

A decade of epigenetic change in aging twins: genetic and environmental contributions to longitudinal DNA methylation

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Summary/Abstract

Background. Epigenetic changes may result from the interplay of environmental exposures and genetic influences and contribute to differences in age-related disease, disability and mortality risk. However, the etiologies contributing to stability and change in DNA methylation have rarely been examined longitudinally. **Methods.** We considered DNA methylation in whole blood leukocyte DNA across a 10-year span in two samples of same-sex aging twins: (a) Swedish Adoption Twin Study of Aging (SATSA; N = 53 pairs, 53% female; 62.9 and 72.5 years, SD=7.2 years); (b) Longitudinal Study of Aging Danish Twins (LSADT; N = 43 pairs, 72% female, 76.2 and 86.1 years, SD=1.8 years). Joint biometrical analyses were conducted on 358,836 methylation probes in common. Bivariate twin models were fitted, adjusting for age, sex and country. **Results.** Overall, results suggest genetic contributions to DNA methylation across 358,836 sites tended to be small and lessen across 10 years (broad heritability $M=23.8\%$ and 18.0%) but contributed to stability across time while person-specific environments explained emergent influences across the decade. Aging-specific sites identified from prior EWAS and methylation age clocks were more heritable than background sites. The 5,037 sites that showed the greatest heritable/familial influences ($p<1E-07$) were enriched for immune and inflammation pathways while 2,020 low stability sites showed enrichment in stress-related pathways. **Conclusions.** Across time, stability in methylation is primarily due to genetic contributions, while novel experiences and exposures contribute to methylation differences. Elevated genetic contributions at age-related methylation sites suggest that adaptations to aging and senescence may be differentially impacted by genetic background.

Keywords: DNA methylation, heritability, aging, longitudinal

Key messages

- Individual differences in late-life methylation are due partly to genetic influences which contribute to stability in methylation patterns across late life.
- Environmental experiences unique to individuals contribute to new influences on methylation patterns across late life.
- Age-related CpG sites show greater heritable influences on methylation consistent with genetic regulation of biological aging rates.

Introduction

The functional profiles of genes are not static and vary across time, and indeed across the lifespan, in part as a result of different environmental exposures and contexts (M. J. Jones, Goodman, & Kobor, 2015; Lappe & Landecker, 2015; McClearn, 2006; van Dongen et al., 2016). Measurable gene-environment dynamics for behavioral traits are possible due to advances in biotechniques for global epigenetic profiling at, e.g. specific methylation sites in the human genome. Epigenetic changes may be critical to the development of complex diseases, accelerated aging, or steeper declines in cognitive and physical functioning with age (Lappe & Landecker, 2015). Understanding epigenetic changes over time in the elderly may identify pathways of decline or plasticity (e.g., maintenance or even boosts in functioning) during the aging process and help with elucidating the biology of aging and survival.

Epigenetic modifications resulting in altered gene expression may occur due to a number of processes, including direct methylation of DNA (P. A. Jones & Takai, 2001). DNA methylation results from *nongenetic* (i.e., epigenetic) processes that may arise due to prenatal or early life exposures or at later points in development (Gottesman & Hanson, 2005; Kanherkar, Bhatia-Dey, & Csoka, 2014). DNA methylation is characteristically produced by the addition of a methyl group to the DNA molecule cytosine within cytosine-guanine dinucleotides (CpGs), estimated at 28 million sites across the human genome (Lovkvist, Dodd, Sneppen, & Haerter, 2016). Dense regions of CpGs referred to as ‘islands’ and represent about 5% of CpGs occurring in the genome (about 20,000 total) and often reside in promotor regions (Vinson & Chatterjee, 2012); in addition, surrounding ‘shores’ and ‘shelves’ to these islands are of interest and may be differentially methylated compared to islands (M. J. Jones et al., 2015). The addition of methylation tags to CpG sites is associated with altered gene expression, typically by interfering with or silencing gene

transcription although upregulation of gene expression has been documented (C. Wang et al., 2019), and may differentially occur in cells across multiple tissue types including brain, muscle and leukocytes (Fernandez et al., 2012). Methylation tags can be removed as a consequence of exposures as well, leading to dynamics in expression across time (Kanherkar et al., 2014).

Although epigenetic variation is largely attributed to environmental factors, there is evidence for genetic contributions to variation in methylation across the epigenome (Hannon et al., 2018; van Dongen et al., 2016), with an average of 16.5 – 18.0% across sites in the Illumina 450k chip array from whole blood and common environmental influence of 3 to 16.7%, with stronger evidence of common environment in young adulthood (Hannon et al., 2018).

Epigenetic changes may accelerate over time, whereby changes in gene expression due to exposures become more abundant and salient to phenotypic changes, hence potentiating the development of health and aging conditions earlier in life. Indeed, methylation is correlated with age (Ciccarone, Tagliatesta, Caiafa, & Zampieri, 2018; van Dongen et al., 2016), is used to define biological clocks that may more closely track biological aging (Field et al., 2018), and is associated with mortality (Y. Zhang et al., 2017) and a number of physical and neuropsychiatric health traits (Kanherkar et al., 2014; Lappe & Landecker, 2015). Longitudinal studies of twins represent a valuable approach to evaluate genetic and environmental contributions to stability and change in methylation across the methylome (Tan, Christiansen, von Bornemann Hjelmberg, & Christensen, 2015). Investigations of etiological contributions have relied primarily on cross-sectional data (Hannon et al., 2018; van Dongen et al., 2016) and have addressed age-related differences (van Dongen et al., 2016) but not change. We evaluate individual differences in DNA methylation at individual CpG sites across the methylome across 10 years in two Scandinavian samples of same-sex aging twins, estimating the genetic and environmental contributions to stability as well as to

novel influences that emerge. Moreover, we examine whether surrounding ‘shores’ and ‘shelves’ are differentially heritable compared to islands, and, whether sites identified as associated with rate of aging in epigenome-wide association study (EWAS) or individual CpG clock sites are differentially heritable.

