



Seven-CpG DNA Methylation Age Determined by Single Nucleotide Primer Extension and Illumina's Infinium MethylationEPIC Array Provide Highly Comparable Results

Valentin Max Vetter^{1,2}, Christian Humberto Kalies¹, Yasmine Sommerer³, Lars Bertram^{3,4} and Ilja Demuth^{1,5*}

¹Department of Endocrinology and Metabolic Diseases (Including Division of Lipid Metabolism), Biology of Aging Working Group, Charité-Universitätsmedizin Berlin, Corporate Member of Freie Universität Berlin and Humboldt-Universität zu Berlin, Berlin, Germany, ²Department of Psychology, Humboldt University Berlin, Berlin, Germany, ³Lübeck Interdisciplinary Platform for Genome Analytics (LIGA), University of Lübeck, Lübeck, Germany, ⁴Center for Lifespan Changes in Brain and Cognition (LCBC), Dept of Psychology, University of Oslo, Oslo, Norway, ⁵Charité - Universitätsmedizin Berlin, BCRT - Berlin Institute of Health Center for Regenerative Therapies, Berlin, Germany

OPEN ACCESS

Edited by:

Kai Tang,
Purdue University, United States

Reviewed by:

Osman A. El-Maari,
University of Bonn, Germany
Lei Gu,
Max Planck Institute for Heart and
Lung Research, Germany

*Correspondence:

Ilja Demuth
ilja.demuth@charite.de

Specialty section:

This article was submitted to
Epigenomics and Epigenetics,
a section of the journal
Frontiers in Genetics

Received: 16 August 2021

Accepted: 14 December 2021

Published: 17 January 2022

Citation:

Vetter VM, Kalies CH, Sommerer Y, Bertram L and Demuth I (2022) Seven-CpG DNA Methylation Age Determined by Single Nucleotide Primer Extension and Illumina's Infinium MethylationEPIC Array Provide Highly Comparable Results. *Front. Genet.* 12:759357. doi: 10.3389/fgene.2021.759357

DNA methylation age (DNAm age, epigenetic clock) is a novel and promising biomarker of aging. It is calculated from the methylation fraction of specific cytosine phosphate guanine sites (CpG sites) of genomic DNA. Several groups have proposed epigenetic clock algorithms and these differ mostly regarding the number and location of the CpG sites considered and the method used to assess the methylation status. Most epigenetic clocks are based on a large number of CpGs, e.g. as measured by DNAm microarrays. We have recently evaluated an epigenetic clock based on the methylation fraction of seven CpGs that were determined by methylation-sensitive single nucleotide primer extension (MS-SNuPE). This method is more cost-effective when compared to array-based technologies as only a few CpGs need to be examined. However, there is only little data on the correspondence in epigenetic age estimation using the 7-CpG clock and other algorithms. To bridge this gap, in this study we measured the 7-CpG DNAm age using two methods, via MS-SNuPE and via the MethylationEPIC array, in a sample of 1,058 participants of the Berlin Aging Study II (BASE-II), assessed as part of the GendAge study. On average, participants were 75.6 years old (SD: 3.7, age range: 64.9–90.0, 52.6% female). Agreement between methods was assessed by Bland-Altman plots. DNAm age was highly correlated between methods (Pearson's $r = 0.9$) and Bland-Altman plots showed a difference of 3.1 years. DNAm age by the 7-CpG formula was 71.2 years (SD: 6.9 years, SNuPE) and 68.1 years (SD: 6.4 years, EPIC array). The mean of difference in methylation fraction between methods for the seven individual CpG sites was between 0.7 and 13 percent. To allow direct conversion of DNAm age obtained from both methods we developed an adjustment formula with a randomly selected training set of 529 participants using linear regression. After conversion of the Illumina data in a second and independent validation set, the adjusted DNAm age was 71.44 years (SD: 6.1 years, $n = 529$). In summary, we found the results of DNAm clocks to be highly comparable. Furthermore, we developed an adjustment formula that allows for direct conversion of DNAm age estimates between methods

and enables one singular clock to be used in studies that employ either the Illumina or the SNUPE method.

Keywords: epigenetic clock, biological age, aging, methylation age, single-nucleotide primer extension assay, Berlin Aging Study II, BASE-II, GendAge study

INTRODUCTION

DNA methylation age (epigenetic clock, DNAm age) and its deviation from chronological age, DNAm age acceleration (DNAmAA), are novel and intensively studied biomarkers of biological aging. They are calculated from the methylation fraction of specific cytosine phosphate guanine (CpG) sites of genomic DNA. Numerous epigenetic clocks are available that differ in location and number of analyzed CpG sites and how these sites were selected. Their associations with mortality (reviewed in (Fransquet et al., 2019) and meta-analysis (Chen et al., 2016)), morbidity and age associated phenotypes (reviewed in (Jylhävä et al., 2017)) are well documented.

