3 was positively correlated with cord blood TL (r = 0.18, p = 0.013). However, no CpG loci were selected as relevant to this factor.

For SIRT1 methylation data, no significant correlations were found with any of the measured markers.

Covariate adjusted linear models

The associations between *TP53*, *PGC1A* and *SIRT1* methylation data and cord blood age-related markers and cord plasma protein levels were further evaluated using regression models. All models were adjusted for technical covariates (sample storage and batch effects),

newborn's sex, gestational age, maternal BMI, maternal and paternal age, ethnicity, parity, smoke status, maternal education and month of delivery (Supplementary Table 5). The observed unadjusted correlations between methylation levels and both cord blood age-related markers and cord plasma protein levels could be confirmed after adjustment.

DISCUSSION

Aging is a complex universal and unavoidable physiological phenomenon. Both telomere attrition and mitochondrial damage are main factors in this biological process, but have mostly been studied independently. We [9, 17] and others [18–20] already observed large variability in TL and mtDNAc among newborns and we found that TL at birth predicts later life TL [17]. In the current study with 613 mothernewborn pairs, we evaluated in cord blood and placenta if TL is connected to mtDNAc, whether p53 and PGC- 1α are connected to these age-related markers and whether we may confirm the core axis of aging hypothesis in newborns.

Our study has two main findings: First, TL in cord blood and placenta is positively associated with mtDNAc in the respective tissue. Second, TL in cord blood and placenta is negatively associated with cord plasma p53 level, and the TL connection with p53 regulation is furthermore strengthened with the observed association with *TP53* methylation levels. These findings were confirmed after adjustment for potential confounding factors, and are in line with the proposed hypothesis as displayed in Figure 1. Our results partly confirmed the contribution of p53 in the telomere-mitochondrial axis of aging in early-life, however, the connection between p53 and mtDNAc and between the protein levels remains unconfirmed in our study.

Negative correlations between placental mtDNAc and both p53 and PGC1-α protein levels were found, but did not survive full adjustment for potential confounders. Also, p53 and PGC1- α were not correlated. However, methylation levels of the genes corresponding to the measured proteins show correlations with all markers that are in line with our hypothesis. As for the methylation levels, the highest contributing CpG loci were either located in the 5'UTR or in the gene body. In the 5'UTR, essential promotor elements are located and methylation of these promotor sequences will downregulate methylation, while gene-body methylation has been observed to be positively correlated with gene expression levels [21].

There was furthermore a strong positive association between cord blood and placental TL, but not between cord blood and placental mtDNAc, which may indicate that tissue-specific differences in mtDNAc are larger than tissue-specific differences in TL. This finding is in line with observations that TLs are highly correlated in different tissues within the same individual [17, 22]. For mtDNAc, the difference largely depends on the difference in energy demand of each cell type [23].

How can our observations be explained based on the proposed hypothesis (Figure 1)? DNA damage and excessive telomere shortening caused by intracellular stresses or environmental signals activate p53 [12, 24], which has commonly been referred to as the 'guardian of the genome' due to its role in protecting the cell from DNA damage and acting as a central hub in many biological downstream pathways [25, 26]. It regulates the expression of a variety of genes involved in different cellular functions, including cell-cycle regulation, apoptosis, DNA replication and repair, cell proliferation, cellular stress response and negative regulation of p53 [27]. Its activation modulates cellular senescence and organismal aging [28], whereas loss or mutation within the TP53 gene (which encodes p53) prevent cell death and increase cancer risk [29]. In this model, activated p53 directly suppresses PGC-1a, which alters the mitochondria (decreased mitochondrial function, impaired ATP generation and increased reactive oxygen species (ROS) production) which can lead to accelerated biological aging. Contradictory, there is growing evidence that p53 helps maintain the mitochondrial genome through translocation into mitochondria and interactions with mtDNA repair proteins. Park et al. [30] suggest that in unstressed cells, p53 functions as mito-checkpoint protein and regulates mtDNA copy number and mitochondrial biogenesis. Conversely, stress activated p53 (through DNA damage telomere shortening) results or in impaired mitochondrial biogenesis [30].

