

Longitudinal studies at birth and age 7 reveal strong effects of genetic variation on ancestry-associated DNA methylation patterns in blood cells from ethnically admixed children

Chris McKennan^{1*}, Katherine Naughton², Catherine Stanhope², Meyer Kattan³, George T. O'Connor⁴, Megan T. Sandel⁴, Cynthia M. Visness⁵, Robert A. Wood⁶, Leonard B. Bacharier⁷, Avraham Beigelman⁷, Stephanie Lovisky-Desir³, Alkis Togias⁸, James E. Gern⁹, Dan Nicolae^{1,2¶}, Carole Ober^{2¶} for the NIAID-sponsored Inner-City Asthma Consortium

¹Department of Statistics, University of Chicago, Chicago, IL

²Department of Human Genetics, University of Chicago, Chicago, IL

³Department of Pediatrics, Columbia University Medical Center, New York, NY

⁴Department of Medicine, Boston University School of Medicine, Boston, MA

⁵Rho Federal Systems Division, Chapel Hill, NC

⁶Department of Pediatrics, Johns Hopkins University Medical Center, Baltimore, MD

⁷Department of Pediatrics, Washington University School of Medicine and St Louis Children's Hospital, St. Louis, MO

⁸National Institute of Allergy and Infectious Disease, Bethesda, MD

⁹Departments of Pediatrics and Medicine, University of Wisconsin School of Medicine and Public Health, Madison WI

¶ Equal contributions

*Corresponding Author:

Email: cgm29@uchicago.edu

Abstract

The epigenetic architecture in humans is influenced by genetic factors, exposure histories and biological factors such as age, but little is known about their relative contribution or their longitudinal dynamics. Here, we studied DNA methylation levels at over 750,000 CpG sites in mononuclear blood cells collected at birth and age 7 from 196 children of primarily self-reported Black and Hispanic ethnicities to study age- and ancestry-related patterns in DNA methylation. We developed a novel Bayesian inference method for longitudinal data and showed that even though average methylation levels changed from birth to age 7, the vast majority of the ancestry-associated methylation patterns present at birth are also present at age 7. A large proportion of ancestry-associated CpGs (59%) had a nearby methylation quantitative trait locus (meQTL) and we show that at least 13% of the ancestry-associated methylation patterns were mediated through local genotype. These combined results indicate that ancestry-associated methylation patterns in blood are in large part genetically determined. Our results further suggest that DNA methylation patterns in blood cells are robust to many environmental exposures, at least during the first 7 years of life.

Introduction

Epigenetic patterning in human genomes reflects the contributions of genetic variation [1, 2] exposure histories [3-8], and biological factors, such as age [9-18], ethnicity [19-24] and disease status [25-28], among others. However, little work has been done to elucidate the relative contributions or longitudinal dynamics of each on epigenetic patterning.

To directly examine the relationship between age, ethnicity, genetic variation, early life exposures and allergic phenotypes and an epigenetic mark, we studied global DNA methylation patterns at over 750,000 CpG sites on the EPIC array in cord blood mononuclear cells (CBMCs) collected at birth and in peripheral blood mononuclear cells (PBMCs) collected at 7 years of age from 196 children participating in the Urban Environment and Childhood Asthma (URECA) birth cohort study[29, 30]. This cohort is part of the NIAID-funded Inner-City Asthma Consortium and is comprised of children primarily of Black and Hispanic self-reported ethnicity, with a mother and/or father with a history of at least one allergic disease living in poor urban areas (see Gern et al. [30] for details of enrollment criteria). Mothers of children in the URECA study were enrolled during pregnancy and children were followed from birth through at least 7 years of age.

The longitudinal design of the URECA study provided us with the resolution to partition genetic from non-genetic effects on ancestry-associated DNA methylation patterns, and yielded new insight into the factors affecting DNA methylation patterns at CpG sites in mononuclear (immune) cells during early life in ethnically admixed children. Using a novel statistical inference method that provides a general framework for analyzing longitudinal genetic and epigenetic data, we show that ancestry-dependent methylation patterns are conserved over the first 7 years of life and that these patterns are strongly influenced, and often mediated, by local

genotype. Further, chronological age, but not measured exposures during pre- or post-natal periods or disease status by age 7, was associated with methylation patterns in this sample of children. Considering the results of our study and those of a recently published comprehensive review on environmental epigenetics research [31], we suggest that methylation levels in blood are not as responsive to environmental exposures as previously suggested [20], at least during the first 7 years of life.