The most frequently used clocks include 71 (Hannum et al., 2013) or more CpG sites (Horvath, 2013; Levine et al., 2018) and therefore have to rely on epigenome-wide measurements, mostly carried out by Illumina's array-based "Infinium Methylation Assays". As an addition to these "big" epigenetic clocks, we recently reported a novel 7-CpG epigenetic clock whose underlying methylation data were measured by the methylation-sensitive nucleotide primer extension method (MS-SNUPE) (Vetter et al., 2018). This methodological approach was originally described by Kaminsky et al. (2005) and modified for the calculation of methylation age by Vidal-Bralo and others (Vidal-Bralo et al., 2016; Vidal-Bralo et al., 2017). When the interest is only seven CpGs this method is more cost effective compared to methylation arrays producing genome-wide methylation data. Consequently, the SNUPE method has been used in several studies (Vidal-Bralo et al., 2016; Banszerus et al., 2019; Vetter et al., 2020). However, the comparability of the findings described in these reports with reports relying on the identical CpG sites determined by an array-based genome-wide [e.g., Liu et al. (2020)] approach is uncertain.

This study aims to close this gap by measuring the methylation fractions of the regarding CpG sites with both the MS-SNUPE method and the "Infinium MethylationEPIC" array (Illumina Inc.) in a cohort of 1,058 adults (female: 52.6%). We subsequently compared the 7-CpG DNAm age of both methods. Due to the strong linear association of the results generated by both methods, we propose an adjustment formula that allows for direct conversion between 7-CpG methylation age measured with the EPIC array and the SNUPE method.

MATERIALS AND METHODS

BASE-II/GendAge Study

The multi-disciplinary and longitudinal BASE-II study aims to identify factors that are associated with "healthy vs. unhealthy" aging (Bertram et al., 2014; Gerstorf et al., 2016). The medical follow-up assessments took place between 2018 and 2020 and

were part of the GendAge study (Demuth et al., 2021). The current study included 1,058 GendAge participants with a mean age of 75.6 years (SD: 3.7 years, age range: 64.9–90.0 years, 52.6% female). For detailed information on BASE-II and GendAge please refer to Bertram et al. (2014) and Demuth et al. (2021).

The GendAge study was executed in accordance with the Declaration of Helsinki and approved by the ethics committee of the Charité—Universitätsmedizin Berlin (approval number: EA2/144/16). All participants gave written informed consent and GendAge is registered in the German Clinical Trials registry (DRKS00016157).

Seven-CpG Epigenetic Clock

The epigenetic clock used in this study employs seven CpG sites: cg09809672, cg02228185, cg19761273, cg16386080, cg17471102, cg24768561 and cg25809905. The formula was trained with SNUPE methylation data obtained from participants at baseline examination of the BASE-II (Vetter et al., 2018) and is referred to as "BII7"-formula. The described CpG sites were previously identified to be the most informative on chronological age and to be measurable in a SNUPE assay by Vidal-Bralo and others (Vidal-Bralo et al., 2016; Vidal-Bralo et al., 2017). An additional CpG site (cg10917602) was measured as well but is not included in the 7-CpG clock. Its results are shown in **Supplementary Figures S1–S3**.

DNA Methylation Assessment Using: Methylation-Sensitive Single Nucleotide Primer Extension

The analyzed genomic DNA was extracted from EDTA whole blood samples with the LGC "Plus XL manual kit", LGC, United Kingdom, and stored at -20°C . Briefly, 1,000 ng genomic DNA were bisulfite converted with the "EZ-96 DNA Methylation-Lightning Kit", Zymo Research. Subsequently, a multiplex PCR was conducted to amplify DNA sections surrounding the CpG sites of interest. The sample was cleaned from remaining oligonucleotides and dNTPs with "Shrimp Alkaline phosphatase", Affymetrix, and "Exonuclease I", New England Biolabs. The "SNaPshot Multiplex Kit", Applied Biosystems, was used for the single nucleotide primer extension (SNUPE). After an ultimate cleaning step with "Shrimp Alkaline phosphatase", the SNUPE-products were measured with an "3730 DNA Analyzer", Applied Biosystems and HITACHI. Raw data was inspected and processed with the "GeneMapper" software package, Applied Biosystems. The peak height was used to calculate the individual methylation fraction. For a more detailed description of the MS-SNUPE protocol, please refer to reference (Vetter et al., 2018) and (Vetter et al., 2021).

Extrinsic epigenetic age acceleration (EEAA) was calculated as residuals of a linear regression analysis of DNAm age on chronological age. Intrinsic epigenetic age acceleration (IEAA) was

TABLE 1 | Cohort characteristics.