The findings of our study are supported by both experimental and population-based human studies. First of all, the link between telomeres and mitochondria was initially proposed in the experimental study of Sahin et al. [31], where telomerase-deficient mice (with a high level of dysfunctional telomeres) showed a strong activation of the TP53 gene, which resulted in suppression of PGC1A and PGC1B genes - the master regulators of mitochondrial biogenesis and metabolism - leading to comprised mitochondrial biogenesis. The link between these age-related markers was also made in several human studies [6, 32]. In a community sample of 392 healthy adults, Tyrka et al. [32] showed a positive correlation between leukocyte TL and mtDNAc, but the underlying mechanism remained undetermined. In 166 elderly, an association has been demonstrated between leukocyte TL and mtDNAc with *SIRT1* as a key role player in the telomere-mitochondrial interactome [6]. *SIRT1* expression was also shown to be inversely associated with *TP53* expression, which subsequently altered the expression of *PGC1A* [6].

What is the importance of our findings? Excessive telomere shortening to a critical length (Hayflick limit) [9] can lead to genome instability and senescence. It is also associated with an increased risk of age-related diseases, such as cardiovascular diseases [10], metabolic diseases [11] and cancer [12]. Mitochondria are the biochemical power plants of eukaryotic cells. Compared with nuclear DNA (nDNA), mtDNA is more vulnerable to damage induced by endogenous and exogenous agents. This due to the lack of protective histones, the small mitochondrial genome, its close proximity to the respiratory chain and its limited DNA repair system [15]. Mitochondrial damage might result in metabolic changes such as impaired ATP generation, increased production of ROS and decreased levels of ROS-detoxifying enzymes, which cause additional genomic instability as observed in aging and cancer [16]. By unraveling the mechanisms underlying the association between age-related markers in an early life context, we can further investigate how these important regulators may be influenced by early life exposures, and how this may lead to vulnerability for disease in later life. In addition, more research is needed to determine whether tracking and fixed ranking of TL among newborns, as evidence by Martens et al. [17], can be explained by variations of key-regulator levels in the core axis of aging. By strengthening our knowledge about tracking and ranking mechanisms involved in the complete axis of aging, improved measures can be taken to promote healthy aging across the life course.

Our study has several strengths. First of all, this is to our knowledge the first study that has investigated the key regulators of the telomere-mitochondrial axis of aging at birth. Second, we used a relatively large sample size (n = 613) within the ENVIRONAGE birth cohort, which is representative for the ENVIRONAGE birth cohort at large [7]. The third strength is the availability of different biological matrices (placenta, cord blood and cord plasma), in which the age-related markers were measured over different biological levels (TL, mtDNAc, proteins levels and methylation levels). This made it possible to investigate the telomeremitochondrial axis of aging independent of the biological matrix.

Besides our strengths, we also had to deal with some limitations. First, in addition to p53 and PGC-1 α , SIRT-1 is also considered to be a major contributor in the telomere-mitochondrial axis of aging pathway [6].

Unfortunately, this could not be confirmed in our study, since were not able to detect cord plasma SIRT-1 levels above the limit of detection of several commercially available ELISA kits. As evidenced by other studies [6, 33], SIRT-1 plays a role in the molecular axis of aging and might provide interesting information to confirm our hypothesis. A second limitation is that p53 and PGC-1 α were only measured in cord plasma but not in placental tissue, due to incompatibility with the measurement assays and due to limited placental tissue availability. As the aging process may be different for different tissues, the availability of placental protein levels of p53 and PGC-1a would provide more insight in the placental aging processes, this especially as placental tissue is a temporary end of life organ. Third, other non-studied markers playing a role in the telomere-mitochondrial axis of aging were not investigated and we can therefore not exclude that our findings are influenced by other important gene regulators. Fourth, our results only give an indication that p53 protein and methylation is linked with TL and mtDNAc.

In conclusion, we show that TL is connected to mtDNAc and that epigenetic and protein differences related to p53 might be involved in connecting these age-related markers at birth. This might be in line with the experimentally proposed telomere-mitochondrial axis of aging and gains important insight into the early life aging process.

AUTHOR CONTRIBUTIONS

TSN conceived and coordinates the ENVIRONAGE birth cohort and designed the current study together with DSM, BGJ and CVDS. BGJ and DSM performed telomere and ELISA measurements. RA, MP and SASL performed quality control of the methylation data. CVDS and DSM processed and statistically analyzed all data and performed the quality control of the database. CVDS, TSN and DSM wrote the first draft of the manuscript. All authors were involved in data interpretation and critical revision of the manuscript.