Methods

Sample. We considered DNA methylation across a 10-year span in 96 pairs of same-sex aging twins (40 monozygotic, MZ pairs; 56 dizygotic, DZ pairs). Across two samples, the average age at time 1 was 68.89 years ($SD=8.58$) and at time 2 was 78.59 years ($SD=8.70$). Specifically, the Swedish Adoption Twin Study of Aging (SATSA) included 53 pairs (22 MZ, and 31 DZ pairs; 53% female), selected with measurements about 10 years apart (range = 8.00 to 11.82 years) at ages 62.9 and 72.5 years at time 1 and time 2 respectively ($SD=7.2$). In 4 of 53 SATSA pairs, one twin partner had methylation data from one time point instead of both time points, but all data were included for these pairs. The Longitudinal Study of Aging Danish Twins (LSADT) included 43 pairs (18 MZ, and 25 DZ pairs; 72% female) at ages 76.2 and 86.1 years at time 1 and time 2 ($SD=1.8$).

Materials. Methylation measurements from the Illumina HumanMethylation450 array (Illumina, San Diego, CA, USA) were preprocessed and normalized with adjustments for cell counts and batch effects. Processing of the SATSA sample probes has been described previously (Jylhävä et al., 2019; Y. Wang et al., 2018) and in brief included: (a) preprocessing with the R package *RnBeads* (Assenov et al., 2014) where filtering of samples and probes proceeded with a greedy-cut algorithm maximizing false positive rate versus sensitivity at a detection p-value of 0.05; (b) removal of sites that overlap with a known SNP site or reside on sex chromosomes; (c) normalization of data using *dasen* (Pidsley et al., 2013); (d) applying a Sammon mapping

method (Sammon, 1969) to remove technical variance; (e) adjustment for cell counts (M. J. Jones, Islam, Edgar, & Kobor, 2017); (f) correction for batch effects using the ComBat approach in the *sva* package (Leek et al., 2012).

Processing of the LSADT data has been described previously (Svane et al., 2018) and in brief included: (a) preprocessing with the R-package MethylAid (van Iterson et al., 2014) where samples below quality requirements were excluded and probes with detection p-value >0.01 , no signal, or bead count <3 were filtered out; (b) removal of probes with $>5\%$ missing values, removal of sites that reside on sex chromosomes or cross-reactive probes; (c) normalization and batch-correction using functional normalization (Fortin et al., 2014) with four principal components.

Although Beta-values are preferred for interpretation of methylation, Beta-value units were translated into M-values via a log₂ ratio for improved distributional properties for the analysis of individual differences (Du et al., 2010). After performing the preprocessing steps, 390,894 probes remained for SATSA and 452,920 CpG sites remained for LSADT.

Altogether 368,391 sites were in common across the Swedish and Danish samples. After the described QC pre-processing in SATSA, 49 of 53 pairs had methylation data available for both members of each pair at both time points, while in 4 pairs one cotwin member had data at both time points while their twin partner had data at one timepoint but not both. After pre-processing, LSADT sample had methylation data represented for both cotwins at both timepoints among the 43 pairs.

Filtering of sites post-analysis. We conducted additional filtering of probes where model-fitting results evidenced means or variances outside of expected values. Specifically, we filtered based on the typical range of M-values (c.f., Du et al., 2010), with expected mean values

falling outside the range -6.25 to 6.25 for 1812 sites under either the ACE or ADE models at either timepoint. Likewise, we filtered based on expected standard deviations exceeding 1.5 under either the ACE or ADE models (Du et al., 2010) resulting in 9554 sites out of range under either the ACE or ADE models at either timepoint. The effective reduction in sites was from 368,391 to 358,836 after dropping 9555 unique sites from the analysis set.

Analysis. Bivariate twin models of M-values were fitted to all available data across the pairs using full-information maximum likelihood (FIML), adjusting for centered age (centered at the average age across time = age - 74 years), sex (0=males, 1=females), and country (0=Sweden, 1=Denmark). Bivariate ACE and ADE Cholesky models evaluated the degree to which additive genetic (A), dominance or non-additive genetic (D), common environmental (C), and non-shared factors (E), encompassing non-shared environmental influences, measurement error, and stochastic factors, contributed to variation and covariation in M-values within and across time (see Figure 1). The resolution of the genetic and environmental effects are done by comparing the relative similarity of monozygotic (MZ) twins who share 100% of their genes in common, including all additive effects and dominance deviations, versus dizygotic (DZ) twins who share on average 50% of segregating genes in common leading to expectations of 50% for additive effects and 25% for dominance deviations. Both twin types are presumed to have the same contribution of common environmental effects that contribute to similarity. We fitted ADE and ACE models as dominance (D) and common environment (C) could not be simultaneously estimated (see Figure 1).

Fit comparison between the ACE and ADE models was done via Akaike Information Criterion (AIC; Akaike, 1974). If the fit of the ADE model was as good or better than the ACE model it was retained as 'best' fitting, and otherwise the ACE model was retained as best. We

evaluated submodels including AE, CE and E models. Differences in nested model deviance statistics $[-2\ln(L)]$ are distributed as chi-square (χ^2) with the difference in the number of parameters between the full and constrained models as the degrees-of-freedom (df). LSADT samples tended to show lower variability in methylation at any given probe compared to SATSA, hence we allowed for scalar differences at each time-point (k_1, k_2) in standard deviations between the two samples (see Figure 1). Thus, the relative contributions of A, C or D, and E were equated across LSADT and SATSA, but the scalar allowed for the variance components to differ by a constant at each assessment. Scalar differences in standard deviations were on average $k_1 = .90$ (SD=.93) and $k_2 = .88$ (SD=.89).