	n	Mean	SD	Min	Max
Chronological age	1058	75.59	3.73	64.91	90.03
DNAmAge (BII7, SNUPE)	1058	71.15	6.86	41.65	98.86
EEAA (BII7, SNUPE)	1058	0.00	6.59	-30.14	27.84
IEAA (BII7, SNUPE)	1040	0.00	6.21	-24.71	25.43
DNAmAge (BII7, Illumina)	1058	68.09	6.37	46.00	91.20
EEAA (BII7, Illumina)	1058	0.00	6.17	-22.62	23.29
IEAA (BII7, Illumina)	1040	0.00	5.78	-20.46	21.01
DNAmAge (BII7adj, Illumina)	529	71.44	6.10	50.69	93.09
EEAA (BII7adj, Illumina)	529	0.00	5.95	-21.18	20.31
IEAA (BII7adj, Illumina)	519	0.00	5.52	-15.68	18.42
male					
Chronological age	502	75.49	4.00	64.91	90.03
DNAmAge (BII7, SNUPE)	502	72.23	6.72	41.65	95.29
EEAA (BII7, SNUPE)	502	1.13	6.41	-30.14	22.23
IEAA (BII7, SNUPE)	495	1.01	5.95	-24.71	20.14
DNAmAge (BII7, Illumina)	502	69.22	6.45	46.00	91.20
EEAA (BII7, Illumina)	502	1.17	6.20	-22.62	23.29
IEAA (BII7, Illumina)	495	1.01	5.76	-15.72	21.01
DNAmAge (BII7adj, Illumina)	254	72.52	6.29	50.69	93.09
EEAA (BII7adj, Illumina)	254	1.11	6.10	-21.18	20.31
IEAA (BII7adj, Illumina)	250	1.03	5.50	-14.36	18.42
female					
Chronological age	556	75.69	3.47	66.41	87.41
DNAmAge (BII7, SNUPE)	556	70.17	6.85	49.07	98.86
EEAA (BII7, SNUPE)	556	-1.02	6.59	-24.46	27.84
IEAA (BII7, SNUPE)	545	-0.92	6.30	-23.58	25.43
DNAmAge (BII7, Illumina)	556	67.07	6.12	48.86	88.19
EEAA (BII7, Illumina)	556	-1.06	5.95	-21.21	19.21
IEAA (BII7, Illumina)	545	-0.92	5.65	-20.46	17.27
DNAmAge (BII7adj, Illumina)	275	70.45	5.76	55.19	88.95
EEAA (BII7adj, Illumina)	275	-1.02	5.63	-16.73	17.19
IEAA (BII7adj, Illumina)	269	-0.96	5.37	-15.68	15.94

IEAA, Intrinsic Epigenetic Age Acceleration; EEAA, Extrinsic Epigenetic Age Acceleration; SNUPE, Single Nucleotide Primer Extension; BII7, BASE-II 7-CpG formula; BII7adj, BII7 adjustment formula for EPIC array (Illumina) data.

calculated as residuals of a linear regression analysis of DNAm age on chronological age and cell counts of neutrophils, monocytes, lymphocytes and eosinophiles. The latter is a modified version of the IEAA proposed by Quach and others (Quach et al., 2017).

DNA Methylation Assessment Using: Infinium MethylationEPIC Array

DNA methylation data was additionally obtained from the same DNA samples with the “Infinium MethylationEPIC” array by Illumina. Recommended default parameters were used for data pre-processing with the R-package “Bigmelon” (Gorrie-Stone et al., 2019) and R 3.6.1 (Team, 2017).

Briefly, probes were removed from the analyses if they had 1% or more samples with a detection p -value of 0.05 or a bead count below three in more than 5% of the samples. Outliers were identified by the *outlyx* and *pcout* (Filzmoser et al., 2008) function. Bisulfite conversion efficiency was estimated by *bscon* and samples with values < 80% were excluded from all following analyses. Subsequently, the samples were reloaded without the identified outliers and the function *dasen* was used for normalization. The amount of change in beta values due to normalization was determined by

the function *qual*. Samples with a root-mean-square deviation of ≥ 0.1 in beta value after normalization were removed and loading and normalization were repeated with the new sample set. DNAmAA with the Illumina data was calculated as EEAA and IEAA in the same way as described above.

Statistical Analysis

The statistical analyses and figures presented in this study were executed and designed with R 3.6.2 (Team, 2017) and the “ggplot2” package (Wickham, 2009). Bland-Altman plots and statistics were computed with the “blandr” package (Datta, 2017). Linear Regression analyses were calculated with R’s “lm” function. Methylation data of both methods was available in 1,087 of the older participants. Participants with difference between measurement methods of 3 SD or more were excluded from all analyses ($n = 29$). For the purpose of developing an adjustment formula converting the DNAm age from EPIC data into SNUPE data, we split the full BASE-II dataset as assessed in GendAge into a “training” and “validation” subsample. Individuals in either subset were randomly selected with the “sample” function in R. Throughout the study, statistical significance was assumed for p -values < 0.05.