CONFLICTS OF INTEREST

The authors declare they have no competing financial interests. None of the funding agencies had a role in the design and conduct of the study; in the collection, analysis, and interpretation of the data; or in the preparation, review, or approval of the manuscript.

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SUPPLEMENTARY MATERIALS

Supplementary Methods

Reaction mixtures and PCR cycling conditions for telomere length (TL) and mitochondrial DNA content (mtDNAc) measurements

The master mix for TL measurement contained 1× OuantiTect SYBR Green PCR master mix (Oiagen, Inc., Venlo, the Netherlands), 2 mM dithiothreitol (DTT), 300 nM telg primer (ACACTAAGGTTTGGGTTTGGGTT TGGGTTTGGGTTAGTGT) and 900 nM telc primer (TGTTAGGTATCCCTATCCCTATCCCTATCCCTAT CCCTAACA). Used cycling conditions were: 1 cycle at 95°C for 10 min, followed by 2 cycles at 94°C for 15 sec and 49°C for 2 min and 30 cycles at 94°C for 15 sec, 62°C for 20 sec, and 74°C for 1 min and 20 sec. The single-copy gene reaction mixture contained $1 \times$ QuantiTect SYBR Green PCR master mix, 300 nM 36B4u primer (CAGCAAGTGGGAAGGTGTAATCC) and 500 nM 36B4d primer (CCCATTCTATCAT CAACGGGTACAA). Used cycling conditions were: 1 cycle at 95°C for 10 min, followed by 40 cycles at 95°C for 15 sec, and 58°C for 1 min and 20 sec [1].

The master mix for mtDNAc measurement consisted of Fast SYBR[®] Green I dye 2× (Applied Biosystems; 5 µL/reaction), 300 nm forward (5'-CACCCAAGAA CAGGGTTTGT-3' for MTF3212/3319, and 5'-ATGG CCAACCTCCTACTCCT-3' for MT-ND1) and 300 nm reverse primer (5'-TGGCCATGGGTATGTTGTTAA-3' for MTF3212/3319 and 5'-CTACAACGTTGGGGGCC TTT-3' for MT-ND1) and RNase free water (1.9 μ L/ reaction). thermal cycling profile was the same for all transcripts: 20 sec at 95°C for activation of the AmpliTag Gold[®] DNA-polymerase, followed by 40 cycles of 1 sec at 95°C for denaturation and 20 sec at 60°C for annealing/extension. Amplification specificity and absence of primer dimers was confirmed by melting curve analysis at the end of each run. After thermal cycling, raw data were collected and processed. CT (cycle threshold)-values of the two mitochondrial

genes were normalized relative to the nuclear reference genes according to the qBase software (Biogazelle, Zwijnaarde, Belgium). The program uses modified software from the classic comparative CT method ($\Delta\Delta$ CT) that takes multiple reference genes into account and uses inter-run calibration algorithms to correct for run-to-run differences [2, 3].

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Supplementary Figures



Supplementary Figure 1. Scree plots presenting eigenvalues of each Principal Component (PC). Eigenvalues represent the total amount of variance that can be explained by a given PC. Panel (A) displays the variance in *TP53* methylation explained by each PC. Panel (B) shows the variance in *PGC1A* methylation explained by each PC and panel (C) shows the variance in *SIRT1* methylation explained by each PC. Only PCs on the left side of the elbow in the curve were retained in the analysis.



Supplementary Figure 2. Correlation between cord blood and placental age related markers. Panel (A) shows the correlation between average relative cord blood TL and average relative placental TL. Panel (B) shows the correlation between average relative cord blood mtDNAc and average relative placental mtDNAc. Abbreviations: TL: telomere length; T/S: telomere/single copy gene ratio; mtDNAc: mitochondrial DNA content; M/S: mitochondrial DNA/single copy gene ratio.



Supplementary Figure 3. Correlation between TL and mtDNAc. Panel (A) shows the correlation between average relative cord blood TL and mtDNAc. Panel (B) shows the correlation between average relative placental TL and mtDNAc. Abbreviations: TL: telomere length; T/S: telomere/single copy gene ratio; mtDNAc: mitochondrial DNA content; M/S: mitochondrial DNA/single copy gene ratio.