In comparing relative heritabilities across sites by location, as well as aging/clock CpGs sets to remaining CpGs, we fitted random effects regression models to age 69 and 79 biometrical estimates using *lme* (version 1.1-21; Bates, Mächler, Bolker, & Walker, 2015). We allowed for random effects between and within sites, reflecting consistency of effects by CpG sites across age and nonsystematic variation across time.

To compare time1-time2 correlations from the biometrical estimates we rescaled the a_{12} , d_{12} or c_{12} and e_{12} paths into correlations (r_A, r_D or r_C , and r_E) and performed Fisher Z-transformations before submitting each to a skew-normal regression analysis using the *sn* package (Azzalini, 2020). Regression analyses compared low stability sites to remaining CpGs, after which regression weights were inverse-transformed into correlation units for interpretation.

Enrichment analyses were conducted GREAT 4.0.4 (McLean et al., 2010). Selected sites were mapped to the Human GRCh37 build and default settings were used for association rules (i.e., basal+extension: 5000 bp upstream, 1000 bp downstream, 1000000 bp max extension, curated regulatory domains included). We present results of both biomial and hypergeometric

tests where the False Discovery Rate (FDR) achieved $p < .05$ and where fold enrichment (FE) tests exceeded 2.0. We followed up the enrichment analyses using the mQTL Database (Gaunt et al., 2016) to annotate associations with methylation quantitative trait loci, noting the number of *cis* or *trans* variants.

Results

Broad-sense heritable contributions (A+D, N=358,836) were on average small at age 69 years ($M = 0.238 * 100 = 23.8\%$, time 1) and decreased across 10 years ($M = 0.180 * 100 = 18.0\%$, time 2) (see Table 1, Variance Components). The decrease in heritability is due to absolute increase in non-shared influences (E) compared to genetic (or common environmental) influences (see Table 1, Absolute Variances). Across time, heritabilities showed divergence by location [ADE best (A+D): $\chi^2(5) = 618.3$, $p = 2.25E-131$; ACE best (A): $\chi^2(5) = 339.5$, $p = 3.19E-71$] (see Table S1, Figure 2). In ADE best results, islands and shelves showed lower broad (A+D) heritabilities than open seas by -.01 or -1% ($p \leq 1.55E-07$) whereas shores were higher by .01 or 1% than open seas ($p \leq 3.79E-15$). In ACE best results, comparably lower heritabilities (A) were observed for islands versus open seas ($p = 3.54E-58$).

At $p < 1E-07$, 5037 or 1.4% of sites showed broad genetic (A, D) or familial effects (A, C) within or across time ($df = 6$), and 35,762 or 10.0% of sites met $p < 1E-02$. Among the 358,836 sites, 52% of sites showed the better-fitting model was ADE (N=187,535) while 48% showed ACE as better-fitting (N=171,301) (see Table 1, Figure 3). A total of 16.4% or 58,676 sites (19308 ADE best, 16454 ACE best) achieved nominal significance comparing the ADE or ACE versus an E model ($p < .05$, 6 df), and 25.5% or 91,380 sites achieved nominal significance of an AE model over an E model ($p < .05$, $df = 3$). Given that power is low for C even in large samples,

as well as to distinguish D from A, we present full model estimates (Visscher, Gordon, & Neale, 2008).

In terms of contributions to stability and change in methylation due to genetic or environmental influences, across 358,836 sites, 58.5% showed cross-time associations at $p < .05$ ($df = 3$, where $a_{12}=[d_{12} \text{ or } c_{12}]=e_{12}=0$) indicative of stability over time due either to genetic and/or environmental mechanisms. As shown in Figure 3, the cross-time stability was largely due to genetic effects in both the ADE best and ACE best models which was most often perfect in correlation.

We observed a correlation between estimated total variability and the extent to which heritable effects were observed: $r_{SD,A+D} = .33$ and $.27$ (187535 sites, ADE best) at time 1 and 2, and $r_{SD,A} = .25$ and $.28$ (171301 sites, ACE best) at time 1 and 2, respectively. Sites in which nonshared factors, E, explained all of the variability of M-values (>99%) at both time points included 8,268 total sites (5,520 ADE best, 2,748 ACE best). In all these cases, we observed that either the MZ twin correlations of M-values were negative (< 0), or DZ correlations were sufficiently negative ($< -.05$), or the difference between MZ and DZ correlations at each time point were sufficiently negative ($< -.1$).

Age-related sites. We evaluated the best-fitting ADE and ACE results of two published CpG sets that were identified in EWAS as related to Age that overlap with the samples used in the presented analysis : (I) 1217 sites from Wang et al. (Y. Wang et al., 2018); (II) 1934 sites from Tan et al. (Tan et al., 2016). Multilevel regression models compared heritabilities by location from the ADE best or ACE best model, fitted to both age 69 and 79 estimates in set I [ADE best (A+D): $\chi^2(5) = 43.7, p = 2.66E-08$; ACE best (A): $\chi^2(5) = 27.9, p = 3.81E-05$], with Islands under ADE or ACE models showing lower heritabilities by .09-.10 or up to a 10%

difference than open seas (both $p \leq 5.13E-07$; see Table S1). In set II, age 69 and 79 heritability estimates also showed divergence by location [ADE best (A+D): $\chi^2(5) = 16.8$, $p = 4.90E-03$; ACE best (A): $\chi^2(5) = 19.4$, $p = 1.62E-03$], with Shores showing higher heritabilities by about .04 or 4% than open seas under ADE or ACE models (all $p \leq 2.56E-02$; see Table S1).