RESULTS

GendAge Study Cohort Characteristics

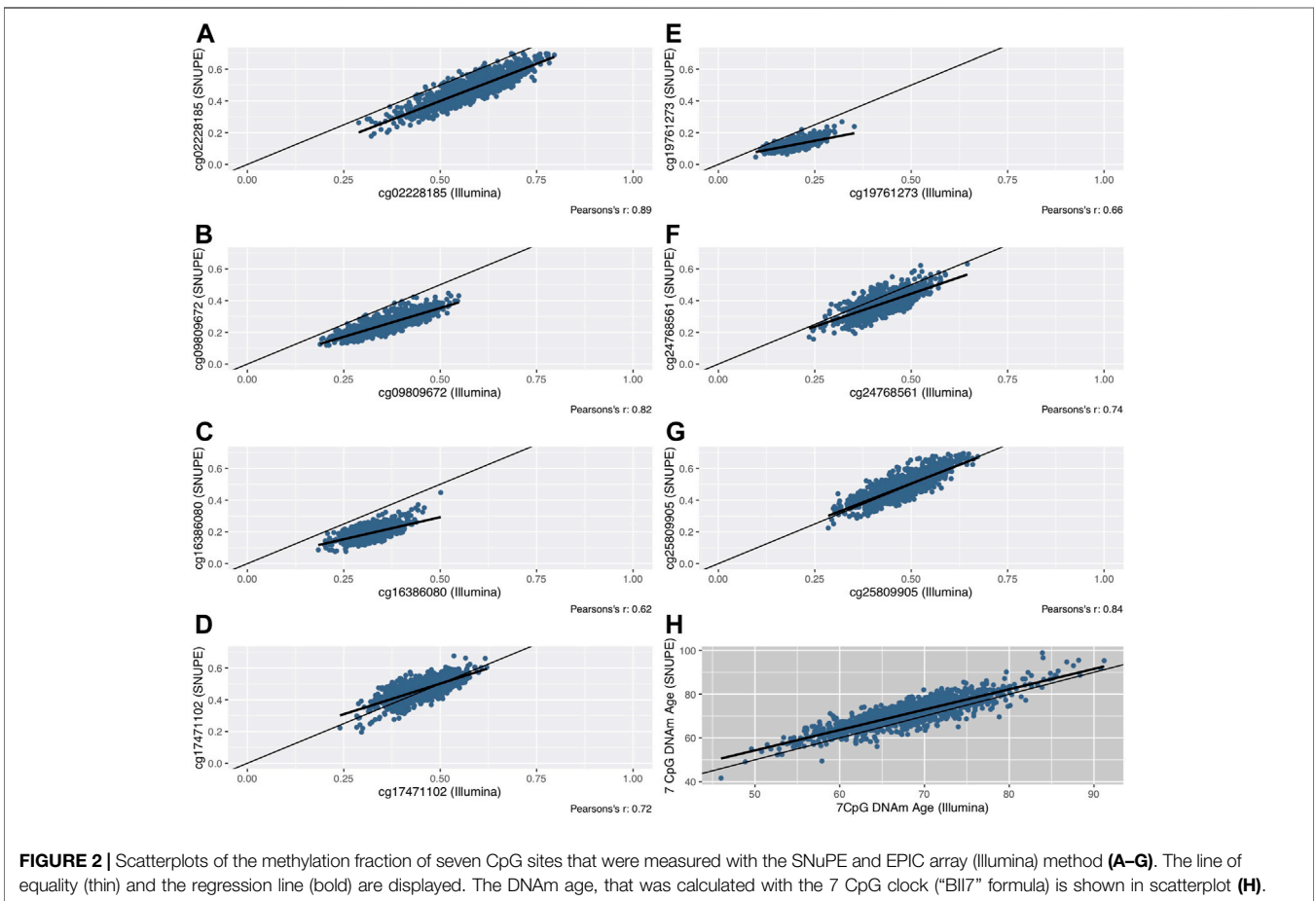
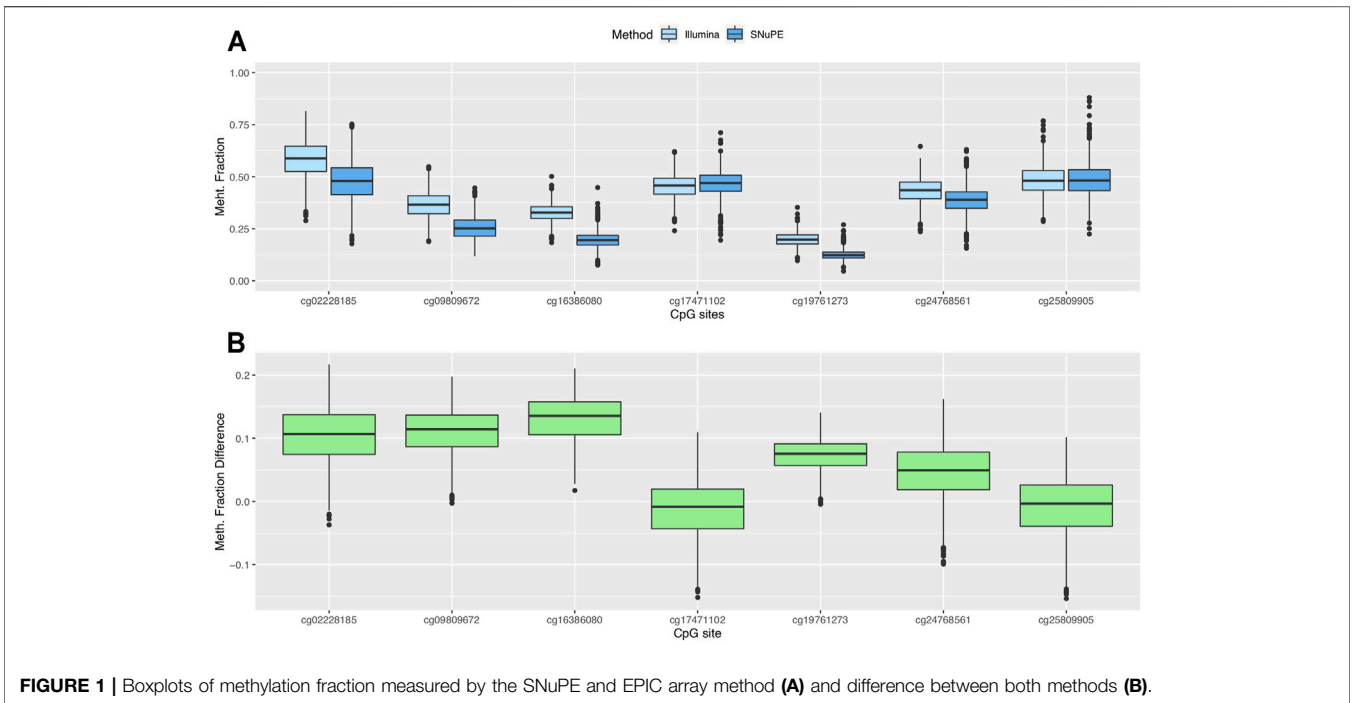
We analyzed 1,058 participants of the GendAge study which included the medical follow-up assessments of BASE-II participants. The chronological age of the participants ranged between 64.9 and 90.0 years and was on average 75.6 years (SD: 3.7) and 52.6% of the analyzed participants were female. Cohort characteristics are displayed in Table 1.