Supplementary Figure 4. Correlation between TL and cord plasma protein levels. Panel (A) shows the correlation between average relative cord blood TL and cord plasma p53 level. Panel (B) shows the correlation between average relative placental TL and cord plasma p53 level. Panel (C) shows the correlation between average relative cord blood TL and cord plasma PGC-1 α level. Panel (D) shows the correlation between average relative placental TL and cord plasma PGC-1 α level. Abbreviations: TL: telomere length; T/S: telomere/single copy gene ratio.



Supplementary Figure 5. Correlation between mtDNAc and cord plasma protein levels. Panel (A) shows the correlation between average relative cord blood mtDNAc and cord plasma p53 level. Panel (B) shows the correlation between average relative placental mtDNAc and cord plasma p53 level. Panel (C) shows the correlation between average relative cord blood mtDNAc and cord plasma PGC-1 α level. Panel (D) shows the correlation between average relative placental mtDNAc and cord plasma PGC-1 α level. Abbreviations: TL: telomere length; T/S: telomere/single copy gene ratio; mtDNAc: mitochondrial DNA content; M/S: mitochondrial DNA/single copy gene ratio.



Supplementary Figure 6. Correlation between cord plasma p53 and cord plasma PGC-1α levels.



Supplementary Figure 7. Pearson correlation heatmap showing correlations between methylation levels of CpG regions for each gene separately. Methylation levels of *TP53*, *PGC1A* and *SIRT1* were obtained from the 850K array.

		Pearson Correlation													
						-1	.0 -0.5	0.0	0.5	1.0					
Cord plasma PGC-1α	0.04	-0.05	-0.01	-0.03	0.06	0.15	-0.06	-0.11	0	0.14	-0.02	0.05	-0.06	0.18	-0.04
Cord plasma p53	-0.17*	-0.13	-0.09	-0.09	0.05	-0.04	-0.05	-0.06	-0.08	0.08	0.05	0.01	-0.06	0	-0.08
Cord blood mtDNAc	-0.01	0.03	0.09	-0.04	-0.16*	-0.08	0	0.08	0.07	-0.08	-0.07	-0.13	-0.01	0.14	0.01
Cord blood TL	-0.09	0.2 *	-0.05	-0.1	-0.04	-0.09	0.05	0.18*	0	0.01	0.04	-0.05	0.02	-0.07	0.09
	PC1	PC2	PC3	PC4	PC5	PC1	PC2	PC3	PC4	PC5	PC1	PC2	PC3	PC4	PC5
		TP53	8 methyl	ation			PGC1	A methy	lation			SIRT	1 methy	lation	

Supplementary Figure 8. Pearson correlation matrix showing correlations between age-related markers and methylation levels in cord blood. Methylation levels of *TP53*, *PGC1A* and *SIRT1* were obtained from the 850K array. All age-related markers are log-transformed. A star indicates *p*-values ≤ 0.05 . Abbreviations: mtDNAc: Mitochondrial DNA content; TL: telomere length.



Supplementary Figure 9. Correlation between PCs of *TP53* methylation levels and cord blood age-related markers. Panel (A) shows a negative correlation between PC1 and cord plasma p53 level. Panel (B) shows a positive correlation between PC2 and average relative cord blood TL. Panel (C) shows a negative correlation between PC5 and average relative cord blood mtDNA content. Abbreviations: TL: telomere length; T/S: telomere/single copy gene ratio; mtDNAc: Mitochondrial DNA content; M/S: Mitochondrial DNA/single copy gene ratio.



Supplementary Figure 10. Correlation between PCs of *PGC1A* **methylation levels and cord blood TL.** A positive correlation is shown between PC3 and average relative cord blood TL. Abbreviations: TL: telomere length; T/S: telomere/single copy gene ratio.

Supplementary Tables

Supplementary Table 1. 850K methylation information. Annotations were obtained using the BiocManager package in R.