Multilevel regression models were fitted to age 69 and 79 estimates to compare the Aging sets' CpG sites to the remaining background CpGs. Stronger heritable influences were apparent for Aging Set I (1217 sites) compared to remaining CpGs with .16 higher broad heritability (.39 vs .24, ADE best; $p = 5.43E-162$) and .12 higher narrow heritability (.18 vs .07, ACE best; $p = 3.87E-146$) and .05 higher common environmentality (.15 vs .10; $p = 1.88E-40$) (see Table S2, Variance Components). Patterns in the absolute variances suggested the greater heritability was due primarily to lower nonshared factors ($p \leq 1.35E-07$) and for ACE models coupled with higher additive genetic and common environmental influences ($p \leq 2.03E-04$; see Table S2, Absolute Variances). Significantly higher heritabilities and common environmentality were also observed for Aging Set II (1934 sites) where the increased heritable and common environmental influences (all $p \leq 1.55E-31$) were driven mainly by amplified genetic and common environmental influences ($p \leq 1.01E-11$) and otherwise comparable non-shared factors between the Aging II set and remaining background CpGs (see Table S2). Thus, the age-related sites showed a significantly higher proportion of variance attributed to heritable and shared environmental influences due to lower nonshared factors in Aging set I and due to higher genetic and common environmental influences in Aging set II.

Methylation clock sites. Available CpG sites from four epigenetic-clocks were evaluated in similar fashion using multilevel regression models fitted to age 69 and 79 estimates: (a) 59 of 71 sites Hannum clock (Hannum et al., 2013), (b) 312 of 353 sites Horvath clock (Horvath,

2013), (c) 443 of 513 sites Levine clock (Levine et al., 2018), and 455 of 514 sites from the Zhang clock (Q. Zhang et al., 2019). Significantly higher heritabilities (A+D, A) and common environmentality (C) were observed for the 1190 unique Clocks sites compared to all remaining CpGs (.02 to .05 higher, $p \leq 8.68E-10$, see Table S2 Variance Components). Comparisons of absolute variances suggested amplified genetic and common environmental influences ($p \leq 3.94E-05$) as well as non-shared factors ($p \leq 3.90E-08$) between the Clock sites and remaining background CpGs (see Table S2, Absolute Variances). Thus, the clock sites showed greater overall variability across sources of variance suggesting greater individual differences in these sites, with a significantly higher portion of variance attributed to heritable and shared environmental influences.

Among the 1190 unique CpG sites compared to one another, the Zhang clock sites tended to show stronger broad (A+D) genetic (.07 to .08 higher, $p \leq 4.00E-07$), and shared environmental (C) contributions (.06 higher, $p \leq 2.19E-08$) than Horvath or Levine clock sites, while Hannum sites were comparable to Zhang sites (within -.018 to .018, $p \geq 4.71E-01$) (see Table S3). The ratio of Intercept variance to total variance (ρ) in heritability estimates was .559 for ADE best models and .680 for ACE best models suggesting 56% and 68% of the variation in heritability, respectively, was CpG site specific across time and less than half of the variation was unique to CpG site and time, consistent with analyses by location (see Table S1) . Likewise, absolute variances showed strong between-site variations (65%-88%, see Table S3).

Low Stability Sites. We identified 2020 CpGs with low stability but meaningful genetic or common environmental contributions at one or both timepoints, i.e., $p > .01$ ($df = 3$, where $a_{12}=[d_{12} \text{ or } c_{12}]=e_{12}=0$) and where e_1 or e_2 accounted for less than 50% of the total variation (1638 ADE best, 382 ACE best). Based on skew-normal analyses, low stability CpGs had lower

correlations among non-shared factors across time than background CpGs (ADE best: $r_{E,background} = .24$ vs $r_{E,low} = .10$, $p = 2.80E-154$; ACE best: $r_{E,background} = .17$ vs $r_{E,low} = .07$, $p \leq 3.09E-27$).

The correlations of genetic (r_A , r_D) and common environmental influences (r_C) across time were comparable (within .02 units) between background and low stability CpGs, albeit significant ($p \leq 1.02E-03$), and otherwise very strong based on skew-normal analyses ($r_{background} = .97 - .99$ vs $r_{low} = .95 - .99$). Variability in these low stability CpGs increased across time with a ratio of SD_2/SD_1 of 1.08 to 1.09 ($SD_{ratio} = .13$) for ACE and ADE best models, respectively. Moreover, heritabilities decreased across time while non-shared components tended to increase (see Figure S1). Compared to background CpGs, low stability CpGs tended to show higher A+D or A and C components (all $p \leq 2.03E-14$) but generally lower overall absolute variances for A+D and E variances ($p \leq 15.51E-09$) in ADE models. Higher absolute variance for A but lower variance for E was observed in ACE models ($p \leq 4.94E-03$) (see Table S2). Altogether, results suggest lower overall phenotypic variance in methylation among the low stability versus background CpGs across time (c.f., Table S2). However, within the set of lower stability CpGs, variance in methylation increased at time 2 mainly due to novel non-shared factors (c.f., Figure S1).

Enrichment analysis: High heritability/familiarity. A set of 5037 CpGs achieving $p < 1E-07$ when evaluating tests of heritability (AD vs E; N = 2049) or familiarity (AC vs E; N=2988) across time were submitted to GREAT 4.0.4 to identify functions of cis-regulatory regions (McLean et al., 2010). This represents 1.4% of the 358,836 sites tested. Specifically, we report the binomial and hypergeometric tests over genomic regions covered by the **5037** CpGs, reporting those that achieved region-based fold enrichment (FE) > 2 and both binomial and hypergeometric FDR Q-Values $< .05$ (see Table 2; for full ontology results see Table S4). The sites that showed the greatest heritabilities showed enrichment in immune and inflammation

pathways as well as neurotransmitter activity pathways. For example, the MHC protein complex pathway in the GO Cellular ontology list includes HLA region genes that code for HLA class II histocompatibility antigens in humans (c.f., GO:0042611, Table S4). Moreover, the interferon-gamma-mediated signaling pathway in the GO Biological ontology list include numerous genes associated with altered cytokine signaling and genes in the HLA region (c.f., GO:0060333, Table S4).