Comparison of DNA Methylation Fraction Measured by the SNUPE Method and Illumina’s EPIC Array

We measured the methylation fraction of the sites included in the 7-CpG clock (cg09809672, cg02228185, cg19761273, cg16386080, cg17471102, cg24768561 and cg25809905) in the same DNA samples with the SNUPE method and with Illumina’s “Infinium MethylationEPIC” array. The methylation fraction of cg1097602 was assessed as well but is not included in the 7-CpG clock. The results are shown in Supplementary Figures S1–S3.

Participants showed methylation fractions between 4.7% (cg19761273) and 88% (cg25809905) in SNUPE data and between 9.7% (cg19761273) and 81.5% (cg02228185) in Illumina data (Figure 1A). The average methylation range per CpG site was 44.9 percentage points (SNUPE) and 39.1 percentage points (Illumina).

The mean of the differences between methods ranged between 0.7 (cg25809905) and 13.1 (cg16386080) percentage points (Figure 1B). The smallest difference was found in CpG sites whose mean methylation fraction was close to 50%. The methylation fractions of individual CpG sites between methods



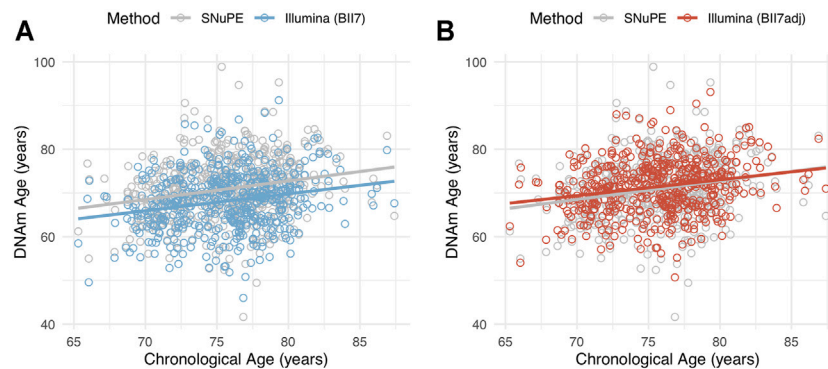


FIGURE 3 | Scatterplot of methylation age, measured with the SNUPE (grey) and EPIC array (Illumina) method (color) and calculated with the “BII7” (A) and “BII7adj” formula (B), on chronological age in the validation set ($n = 529$).

were moderately (cg16386080, Pearson’s $r = 0.62$) to highly correlated (cg02228185, Pearson’s $r = 0.89$) (Figures 2A–H).

To assess the limits of agreement and investigate a possible association between the measurement error and the methylation fraction, Bland-Altman plots were computed (Supplementary Figure S1). A regression analysis of the parameters analyzed in the Bland-Altman plots was generated to objectify a potential proportional bias. All plots showed marginal skewness at most ($|\beta| \leq 0.19$), except for one CpG site that showed a weak negative association ($\beta = -0.42$, cg19761273) between the difference of measurements and the mean of measurements. The latter was proposed by Bland and Altman as a substitute for the unknown true value (Martin Bland and Altman, 1986).

We found a mean difference between measurement methods of 7 percentage points or less in cg25809905, cg17471102, cg24768561 and cg19761273 (Figure 1B). The highest average difference in the methylation fraction between methods was 13 percentage points and was found in cg16386080. The limits of agreement (LOA), which were pinned by Bland and Altman (Martin Bland and Altman, 1986) as mean of difference ± 1.96 SD, contain by definition 95% of the values that represent the differences between methods. The range between the upper LOA (mean of difference $+1.96$ SD) and the lower LOA (mean of difference -1.96 SD) was 18.5 percentage points or smaller.

Seven-CpG DNAm Age Is Highly Correlated Between Methods

The DNAm age was calculated with the 7-CpG formula (“BII7”) that was trained on SNUPE methylation data (Vetter et al., 2018). It was measured to be on average 71.2 years by the SNUPE method and 68.1 years by the Illumina method. We found the DNAm age of men to be on average 2 years higher compared to women although the difference in chronological age was only 0.2 years, a finding that was reported for other epigenetic clocks as well (Horvath et al., 2016; Simpkin et al., 2016).

Both clocks were highly correlated with each other (Pearson’s $r = 0.86$). The deviation of DNAm age around the “line of equality”, indicating the potentially optimal line of identical results in both

methods, is shown in Figure 2H. The Bland-Altman plot (Supplementary Figure S1) shows a consistent bias of 3.1 years with a lower LOA of -3.8 and an upper LOA of 9.9 years. We found no proportional bias (linear regression analysis; $\beta = 0.08$).