		850k	K methylation				
CpG Name	Annotation	Position	Relation to Island	UCSC Ref Gene Name	UCSC Ref Gene Accession	UCSC Ref Gene Group	Regulatory Feature Group
cg25053252	chr17:7589290-7589503	7589358	Island	TP53	NM_000546	5'UTR	PA
cg17461511	chr17:7591494-7591839	7591552	Island	TP53	NM_000546	TSS1500	PA
cg02045224	chr17:7591494-7591839	7591618	Island	TP53	NM_000546	TSS1500	PA
cg07991600	chr17:7589290-7589503	7589380	Island	TP53	NM_000546	5'UTR	PA
cg02087342	NA	7579047	Open Sea	TP53	NM_000546	Body	
cg22949073	NA	7572391	Open Sea	<i>TP53</i>	NM_000546	3'UTR	
cg13169780	chr17:7591494-7591839	7591570	Island	TP53	NM_000546	TSS1500	PA
cg07760161	chr17:7589290-7589503	7588378	N Shore	TP53	NM_000546	5'UTR	PA; CTS
cg02166782	chr17:7591494-7591839	7591564	Island	<i>TP53</i>	NM_000546	TSS1500	PA
cg22175811	chr17:7591494-7591839	7590052	N Shore	<i>TP53</i>	NM_000546	5'UTR	PA
cg21050342	chr17:7591494-7591839	7591644	Island	TP53	NM_000546	TSS1500	PA
cg13468400	NA	7579546	Open Sea	TP53	NM_000546	Body	
cg08119584	chr17:7591494-7591839	7589976	N Shore	TP53	NM_000546	5'UTR	PA
cg18198734	NA	7579263	Open Sea	TP53	NM_000546	Body	
cg23290969	chr17:7589290-7589503	7589272	N Shore	TP53	NM_000546	5'UTR	PA
cg16397722	NA	7577090	Open Sea	TP53	NM_000546	Body	
cg12041075	NA	7579312	Open Sea	TP53	NM_000546	Body	
cg08691422	chr17:7589290-7589503	7588787	N Shore	TP53	NM_000546	5'UTR	
cg15206330	chr17:7591494-7591839	7591591	Island	TP53	NM_000546	TSS1500	PA
cg12041429	chr17:7589290-7589503	7585875	N Shelf	TP53	NM_000546	5'UTR	
cg02842899	chr17:7589290-7589503	7589491	Island	TP53	NM_000546	5'UTR	PA
cg01620719	chr17:7589290-7589503	7589251	N Shore	TP53	NM_000546	5'UTR	PA
cg10792831	NA	7578689	Open Sea	TP53	NM_000546	Body	
cg06365412	NA	7580709	Open Sea	TP53	NM_000546	5'UTR	
cg27182172	chr4:23890120-23890955	23890890	Open Sea	PGC1A	NM_013261	Body	
cg09082664	chr4:23499647-23499989	23890701	Open Sea	PGC1A	NM_013261	Body	
cg27514608	chr4:23892305-23892675	23892540	Open Sea	PGC1A	NM_013261	TSS1500	
cg05158538	chr4:23891440-23891870	23891486	Open Sea	PGC1A	NM_013261	Body	
cg10955995	chr4:23857865-23858250	23858146	Open Sea	PGC1A	NM_013261	Body	
cg27461259	chr4:23892305-23892675	23892621	Open Sea	PGC1A	NM_013261	TSS1500	
cg07744449	NA	23887314	Open Sea	PGC1A	NM_013261	Body	
cg08550435	chr4:23829600-23829910	23829879	Open Sea	PGC1A	NM_013261	Body	
cg02896941	chr4:23499647-23499989	23890659	Open Sea	PGC1A	NM_013261	Body	
cg19701879	chr4:23880820-23881155	23881105	Open Sea	PGC1A	NM_013261	Body	
cg11270806	chr4:23892305-23892675	23892515	Open Sea	PGC1A	NM_013261	TSS1500	
-			-		_		