The set of 5037 CpGs were then submitted to the mQTL Database (Gaunt et al., 2016). The search resulted in 1435 unique CpG matches to 155,177 SNP variants from the Middle Age timepoint (see Table S6). Of the 1435 CpG matches, 1256 were associated with *cis*-mQTLs and 304 were associated with *trans*-mQTLs suggesting an abundance of associations with *cis*-mQTLs. The maximum number of mQTLs associated with any given CpG was for cg03202060 with 5230 *cis*-mQTLs variants plus 575 *trans*-mQTLs. The *cis*-mQTLs for cg03202060 reside in the HLA region on chromosome 6 (e.g., <https://www.genecards.org/cgi-bin/carddisp.pl?gene=HLA-DQB1&keywords=HLA-DQB1>), and the *trans*-mQTLs traverse genes such as *DDAH2* related to metabolism of nitric oxide (<https://www.genecards.org/cgi-bin/carddisp.pl?gene=DDAH2>) and *BAG6* (<https://www.genecards.org/cgi-bin/carddisp.pl?gene=BAG6&keywords=BAG6>) residing within the major histocompatibility class III region (MHCIII) and involved in the control of apoptosis. A scatterplot of cg03202060 M-values of twin 1 by twin 2 across time is shown in Figure S2a,b showing greater similarity for MZ than DZ pairs.

As polycomb repression may relate to age-related changes in DNA methylation, we filtered our set of 5037 CpGs to reflect genes annotated on the 27k array and evaluated whether our set mapped to 1861 PolyComb Group Target genes (PCGTs) identified using the Illumina

27k chip probes (Zhuang et al., 2012). We observed 493 CpGs within a set of 293 PCGTs overlapped, or a 15.7% overlap of PCGTs (see Table S6). A hypergeometric test of the 293 overlapping PCGTs was significant at $p = 1.004E-11$ suggesting overrepresentation, when considering the number of unique PCGTs in Zhuang et al. (2012), and the number of genes represented in the Illumina 27k chip.

Enrichment analysis: Low stability sites. The 2020 low stability CpGs were submitted to GREAT 4.0.4, showing enrichment for stress-related DNA and RNA transcription pathways (see Tables S7-S8). Hence, these sites may lie in genes/gene pathways that are sensitive to exogenous exposures to stress leading to increasing divergence in methylation profiles across time. The GO Biological RNA and DNA pathways noted relate to heat shock and response to hypoxia in a number of plant and animal species, including humans (c.f., annotations GO:0043620, GO:0061418; Table S8).

The low stability CpGs were submitted in kind to the mQTL Database (Gaunt et al., 2016) producing 397 unique CpG matches to 7103 mQTLs at the Midlife timepoint. Of the 397 CpG matches, 58 annotations were to *cis*-mQTLs and 347 were to *trans*-mQTLs (see Table S9), suggesting an abundance of associations with *trans*-mQTLs. The maximum number of mQTLs linked with any given CpG was for cg07677296 matched with 576 *cis*-mQTLs. The *cis*-mQTLs variants associated with cg07677296 traverse *FAHD1* and *NUBP2* on chromosome 16 and have been implicated in aging pathways related to insulin-like growth factor (Teumer et al., 2016). A scatterplot of cg07677296 M-values of twin 1 by twin 2 across time shows comparable similarity for MZ and DZ pairs (see Figure S2c,d).

Discussion

Overall, results suggest genetic contributions to DNA methylation tended to be small, vary by location, and decrease across a decade; however, genetic influence mainly contributed to the stability of methylation. Unique person-specific influences not shared by co-twins were emergent across 10 years suggesting that non-shared factors become more salient to DNA methylation in late life. The extent of variation in methylation at any given CpG site was positively correlated with observing stronger heritable effects. Moreover, 58% of sites showed stability across time due to strongly correlated genetic influences and modestly correlated nonshared factors, suggesting continuity of influences across 10 years for more than half the CpG sites. The sites that showed the greatest heritabilities showed enrichment in immune and inflammation pathways and neurotransmitter transporter activity pathways. Low stability sites meanwhile showed increased expression variability across time due to novel nonshared factors, with enrichment in stress-related pathways, suggesting that these sites are responsive to “new” environmental cues even in old age.

Prior studies report average heritabilities of 16.5 – 18.0% across adulthood (17-79 years) (Hannon et al., 2018; van Dongen et al., 2016), and common environmental influences of 3.0 – 16.7%, with stronger evidence of common environment in young adulthood (Hannon et al., 2018). Our results of weakening heritable influences across age is consistent with the Dutch cross-sectional study reporting average heritabilities of 21% and 18% at ages 25 and 50 assuming an AE model, whereas our estimates of broad heritability under an ADE model are 24% and 18% many decades later at age 69 and 79 years, respectively. Where non-additive genetic effects fit best, the average broad heritability was 24% across age. For sites where

including common environment fit best (ACE), lower average heritabilities were observed at 7% and common environment contributed 10% to variation in methylation across age.

CpG sites related to age show a greater impact of heritable influences consistent with genetic regulation of the rate of biological aging. Sites associated with age and longevity generally show higher heritabilities than the total background sites and varied in magnitude of heritabilities by location, where ‘islands’, which often reside in promotor regions (Vinson & Chatterjee, 2012), typically showed lower heritability than those sites residing in surrounding ‘shores’ and ‘shelves’, which have been shown to be differentially methylated compared to islands (M. J. Jones et al., 2015).