Correlation Between DNAm Age and Chronological Age and New Formula for Conversion Between Methods

A moderate correlation was found between chronological age and DNAm age based on SNUPE data (Pearson’s $r = 0.28$) and Illumina data (Pearson’s $r = 0.25$). The slope of the regression line of DNAm age on chronological age was 0.51 (SNUPE) and 0.42 (Illumina).

To adjust the DNAm age obtained with EPIC array-based data, we computed a linear regression analysis of SNUPE-DNAM age on Illumina-DNAM age in a training set of 529 randomly selected participants. The resulting adjustment formula (“BII7adj”) allows for direct conversion between DNAm age estimates:

$$Y_{\text{DNAmAge(SNUPE)}} = 7.555 + 0.938 * X_{\text{DNAmAge(Illumina)}}$$

Descriptive statistics of the adjusted DNAm age obtained through EPIC array-based data in the validation set of our cohort ($n = 529$) are displayed in Table 1 and its association to chronological age is shown in Figure 3. We want to point out that this adjustment formula is primarily designed and evaluated for the conversion of DNAm age between methods and should not be used to convert singular methylation level of specific CpG sites.

DISCUSSION

In this study we compare two methods of DNAm age measurement to construct the 7-CpG DNAm clock, i.e. via the SNUPE method and via high-throughput DNAm profiling using the “Infinium MethylationEPIC” array, in 1,058 participants of the GendAge study. Although DNAm age estimates strongly correlated between

both methods, they showed a deviation of 3.1 years on average. Hence, we propose an adjustment formula to directly convert the DNAm age resulting from both methods to increase their comparability. These findings enable the use of the 7-CpG clock, that was originally developed for SNUPE data only, to be calculated with methylation data that was obtained with the EPIC array as well. To our knowledge, this is the only study to systematically compare these epigenetic age estimations from the 7-CpG clock using two different methodological approaches in one dataset.

To assess LOA between the methylation measurements of the individual CpG sites, we followed the established and widely used approach that was first proposed by Bland and Altman in 1986 (Martin Bland and Altman, 1986). To evaluate a possible association between the difference between measurements and the true value, which is unknown and therefore estimated as the mean between measurements, Bland-Altman plots were drawn. The range between the upper and lower LOA of the other CpG sites, that includes by definition 95% of the values that represent the differences between measurements, was 18.5 percentage points or smaller, a range which can be accepted for the purpose of DNAm age calculation.

Only one of the epigenetic clocks CpG sites (cg19761273) showed a moderate association between difference between measurements and mean of measurements. However, we found the smallest difference between methods in CpG sites, whose methylation fraction was close to 50%. We therefore assume a potential proportional bias between both methods which, however (almost) never becomes detectable in our cohort. That may be due to the comparatively small methylation ranges that occur in each individual CpG site *in vivo*. A series of experiments with DNA whose methylation fraction ranges between 0 and 100% would allow to test for proportional bias in every individual CpG site. Although easy to conduct, this experiment would not influence the evaluation of these methods with regard to calculation of methylation age where only smaller fractions seem to be apparent and to be unaffected by any potential proportional bias. One observation in our analyses relates to the difference in 7-CpG-derived DNAm age in men which on average was 2 years higher compared to women although the difference in chronological age between sexes in this dataset was only 0.2 years. This difference in DNAm age was reported in this (Banszerus et al., 2019; Lemke et al., 2022) and other clocks before (Horvath et al., 2016; Simpkin et al., 2016) and may reflect male-specific behavior (such as increased cigarette smoking or alcohol consumption) or may represent a bias due to other reasons and should be assessed further in independent datasets.

A limitation of this study is the age range of the analyzed cohort (65–90 years) which limits the evaluation of both methods to older adults. However, the mean chronological age of our cohort is close to that of many other cohorts analyzing epigenetic clocks (Marioni et al., 2015; Zhang et al., 2018; Jylhävä et al., 2019). Although the comparison of both methods would be undoubtedly interesting for a younger age range as well as for centenarians, we thought it most important to evaluate them in the area where they are most commonly used. Nevertheless, a comparison of methods in a cohort with a more even age distribution would be desirable. An external validation would be especially interesting, because the formula (“BII7”) that is used to calculate the 7-CpG clock was trained on the same participants but on samples taken on average 7.4 years

earlier (at baseline examination), therefore the individuals analyzed in this study are not independent from the 7-CpG training set. Another potentially limiting factor for comparison between methods is the efficiency of the bisulfite conversion step. While it was not quantified for the 7-CpG clock, it was quantified (and used as QC parameter) in the EPIC-based data. Given that the age estimates provided by both clocks correlate quite highly, we do not believe that incomplete bisulfite conversion has affected our results appreciably.

In conclusion, we report a good degree of agreement between the individual methylation fractions of analyzed CpG sites measured with the SNUPE method and with the EPIC array by Illumina. A difference of 3.1 years between the DNAm age estimations based on the different measurement methods was found, which can be (partially) corrected by our newly developed conversion formula. With this study, we aim to increase the comparability between the 7-CpG clock determined with the SNUPE method and higher throughput methods, such as the EPIC array. Further studies are needed to clarify whether the high degree of agreement between methods can be replicated in different/younger age groups as well.

DATA AVAILABILITY STATEMENT

The datasets presented in this article are not readily available. Data are available upon reasonable request. Interested investigators are invited to contact the study coordinating PI to obtain additional information about the GendAge study and the data-sharing application form. Requests to access the datasets should be directed to ID, ilja.demuth@charite.de.