cg24716152	chr4:23879168-23881165	23879965	Open Sea	PGC1A	NM_013261	Body	
cg12691631	chr4:23891440-23891870	23891835	Open Sea	PGC1A	NM_013261	TSS200	
cg15541091	chr4:23812780-23813175	23813125	Open Sea	PGC1A	NM_013261	Body	
cg06772578	NA	23796624	Open Sea	PGC1A	NM_013261	3'UTR	
cg09427718	chr4:23819243-23820433	23819929	Open Sea	PGC1A	NM_013261	Body	
cg20723350	chr4:23891440-23891870	23891596	Open Sea	PGC1A	NM_013261; NM_013261	1stExon; 5′UTR	
cg10582690	chr10:69644169-69645178	69644422	Island	SIRT1	NM_012238	TSS200	PA
cg07832903	chr10:69644169-69645178	69644926	Island	SIRT1	NM_012238	Body	PA
cg00972288	chr10:69644169-69645178	69644348	Island	SIRT1	NM_012238	TSS200	PA
cg19681528	chr10:69644169-69645178	69644405	Island	SIRT1	NM_012238	TSS200	PA
cg02720207	chr10:69644169-69645178	69644399	Island	SIRT1	NM_012238	TSS200	PA
cg25513531	chr10:69644169-69645178	69644407	Island	SIRT1	NM_012238	TSS200	PA
cg26075202	chr10:69644169-69645178	69645177	Island	SIRT1	NM_012238	Body	
cg10705386	chr10:69644169-69645178	69644512	Island	SIRT1	NM_012238	1stExon	PA
cg01128800	chr10:69644169-69645178	69644938	Island	SIRT1	NM_012238	Body	PA
cg00851987	chr10:69662340-69664884	69664037	Open Sea	SIRT1	NM_012238	Body	
cg03251249	chr10:69644169-69645178	69645012	Island	SIRT1	NM_012238	Body	PA
cg05506809	chr10:69644169-69645178	69648299	S Shelf	SIRT1	NM_012238	Body	
cg19192316	chr10:69644169-69645178	69643020	N Shore	SIRT1	NM_012238	TSS1500	
cg23988827	NA	69676715	Open Sea	SIRT1	NM_012238	3'UTR	
cg19980381	chr10:69644169-69645178	69644128	N Shore	SIRT1	NM_012238	TSS1500	PA
cg13796676	chr10:69644169-69645178	69648687	S Shelf	SIRT1	NM_012238	Body	

Abbreviations: NA: not applicable; PA: Promotor Associated; CTS: Cell Type Specific.

Supplementary Table 2. Rotated loadings. Bold loadings have an absolute value larger than 0.45, meaning the CpG is selected as relevant for the specific PC.

<i>TP53</i>						
CpG	PC1	PC2	PC3	PC4	PC5	
cg01620719	-0.11	0.16	0.35	0.02	0.44	
cg02045224	-0.20	0.27	-0.07	0.17	-0.16	
cg02087342	0.55	0.27	0.06	-0.08	-0.16	
cg02166782	-0.01	0.32	0.31	0.27	-0.26	
cg02842899	-0.12	0.47	-0.05	0.04	-0.35	
cg06365412	0.37	0.36	-0.27	-0.06	0.42	
cg07760161	-0.01	0.18	0.60	-0.25	0.25	
cg07991600	-0.56	0.26	-0.22	0.20	0.32	
cg08119584	0.03	0.16	0.18	0.34	-0.23	
cg08691422	0.04	0.21	0.58	-0.16	0.04	
cg10792831	0.49	0.42	0.14	-0.32	0.03	
cg12041075	0.43	0.46	-0.24	-0.15	0.09	
cg12041429	0.72	-0.25	-0.02	0.24	0.09	
cg13169780	0.04	0.14	0.04	0.26	0.03	
cg13468400	-0.04	0.32	0.04	-0.44	-0.36	