Results

Our study included 196 children participants in the Urban Environment and Childhood Asthma (URECA) cohort who had stored cord blood mononuclear cells (CBMCs) and peripheral blood mononuclear cells (PBMCs) collected at birth and age 7, respectively [29], and passed quality control (QC) checks as described in Methods. The URECA children were classified by parent- or guardian-reported race into one of the following categories: Black, $n = 147$; Hispanic, $n = 39$; White, $n = 1$; Mixed race $n = 7$, and Other, $n = 2$. A description of the study population is shown in Table 1 and in Supplementary Materials. Ancestry, assessed using ancestral PCs, revealed varying proportions of African and European ancestry along PC1. Because there is little separation along PC2 (Figure 1) and no genome-wide significant correlation between PC2 through PC10 and methylation levels at either age, we defined PC1 as inferred genetic ancestry (IGA). The reported races (RR) of the children are also shown in Figure 1. The means and ranges of gestational age stratified by reported race are shown in Table 1; the distribution of gestational age at birth is shown in Figure S1 in the supplement.

Table 1: Covariates for the 196 URECA children in our study, stratified by self-reported race.

	Black	Hispanic	White	Mixed	Other
Sample Size (N)	147	39	1	7	2
Males (%)	71 (48%)	25 (64%)	0 (0%)	4 (57%)	0 (0%)
Asthma diagnosis at age 7 (%)	38 (26%)	12 (31%)	0 (0%)	2 (29%)	0 (0%)
Gestational age at birth, in weeks (mean [range])	39.0 [34,42]	38.9 [35,41]	36.0	39.1 [37,40]	39.0 [38,40]
Sample Collection Site					
Baltimore (%)	64 (44%)	1 (3%)	1 (100%)	3 (43%)	2 (100%)
Boston (%)	17 (12%)	5 (13%)	0 (0%)	3 (29%)	0 (0%)
New York (%)	23 (16%)	32 (82%)	0 (0%)	1 (14%)	0 (0%)
St. Louis (%)	43 (29%)	1 (3%)	0 (0%)	1 (14%)	0 (0%)

Inferred genetic ancestry effects on DNA methylation patterns are conserved in magnitude and direction between birth and age 7

Previous cross-sectional studies have revealed associations between ancestry and DNA methylation at birth [19, 23] and later in life [20-22, 24, 25]. These correlations were generally attributed to the combined effects of genetic variation and environmental exposures [20-23]. However, because of the cross-sectional nature of these studies, it is not known if the association between ancestry and methylation patterns present at birth persist (or change) in childhood. Moreover, because ancestry is typically confounded with environmental exposures [41], it has been proposed that ancestry effects on methylation levels may reflect the effects of exposure

histories, which also may vary by race or ethnicity [20]. Alternatively, ancestry effects on DNA methylation patterns could also be due to genetic differences. In these cases, we would expect ancestry-associated methylation patterns to be conserved from birth to later childhood. Using the longitudinal data from the URECA cohort, we tested this hypothesis by addressing three questions. What is the correlation between ancestry and methylation levels at individual CpG sites at birth and age 7? Is the direction and magnitude of the correlation between ancestry and methylation levels conserved between birth and age 7? Are there any CpGs for which the correlation between methylation and ancestry at birth is significantly different from the correlation between methylation and ancestry at age 7?

Standard hypothesis testing can be used to answer the first question but is not appropriate for answering the second or third because failure to reject the null hypothesis that the effects are equal at birth and age 7 does not imply the null hypothesis is true. Additionally, because our studies were conducted in cord blood cells at birth and peripheral blood cells at age 7, effects at birth and age 7 may differ slightly due to differences in cell composition [42]. To circumvent these issues, we built a Bayesian model (see model (S3) in the Supplement) and let the data determine both the strength of the correlation between inferred genetic ancestry or reported race and methylation, and how similar the correlations were at birth and age 7. We then answered the first, second and third questions by defining and estimating the correct (*cor*), conserved (*con*) and discordant (*dis*) sign rates for each CpG $g = 1, \dots, 784,484$:

$cor_g^{(0)}$ = Posterior probability that the estimate for the direction of CpG g 's ancestry effect at birth was correct.

$con_g^{(0,7)}$ = Posterior probability that the directions of CpG g 's ancestry effects at birth and age 7 were the same AND the directions were estimated correctly.