Moreover, the set of methylation clocks sites are likewise more heritable than background CpG sites, with Zhang sites more heritable than Horvath and Levine sites, and Hannum sites comparable to the Zhang sites. We have recently reported heritability estimates of methylation clock ages of 52% for the Horvath clock and 36% for the Levine clock (Jylhava et al., 2019), where, consistent with our current site-specific effects, stability across time was mediated primarily by genetic factors, whereas the person-specific environmental factors contributed to differences across time. The 353 Horvath clock sites were selected as best predictors of chronological age using multiple tissues (Horvath, 2013) similar to the 513 Levine clock sites that were selected based on prediction of chronological age and nine biomarkers of phenotypic aging with models trained on multiple tissues (Levine et al., 2018). The 71 Hannum clock sites best predicted age (adjusted for sex, BMI) based on methylation observed in whole blood while the 514 sites from the Zhang prediction model relied on methylation observed in blood and saliva samples (Q. Zhang et al., 2019). The current findings of moderately higher heritabilities in

the Zhang and Hannum sites versus the other clock sites may be in part due to our use of blood tissue.

Enrichment analyses of the 1.4% of sites meeting $p < 1E-07$ suggest immune and inflammation pathways and neurotransmitter transporter activity pathways may feature in sites with strong heritable or familial components. Moreover, the analysis of mQTL associations suggest that a number these high heritability CpGs are associated largely with *cis*-mQTLs, including those in the HLA region. Previous studies have identified methylation changes associated with altered immune functioning, including age-related hypermethylation and reduced expression in CD8+ cells for genes involved in T cell mediated immune response and differentiation (Tserel et al., 2015). Indeed, five CpGs in our set identified as associated with *cis*-mQTLs at midlife, lie within the *BCL11* gene (cg26396443) or *RUNX3* gene (cg05162523, cg13566436, cg20674490, cg22509179) involved in T cell differentiation (Tserel et al., 2015). A related study of German and Danish individuals (including an overlapping sample of twins herein) evaluating RNA-sequencing expression patterns and longevity identified expression patterns in biological processes contributing to immune system and response pathways (Häsler et al., 2017), and observed high heritabilities (30-99%) among 20% of *cis*-eQTLs.

Immunosenescence describes an age-associated decline in elderly individuals' immune functioning, such as mounting less effective responses to vaccines, and lowered resistance to illnesses, with concomitant up-regulation of pro-inflammatory cytokines, among several other cellular and physiological changes in the immune system (Accardi & Caruso, 2018). It has been proposed that heritable factors may be partly associated with differential immune responses (Derhovanessian et al., 2010; Poland et al., 2014) and may predict influenza-related susceptibility and mortality (Poland et al., 2014), for example, and, broadly, successful aging and

longevity (Derhovanessian et al., 2010). Hence, differential adaptations to aging processes including immunosenescence reflect gene-environment dynamics with some individuals showing better adaptations than others due to genetic influences.

High heritability CpGs were also enriched for PCGTs – a group of genes that are epigenetically regulated by polycomb-group proteins and involved in developmental processes and cell-fate decisions (Lanzuolo & Orlando, 2012). Enrichment of hypermethylated of PCGT has also been implicated in cancer and aging and show consistent patterns across different cell types (Teschendorff et al., 2010). Our findings would thus support the role of heritable/familial factors in the epigenetic regulation of these fundamental cellular processes.

Enrichment analyses of low stability CpG sites suggest that stress-related DNA and RNA transcription pathways may be relevant for these environmentally responsive sites which showed increased novel environmental contributions to methylation. It is notable that unlike the high heritability set, the low stability set showed more associations with *trans*-mQTLs. That said, cg07677296 matched with 576 *cis*-mQTLs, with variants spanning *FAHDI* and *NUBP2*, both implicated in metabolic and aging pathways related to insulin-like growth factor (IGF) (Teumer et al., 2016). Specifically, *FAHDI* was identified as a *cis*-eQTL associated with a variant in *NUBP2* (rs1065656) that may contribute to circulating IGF-I and IGFBP-3 concentrations (Teumer et al., 2016). Moreover, IGF-I is implicated in oxidative stress pathways (Gubbi et al., 2018).

The current study establishes the extent to which the genetic and environmental influences contribute to site-specific methylation across a 10-year span in a longitudinal sample of Swedish and Danish twins. While stability of methylation was largely due to genetic influences, person-specific environmental influences were emergent across time and explained change. By and

large, the dynamics of methylation may be influenced by experiences and exposures, suggesting possible mediation of gene expression; however, the most heritable sites may participate in immune and inflammation pathways and neurotransmitter transporter activity pathways which suggest that adaptations to aging and senescence may be differentially impacted by genetic background.

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Author contributions

CAR drafted the manuscript. CAR and EM analyzed data. QT and JH contributed to scripting and QT, SH and JJ advised on enrichment analyses. NLP and SH contributed to the coordination of the study and acquisition of the SATSA methylation data. JJ contributed to preparation of SATSA data and interpretation of results. LC, QT and JH coordinated the LSADT data

acquisition. All authors participated in interpretation of the data, have read and commented on the manuscript and approved the final version.

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Table 1. Variance components and absolute variances at time 1 (69 years) and time 2 (79 years).