2215206220	-0.07	0.22	0.36	0.20	-0.03
cg15206330		0.23			
cg16397722	0.54	-0.21	0.17	0.00	0.03
cg17461511	-0.42	0.44	-0.25	-0.15	0.13
cg18198734	0.35	0.61	-0.25	-0.06	-0.21
cg21050342	0.05	0.25	0.31	0.47	0.25
cg22175811	-0.17	0.17	0.38	-0.06	0.33
cg22949073	0.64	-0.10	-0.05	0.34	0.07
cg23290969	-0.03	0.20	0.22	0.00	-0.40
cg25053252	-0.62	0.29	-0.19	0.10	-0.01
0.0	DC1	PGC1A		DC4	D.C.F.
CpG	PC1	PC2	PC3	PC4	PC5
cg02896941	0.04	0.06	0.02	-0.07	0.02
cg05158538	0.08	0.07	-0.09	-0.10	-0.06
cg06772578	0.30	0.21	0.21	-0.12	-0.07
cg07744449	0.20	0.19	0.19	-0.09	-0.09
cg08550435	0.45	0.05	0.10	-0.08	-0.11
cg09082664	-0.04	-0.14	-0.14	0.07	-0.11
cg09427718	0.54	-0.05	0.10	0.06	0.08
cg10955995	0.20	-0.02	0.10	0.08	0.05
cg11270806	0.04	-0.02	-0.05	-0.08	0.04
cg12691631	0.06	0.03	-0.02	0.04	-0.16
cg15541091	0.20	0.32	0.20	-0.11	-0.19
cg19701879	0.48	-0.13	-0.14	0.06	-0.02
cg20723350	0.05	0.03	0.00	0.02	-0.09
cg24716152	0.29	-0.26	-0.22	0.14	0.02
cg27182172	-0.02	-0.07	-0.09	-0.01	0.03
cg27461259	0.04	0.12	0.01	-0.01	0.03
cg27514608	0.20	-0.11	-0.01	0.00	-0.08
		SIRT1			
CpG	PC1	PC2	PC3	PC4	PC5
cg00851987	-0.51	0.14	-0.10	-0.15	0.28
cg00972288	-0.01	0.28	-0.08	0.68	-0.22
cg01128800	0.69	0.45	-0.32	-0.08	-0.16
cg02720207	0.49	0.40	0.18	-0.17	0.04
cg03251249	0.00	-0.03	-0.22	0.59	0.48
cg05506809	-0.56	0.60	-0.12	0.13	-0.06
cg07832903	0.01	0.00	0.55	0.27	-0.62
cg10582690	0.73	0.34	-0.15	-0.21	-0.16
cg10705386	0.41	0.36	0.19	0.02	0.11
cg13796676	-0.66	0.57	-0.05	0.21	-0.01
cg19192316	-0.65	0.18	0.03	-0.07	0.25
cg19681528	0.35	0.23	0.25	-0.24	-0.08
cg19980381	-0.02	0.05	0.64	0.03	0.28
cg23988827	-0.46	0.61	-0.17	-0.09	-0.13
cg25513531	0.08	0.35	0.42	0.11	0.03
cg26075202	0.76	0.31	-0.27	-0.21	-0.14

Characteristic	Analyzed population Mean \pm SD or <i>n</i> (%) (<i>n</i> = 613)	Total population Mean ± SD or n (%) (n = 1530)	<i>P</i> -value	
Mothers				
Age, y	29.3 ± 4.6	29.4 ± 4.55	0.64	
Pre-pregnancy BMI, kg/m ²	24.6 ± 4.8	24.5 ± 4.76	0.66	
Educational level				
Low	79 (12.9%)	160 (10.5%)	0.13	
Middle	227 (37.0%)	500 (32.7%)	0.064	
High	307 (50.1%)	726 (47.5%)	0.32	
Smoking status				
Never smoker	391 (63.8%)	897 (58.6%)	0.030	
Former smoker	154 (25.1%)	326 (21.3%)	0.064	
Current smoker	68 (11.1%)	173 (11.3%)	0.95	
Parity				
1	337 (55.0%)	751 (49.1%)	0.015	
2	206 (33.6%)	507 (33.1%)	0.86	
2	70 (11.4%)	172 (11.2%)	0.95	
Newborns				
Sex				
Female	321 (52.4%)	736 (48.1%)	0.10	
Gestational age, wk	39.2 ± 1.7	39.2 ± 1.6	0.91	
Birth weight, g	3420 ± 496	3400 ± 495	0.52	
Ethnicity				
European-Caucasian	533 (86.9%)	1221 (79.8%)	0.0001	
Season of birth				
Winter	150 (24.5%)	372 (24.3%)	0.97	
Spring	149 (24.3%)	344 (22.5%)	0.40	
Summer	151 (24.6%)	329 (21.5%)	0.13	
Autumn	163 (26.6%)	385 (25.2%)	0.54	

Supplementary Table 3. Comparison of the characteristics between our study population (n = 613) and full ENVIRONAGE data (n = 1530).

Supplementary Table 4. Association between mitochondrial DNA content (mtDNAc) and cord plasma protein levels (p53 and PGC-1 α) and between the cord plasma protein levels.