ETHICS STATEMENT

The GendAge study was executed in accordance with the Declaration of Helsinki and approved by the ethics committee of the Charité—Universitätsmedizin Berlin (approval number: EA2/144/16). All participants gave written informed consent and GendAge is registered in the German Clinical Trials registry (DRKS00016157).

AUTHOR CONTRIBUTIONS

Conceived and designed the study: ID and VV. Contributed study specific data: all authors. Analyzed the data: VV. Wrote the manuscript: VV. All authors revised and approved the manuscript.

FUNDING

This work was supported by a grant of the Deutsche Forschungsgemeinschaft (grant number DE 842/7-1) to ID, the Cure Alzheimer’s Fund (as part of the “CIRCUITS-AD” consortium project) and the European Research Council’s “Horizon 2020” funding scheme (as part of the “Lifebrain” consortium project; both to LB). This article uses data from the Berlin Aging Study II (BASE-II) and the GendAge study which were supported by the German Federal Ministry of Education and

Research under grant numbers #01UW0808; #16SV5536K, #16SV5537, #16SV5538, #16SV5837, #01GL1716A and #01GL1716B.

ACKNOWLEDGMENTS

We thank Sanaz Sedghpour Sabet and Tanja Wesse as well as Andre Franke and Michael Wittig for their help in generating the EPIC array data used in this study. We also acknowledge the high-performance compute environment (“OmicsCluster”) at

University of Lübeck where the initial EPIC-data processing and analysis steps were run.

REFERENCES

- Banszerus, V. L., Vetter, V. M., Salewsky, B., König, M., and Demuth, I. (2019). Exploring the Relationship of Relative Telomere Length and the Epigenetic Clock in the LipidCardio Cohort. *Ijms* 20 (12), 3032. doi:10.3390/ijms20123032
- Bertram, L., Böckenhoff, A., Demuth, I., Düzel, S., Eckardt, R., Li, S.-C., et al. (2014). Cohort Profile: The Berlin Aging Study II (BASE-II)†. *Int. J. Epidemiol.* 43 (3), 703–712. doi:10.1093/ije/dyt018
- Chen, B. H., Marioni, R. E., Colicino, E., Peters, M. J., Ward-Caviness, C. K., Tsai, P. C., et al. (2016). DNA Methylation-Based Measures of Biological Age: Meta-Analysis Predicting Time to Death. *Aging (Albany NY)* 8 (9), 1844–1865. doi:10.18632/aging.101020
- Datta, D. (2017). *Blandr: A Bland-Altman Method Comparison Package for R*. Zenodo.
- Demuth, I., Banszerus, V., Drewelies, J., Düzel, S., Seeland, U., Spira, D., et al. (2021). Cohort Profile: Follow-Up of a Berlin Aging Study II (BASE-II) Subsample as Part of the GenAge Study. *BMJ Open* 11 (6), e045576. doi:10.1136/bmjopen-2020-045576
- Filzmoser, P., Maronna, R., and Werner, M. (2008). Outlier Identification in High Dimensions. *Comput. Stat. Data Anal.* 52 (3), 1694–1711. doi:10.1016/j.csda.2007.05.018
- Fransquet, P. D., Wrigglesworth, J., Woods, R. L., Ernst, M. E., and Ryan, J. (2019). The Epigenetic Clock as a Predictor of Disease and Mortality Risk: A Systematic Review and Meta-Analysis. *Clin. Epigenet* 11 (1), 62. doi:10.1186/s13148-019-0656-7
- Gerstorf, D., Bertram, L., Lindenberger, U., Pawelec, G., Demuth, I., Steinhagen-Thiessen, E., et al. (2016). Editorial. *Gerontology* 62 (3), 311–315. doi:10.1159/000441495
- Gorrie-Stone, T. J., Smart, M. C., Saffari, A., Malki, K., Hannon, E., Burrage, J., et al. (2019). Bigmelon: Tools for Analysing Large DNA Methylation Datasets. *Bioinformatics* 35 (6), 981–986. doi:10.1093/bioinformatics/bty713
- Hannum, G., Guinney, J., Zhao, L., Zhang, L., Hughes, G., Sada, S., et al. (2013). Genome-wide Methylation Profiles Reveal Quantitative Views of Human Aging Rates. *Mol. Cell* 49 (2), 359–367. doi:10.1016/j.molcel.2012.10.016
- Horvath, S. (2013). DNA Methylation Age of Human Tissues and Cell Types. *Genome Biol.* 14 (10), R115. doi:10.1186/gb-2013-14-10-r115
- Horvath, S., Gurven, M., Levine, M. E., Trumble, B. C., Kaplan, H., Allayee, H., et al. (2016). An Epigenetic Clock Analysis of Race/ethnicity, Sex, and Coronary Heart Disease. *Genome Biol.* 17 (1), 171. doi:10.1186/s13059-016-1030-0
- Jylhävä, J., Jiang, M., Foebel, A. D., Pedersen, N. L., and Hägg, S. (2019). Can Markers of Biological Age Predict Dependency in Old Age?. *Biogerontology* 20 (3), 321–329. doi:10.1007/s10522-019-09795-5
- Jylhävä, J., Pedersen, N. L., and Hägg, S. (2017). Biological Age Predictors. *EBioMedicine* 21, 29–36. doi:10.1016/j.ebiom.2017.03.046
- Kaminsky, Z. A., Assadzadeh, A., Flanagan, J., and Petronis, A. (2005). Single Nucleotide Extension Technology for Quantitative Site-specific Evaluation of met/C in GC-Rich Regions. *Nucleic Acids Res.* 33 (10), e95. doi:10.1093/nar/gni094
- Lemke, E., Vetter, V. M., Berger, N., Banszerus, V. L., König, M., and Demuth, I. (2022). Cardiovascular Health Is Associated With the Epigenetic Clock in the Berlin Aging Study II (BASE-II). *Mech. Ageing Dev.* 201, 111616. doi:10.1016/j.mad.2021.111616
- Levine, M. E., Lu, A. T., Quach, A., Chen, B. H., Assimes, T. L., Bandinelli, S., et al. (2018). An Epigenetic Biomarker of Aging for Lifespan and Healthspan. *Aging* 10 (4), 573–591. doi:10.18632/aging.101414
- Liu, Z., Leung, D., Thrush, K., Zhao, W., Ratliff, S., Tanaka, T., et al. (2020). Underlying Features of Epigenetic Aging Clocks *In Vivo* and *In Vitro*. *Aging cell* 19 (10), e13229. doi:10.1111/acel.13229
- Marioni, R. E., Shah, S., McRae, A. F., Ritchie, S. J., Muniz-Terrera, G., Harris, S. E., et al. (2015). The Epigenetic Clock Is Correlated with Physical and Cognitive Fitness in the Lothian Birth Cohort 1936. *Int. J. Epidemiol.* 44 (4), 1388–1396. doi:10.1093/ije/dyu277