Variance Components		A1		D1/C1		E1		A2		D2/C2		E2	
N sites		<i>M</i>	<i>SD</i>	<i>M</i>	<i>SD</i>	<i>M</i>	<i>SD</i>	<i>M</i>	<i>SD</i>	<i>M</i>	<i>SD</i>	<i>M</i>	<i>SD</i>
ADE	358,836	0.111	0.142	0.127	0.160	0.762	0.175	0.091	0.125	0.089	0.130	0.820	0.158
ADE best	187,535	0.057	0.106	0.217	0.168	0.725	0.182	0.048	0.092	0.152	0.148	0.800	0.168
ACE	358,836	0.150	0.166	0.057	0.083	0.793	0.163	0.109	0.142	0.054	0.080	0.837	0.147
ACE best	171,301	0.076	0.122	0.106	0.093	0.817	0.150	0.055	0.103	0.098	0.091	0.846	0.137
Absolute Variances		A1		D1/C1		E1		A2		D2/C2		E2	
N sites		<i>M</i>	<i>SD</i>	<i>M</i>	<i>SD</i>	<i>M</i>	<i>SD</i>	<i>M</i>	<i>SD</i>	<i>M</i>	<i>SD</i>	<i>M</i>	<i>SD</i>
ADE	358,836	0.029	0.079	0.038	0.088	0.162	0.144	0.028	0.079	0.032	0.086	0.214	0.195
ADE best	187,535	0.019	0.063	0.067	0.112	0.174	0.151	0.018	0.063	0.057	0.111	0.235	0.206
ACE	358,836	0.047	0.109	0.012	0.034	0.170	0.152	0.042	0.109	0.014	0.038	0.220	0.200
ACE best	171,301	0.071	0.130	0.003	0.012	0.187	0.162	0.062	0.130	0.004	0.014	0.244	0.214

Note. A = Additive genetic, D = Nonadditive genetic (Dominance), C = Common environment, E = Non-shared factors.

Table 2. GREAT 4.0.4 annotations using binomial and hypergeometric tests over genomic regions covered by the 5037 CpGs showing significant heritability/familiality $p < 1E-07$.

Ontology	Binomial						Hypergeometric					
	Rank	Raw P-Value	FDR Q-Val	Fold Enrichment	Observed Region Hits	Region Set Coverage	Rank	FDR Q-Val	Fold Enrichment	Observed Gene Hits	Total Genes	Gene Set Coverage
GO Biological Process												
interferon-gamma-mediated signaling pathway	3	1.71E-17	7.48E-14	3.23	74	0.015	105	2.55E-02	1.82	30	64	0.006
osteoblast development	104	5.04E-07	6.37E-05	2.48	39	0.008	57	3.91E-03	2.98	13	17	0.003
GO Cellular Component												
MHC protein complex	1	8.53E-45	1.47E-41	17.69	51	0.010	1	7.81E-05	3.31	17	20	0.004
integral component of luminal side of endoplasmic reticulum membrane	4	9.52E-31	4.11E-28	10.42	46	0.009	13	2.61E-02	2.48	14	22	0.003
MHC class II protein complex	5	2.78E-27	9.58E-25	19.42	29	0.006	2	5.32E-05	3.63	14	15	0.003
GO Molecular Function												
neurotransmitter:sodium symporter activity	53	6.16E-06	4.90E-04	2.90	24	0.005	17	2.80E-02	2.66	13	19	0.003

Note. Shown ontology from GREAT 4.0.4. FDR=False Discovery Rate.

Figure 1. Bivariate Cholesky model.

Note. ACE and ADE models were separately fitted to M-values at two waves 10 years apart.

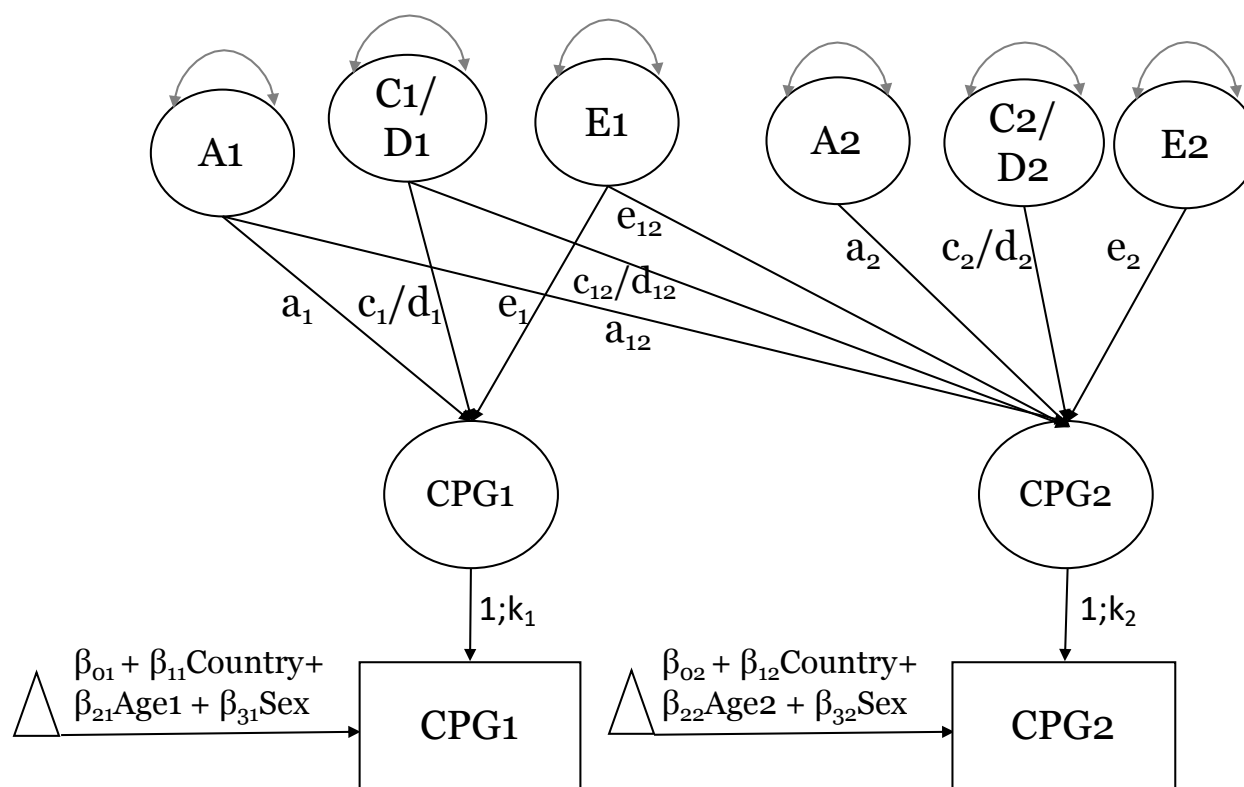


Figure 2. Broad-sense heritability by Location across 10 years (ADE results, 358,836 CpGs)
Note. Site differences shown below are significant across time: $\chi^2(5)=995.48, p=5.72E-213$