$dis_g^{(0)}$ = Posterior probability that the ancestry effect for CpG g at birth was non-zero and was 0 or in the opposite direction at age 7 AND both directions were estimated correctly.

The correct and discordant sign rates at age 7 ($cor_g^{(7)}$, $dis_g^{(7)}$) were defined analogously. Because the correct sign rate at birth and age 7 is always at least as large as the conserved sign rate, we say that the birth and age 7 effects for CpG g overlap if its conserved sign rate ($con_g^{(0,7)}$) is above a designated threshold. Supplemental Figures S2 and S3 provide insight into how the correct and conserved sign rates compare with standard univariate P values. We refer the reader to "Joint modeling of methylation at birth and seven" in the Supplement for a detailed description of our model and estimation procedure.

After fitting the relevant parameters in the model to the data, we were able to estimate the fraction of CpGs with non-zero effects at both ages that fell into one of four possible bins: the two effects were completely unrelated ($\rho = 0$), moderately similar ($\rho = 1/3$), very similar ($\rho = 2/3$), or identical ($\rho = 1$). Note that if a non-trivial fraction of CpG sites had effects that were negatively related, they would be assigned to the first bin ($\rho = 0$). In fact, less than 1% of the CpGs with non-zero inferred genetic ancestry effects at both ages had unrelated or moderately similar IGA effects ($\rho = 0$ or $1/3$), whereas 34% fell in the very similar bin and 66% had identical inferred genetic ancestry effects at birth and age 7 (Supplemental Figure S4). This indicates that when inferred genetic ancestry effects on methylation are non-zero at both birth

and age 7, they tend to be very similar or exactly the same with respect to both direction and magnitude.

We then estimated the correct and conserved sign rates for all 784,484 probes, and identified 2,873 inferred genetic ancestry-associated CpGs (IGA-CpGs) in CBMCs ($cor_g^{(0)} \geq 0.95$), 3,834 in PBMCs at age 7 ($cor_g^{(7)} \geq 0.95$), and 2,659 whose effects were conserved in sign ($con_g^{(0,7)} \geq 0.95$). Methylation tended to increase with increasing African ancestry at 1,494 of the 2,659 conserved CpGs (56%), suggesting that individuals with more African ancestry tend to have more methylation ($P < 10^{-10}$). This is consistent with the study of Moen et al. [22], which used the Illumina 450K array to quantify the differences in methylation between European and African populations. Supplemental Figure S5 shows the IGA-CpG locations in the genome and Figure 2a illustrates the overlap between IGA-CpGs at birth and age 7. This strong overlap corroborates our above observations and answers the second question in the affirmative: if methylation is strongly correlated with inferred genetic ancestry at birth, the magnitude and direction of the correlation is conserved at age 7.

Inferred genetic ancestry is more correlated with methylation than is self-reported race

The observed correlations between ancestry and methylation levels may reflect differences in environmental exposures [20, 22], due to associations between race or ethnicity with socio-cultural, nutritional, and geographic exposures, among others [41]. In fact, Galanter et al. [20] showed in a cross-sectional study that self-reported ethnicity explained a substantial portion of the variability in whole blood DNA methylation patterns from Latino children of diverse ethnicities, even more so than inferred genetic ancestry. They concluded that ethnicity captures genetic, as well as the socio-cultural and environmental differences that influence methylation

levels. If this were the case in the URECA children, we should observe just as large, if not larger, an effect as we did for inferred genetic ancestry when we substitute reported race for inferred genetic ancestry in the analyses presented in the previous section. However, using reported race, we identified only 457 CpGs at birth and 709 CpGs at age 7 whose correct sign rate was at least 0.95, and 424 whose reported effects were conserved from birth to age 7 ($con_g^{(0,7)} \geq 0.95$), far fewer than the 3,991 CpGs whose methylation was significantly correlated with inferred genetic ancestry at birth or at age 7.

To explore this further, we examined the overlap between IGA-CpGs and reported race-associated CpGs (RR-CpGs) in CBMCs at birth and in PBMCs at 7 (Figures 2c-d). Because reported race is an estimate of inferred genetic ancestry, there is a still substantial overlap between IGA-CpGs and RR-CpGs. In fact, almost all of the RR-CpGs are among the IGA-CpGs, but the opposite is not true. This indicates that while IGA-CpGs include most RR-CpGs, reported race does not capture most of the variation in methylation attributable to inferred genetic ancestry in these children.