	Cord p53 ($n = 61$	13)	Cord PGC-1a ($n = 607$)		
	% difference (95% CI)	<i>P</i> -value	% difference (95% CI)	<i>P</i> -value	
Cord mtDNAc	-0.0082 (-0.51, 0.50)	0.99	0.27 (-0.91, 1.47)	0.26	
Placenta mtDNAc*	-0.30 (-0.86, 0.27)	0.31	-0.74(-2.05, 0.58)	0.27	
Cord PGC-1a	0.12 (-0.21, 0.44)	0.51	NA	NA	

Estimates are presented as percentage difference with 95% CI for a 10% change in explanatory variable. All models are adjusted for technical covariates (sample storage and batch effects), newborn's sex, gestational age, maternal BMI, maternal and paternal age, ethnicity, parity, smoke status, maternal education and month of delivery. Cord plasma p53 and PGC-1 α represent the exposure variable, while the variables in the left column represent the outcome variable. ^{*}Data available for *n* = 575. Abbreviation: NA: not applicable.

Supplementary Table 5. Association between Principle Components (PCs) of *TP53*, *PGC1A* and *SIRT1* methylation levels and cord blood telomere length, cord blood mitochondrial DNA content (mtDNAc) and cord plasma protein levels (p53 and PGC-1 α).

	Cord TL	Cord mtDNA	Cord p53	Cord PGC-1a
	(n = 200)	(n = 205)	(n = 205)	(<i>n</i> = 205)
	% difference (95% CI)			
TP53				
PC1	-1.00 (-2.48, 0.50)	-0.48 (-4.43, 3.63)	-5.58 (-1.25, -0.66)	0.27 (-2.72, 3.35)
PC2	1.74 (-0.073, 3.58)	0.32 (-4.51, 5.39)	-4.63 (-9.79, 0.83)	-1.00 (-4.55, 2.68)
PC3	-1.38 (-3.36, 0.65)	-0.11 (-5.62, 5.71)	-3.23 (-9.73, 3.74)	0.13 (-3.90, 4.33)
PC4	-2.46 (-4.84, -0.024)	-2.96 (-9.28, 3.81)	-4.63 (-12.42, 3.85)	-0.20 (-5.10, 4.96)
PC5	-0.63 (-2.84, 1.64)	-10.52 (-15.59,-5.14)	2.59 (-5.16, 10.96)	0.77 (-3.79, 5.54)
PGC1A				
PC1	-0.31 (-1.88, 1.29)	-0.17 (4.06, 3.89)	-2.51 (7.33, 2.55)	1.15 (-1.82, 4.22)
PC2	-0.65 (-2.47, 1.21)	-0.34 (-5.24, 4.81)	-1.39 (-7.30, 4.89)	-0.26 (-3.82, 3.44)
PC3	2.20 (0.11, 4.34)	4.47 (-0.92, 10.78)	-0.18 (-7.08, 7.024)	-0.77 (-4.87, 3.52)
PC4	-0.77 (-2.99, 1.50)	2.50 (-3.38, 8.74)	-3.87 (-11.09, 3.93)	0.98 (-3.57, 5.74)
PC5	0.91 (-1.66, 3.54)	-3.98 (-10.38, 2.88)	3.58 (-5.27, 13.25)	2.71 (-2.55, 8.25)
SIRT1				
PC1	0.12 (-1.27, 1.52)	-2.43 (-5.96, 1.32)	0.73 (-3.80, 5.47)	-0.26 (-2.93, 2.48)
PC2	-0.41 (-2.25, 1.46)	-4.15 (-8.42, 0.80)	-0.76 (-6.84, 5.72)	-0.79 (-4.42, 2.97)
PC3	1.00 (-1.31, 3.37)	-1.18 (-6.97, 4.96)	-3.23 (-10.66, 4.81)	-2.57 (-7.04, 2.12)
PC4	0.52 (-1.99, 3.08)	4.20 (-2.30, 11.14)	0.48 (-7.79, 9.48)	4.41 (-0.70, 9.78)
PC5	1.10 (-1.63, 3.91)	0.11 (-6.92, 7.67)	-5.28 (-13.83, 4.12)	-0.77 (-6.16, 4.94)

Estimates are presented as percentage difference with 95% CI for a one-unit change in exposure variable. All models are adjusted for technical covariates (sample storage and batch effects), newborn's sex, gestational age, maternal BMI, maternal and paternal age, ethnicity, parity, smoke status, maternal education and month of delivery. The variables in columns represent the response variables, while the variables in the rows represent the explanatory variables. Bold values indicate significant estimates.