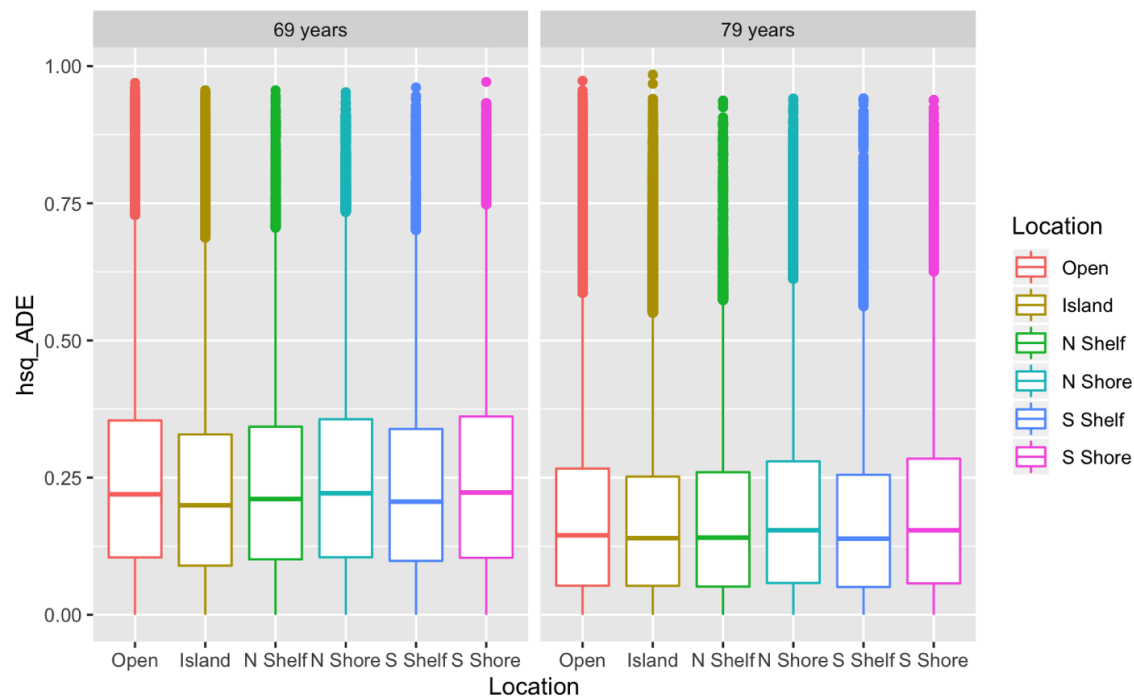


Figure 3. Best-fitting models: ADE (52%) or ACE (48%)

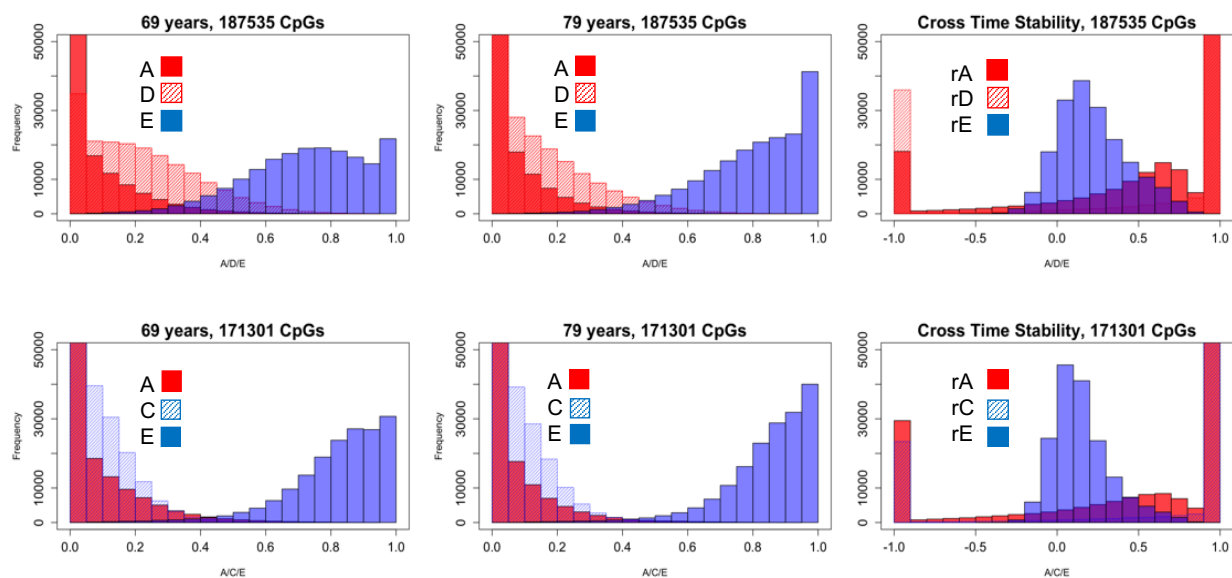


Figure 4. Age-related CpG Sets: Broad heritability by CpG Location