The observed correlations between DNA methylation and ancestry are primarily genetic

To further address the question of whether ancestry effects on methylation at either birth or age 7 were due to genetic variation or to environmental exposures, we used local genetic variation (within 5kb of a CpG site) and DNA methylation data at birth and age 7 in the 147 self-reported Black children in our study. Of the 519,622 CpGs within 5kb of a SNP, 65,068 and 70,898 had at least one meQTL in CBMCs at birth and in PBMCs at age 7, respectively, at an FDR of 5%. In addition, 59% of IGA-CpGs with at least one SNP in the ± 5 kb window had at least one meQTL

at birth or age 7 at an FDR of 5%, indicating IGA-CpGs were enriched for CpGs with meQTLs (Figure 3a-b).

To provide additional evidence that local genotype mediates the effect of inferred genetic ancestry on methylation, we used logistic regression to regress the genotype of each of the 269,622 SNPs in our study set onto inferred genetic ancestry. The goal was to determine the fraction of IGA-CpGs that were mediated through local genotype, i.e. IGA-CpGs with both edges a and c in Figure 3a. Our analysis first revealed that the genotypes at meQTLs whose target CpGs were IGA-CpGs in either CBMCs at birth or PBMCs at age 7 (IGA-meQTLs) were significantly more correlated with inferred genetic ancestry than the genotype of non-IGA-meQTLs (Figure 3c). Moreover, approximately 13% of the IGA-CpGs with at least one SNP in their ± 5 kb windows had an inferred genetic ancestry effect that was mediated through local genotype (i.e. had edges a and c, see Supplement for calculation details), which is likely an underestimate of the true number IGA-CpGs mediated through local genotype because our sample size was relatively small [43]. Nonetheless, this is striking compared to the 0.1% of non-IGA-CpGs whose corresponding SNP had edge c at a 20% FDR.

Lastly, we used DNA methylation data on 573 ethnically diverse U.S. Latino children ages 9 to 16 years old from the Galanter et al. study [20] to further explore the effect of ancestry on DNA methylation in whole blood. Children and teenagers in the Galanter study were classified as Mexican, Puerto Rican, mixed Latino, or other Latino. They reported 916 CpG sites whose methylation was significantly associated with reported race, 773 of which were also significantly associated with estimated percent European, Native American and African ancestry at a Bonferroni P value threshold of 1.6×10^{-7} . A total of 726 of their 916 ethnic-associated CpGs were also in the set of 784,484 probes CpG sites in our study. Our set of IGA-CpGs at

birth or age 7 contained a significant fraction of the 726 ethnic-associated CpGs from the cross-sectional study, but there was considerably less overlap with our RR-CpGs (Figure 4). If the correlation we observed between ancestry and methylation was largely due to responses to environmental exposures, as suggested in the Galanter study, then the overlap with reported race should have been at least as large as the overlap with inferred genetic ancestry. That was not the case, further suggesting that the inferred genetic ancestry effects on methylation in the URECA cohort are primarily genetic in origin.

Non-genetic factors influence the observed ancestry-methylation correlation more at age 7 than at birth

Although most of the variation in methylation levels at ancestry-associated CpGs can be attributed to genetic variation in the URECA children, a small proportion may be due to non-genetic (environmental) factors. To explore this possibility, we further hypothesized that non-genetic effects on methylation levels at ancestry-associated CpGs would be greater at age 7 than at birth, due to accumulated exposures over the first 7 years of life. We note that none of the direct or indirect measures of exposures that were available in this cohort were associated with methylation levels at either age, including maternal asthma, maternal infections during pregnancy, pet ownership, bedroom allergens, mother stress, anxiety and depression metrics, maternal cotinine levels during pregnancy, number of smokers in the household, number of siblings, number of previous live births, daycare attendance, number of colds at age 2 or 3, and allergic sensitization or asthma in the child (see Supplement for details). We did, however, identify 16,172 age-related CpGs (CpGs whose methylation changed from birth to age 7). Besides being strongly enriched for CpGs used to predict gestational age in Knight et al. [13] and

chronological age in Horvath [10] (see Figure S7 in the Supplement), estimates of the age effects among the CpGs that changed from birth to age 7 showed the same direction of change as their corresponding estimated gestational age effects at birth in 97% of the 16,172 CpGs, which included 14,186 gestational age-associated effects that were not significant at the 5% FDR threshold. This concordance in direction of effect is unlikely to occur by chance ($P < 10^{-10}$, see Supplement for calculation), and indicates that the majority of the change in mean methylation levels from birth to age 7 was due to aging-related mechanisms rather than age-dependent environmental exposures.