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fgene.2021.759357/full#supplementary-material>

- Martin Bland, J., and Altman, D. (1986). Statistical Methods for Assessing Agreement between Two Methods of Clinical Measurement. *The Lancet* 327 (8476), 307–310. doi:10.1016/s0140-6736(86)90837-8
- Quach, A., Levine, M. E., Tanaka, T., Lu, A. T., Chen, B. H., Ferrucci, L., et al. (2017). Epigenetic Clock Analysis of Diet, Exercise, Education, and Lifestyle Factors. *Aging* 9 (2), 419–446. doi:10.18632/aging.101168
- Simpkin, A. J., Hemani, G., Suderman, M., Gaunt, T. R., Lyttleton, O., Mcardle, W. L., et al. (2016). Prenatal and Early Life Influences on Epigenetic Age in Children: a Study of Mother-Offspring Pairs from Two Cohort Studies. *Hum. Mol. Genet.* 25 (1), 191–201. doi:10.1093/hmg/ddv456
- Team, R. C. (2017). *R: A Language and Environment for Statistical Computing*. Vienna, Austria: R Foundation for Statistical Computing. Available at: <https://www.R-project.org>.
- Vetter, V. M., Kalies, C. H., Sommerer, Y., Spira, D., Drewelies, J., Bertram, L., et al. (2021). Relationship between Five Epigenetic Clocks, Telomere Length and Functional Capacity Assessed in Older Adults: Cross-Sectional and Longitudinal Analyses. medRxiv, 2021. doi:10.3390/ijms21264547
- Vetter, V. M., Meyer, A., Karbasiyan, M., Steinhagen-Thiessen, E., Hopfenmüller, W., and Demuth, I. (2018). Epigenetic Clock and Relative Telomere Length Represent Largely Different Aspects of Aging in the Berlin Aging Study II (BASE-II). *J. Gerontol. A Biol. Sci. Med. Sci.* 74, 27. doi:10.1093/gerona/gly184
- Vetter, V. M., Spira, D., Banszerus, V. L., and Demuth, I. (2020). Epigenetic Clock and Leukocyte Telomere Length Are Associated with Vitamin D Status, but Not with Functional Assessments and Frailty in the Berlin Aging Study II. *The Journals Gerontol. Ser. A* 75, 2056. doi:10.1093/gerona/glaa101
- Vidal-Bralo, L., Lopez-Golan, Y., and Gonzalez, A. (2017). Corrigendum: Simplified Assay for Epigenetic Age Estimation in Whole Blood of Adults. *Front. Genet.* 8, 51. doi:10.3389/fgene.2017.00051
- Vidal-Bralo, L., Lopez-Golan, Y., and Gonzalez, A. (2016). Simplified Assay for Epigenetic Age Estimation in Whole Blood of Adults. *Front. Genet.* 7, 126. doi:10.3389/fgene.2016.00126
- Wickham, H. (2009). *Elegant Graphics for Data Analysis (Ggplot2)*. New York, NY: Springer-Verlag.
- Zhang, Y., Saum, K.-U., Schöttker, B., Holleczeck, B., and Brenner, H. (2018). Methylopic Survival Predictors, Frailty, and Mortality. *Aging* 10 (3), 339–357. doi:10.18632/aging.101392

Conflict of Interest: 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.

Publisher’s Note: All claims expressed in this article are solely those of the authors and do not necessarily represent those of their affiliated organizations, or those of the publisher, the editors and the reviewers. Any product that may be evaluated in this article, or claim that may be made by its manufacturer, is not guaranteed or endorsed by the publisher.

Copyright © 2022 Vetter, Kalies, Sommerer, Bertram and Demuth. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.