To directly test the hypothesis that non-genetic factors tend to have larger effects on methylation levels at age 7 than at birth, we used our Bayesian model to estimate the proportion of CpGs in our study whose methylation was not associated with ancestry at birth but associated at age seven and the proportion that were associated with ancestry at birth but not at age 7. The former was greater than 14% while the latter was less than 1.5% using either inferred genetic ancestry or reported race as a measure of ancestry. Even though over 14% of all CpGs in our study had ancestry effects present at age 7 but not at birth, we were only able to identify 18 discordant IGA-CpGs and 4 discordant RR-CpGs at age 7 using a liberal threshold of $dis_g^{(7)} \geq 0.80$. That is, for almost all CpGs that are associated with ancestry at age 7 but not at birth, the expected ancestry effect sizes were quite small relative to the statistical error, making it impossible to assign the direction of effect on methylation changes with confidence (Figure S6). Therefore, while some CpG sites may be influenced by exposures that are correlated with ancestry at age 7 but not at birth, their effects were far too small to estimate in this sample size.

Discussion

The relationships between DNA methylation, chronological age, and ancestry have the potential to shed light on disease etiology and may help determine the relative genetic and environmental contributions to the observed inter-individual variability of the epigenome [9-15, 19-24]. While it has previously been shown that ancestry is related to DNA methylation in cross-sectional studies [19-24], and that statistically significant meQTLs are conserved as one ages [44], it has yet to be shown whether or not *ancestry-dependent* methylation marks are conserved as children age.

Even though there was substantial change in blood methylation levels over time among children in this cohort, inferred genetic ancestry and self-reported race effects on methylation were overwhelmingly conserved in both direction and magnitude from birth to age 7. This result is interesting in and of itself because it provides an example of perinatal epigenetic variation that persists later in life, and more generally an example of a persistent effect on DNA methylation levels, which has been cited as a critical area of future epigenetic research [31, 45]. The consistency of our estimates for the effect due to ancestry also demonstrates the fidelity of our processing step to account for unobserved factors like cell composition, since failure to account for latent covariates often leads to biased and irreproducible estimates [46, 47]. Furthermore, the novel statistical framework we used to infer effects that are conserved versus those that vary over time can be easily applied to other longitudinal DNA methylation data, as a way to avoid the spurious logic often used in applications of frequentist hypothesis testing that failing to reject the null hypothesis implies the null is true.

While the observation that inferred genetic ancestry and reported race effects are conserved from birth to age 7 gives credence to the hypothesis that the effects are genetic in nature, it does not rule out the possibility of environmental components or gene-environment interactions that could determine ancestry-related methylation prior to birth and persist as the

child ages. To further explore this, we showed that the IGA-CpGs were enriched among CpGs with meQTLs, and that methylation levels at many of the IGA-CpGs are mediated by local genotype, indicating that much of the ancestry-methylation correlation could be attributed to genetic variation. Moreover, the RR-CpGs were only a small subset of IGA-CpGs in our study. This is opposite to the findings of Galanter et al. [20], who argued that ancestry-dependent methylation patterns in admixed populations are in large part determined by differences in exposure histories. Because their data were cross-sectional they could not evaluate whether the observed patterns arose during childhood or were also present at birth. Our results provide evidence for genetics accounting for most of the correlation between methylation and ancestry, and implies that the genetic contribution to variability in blood methylation is substantial.

Our observations in support of strong genetically – and weak environmentally – determined ancestry-associated methylation patterns in blood may seem paradoxical to the plethora of studies showing that DNA methylation levels in blood cells are associated with environmental exposures, such as cadmium, arsenic and smoking, to name a few [5-8, 20, 48-52]. Whereas the estimated genetic effect sizes in our study are substantially larger than many of the environmentally-associated effects on methylation patterns previously reported, the effects of environmental exposures on methylation in blood are probably too small to estimate with even moderate to large sample sizes [31]. For example, it was only by performing a meta-analysis in 6,685 individuals that Joubert et al. [5] were able to identify 6,000 CpGs whose DNA methylation levels in blood from infants and adolescents were associated maternal smoking exposure. In one sense, we were able to corroborate previous observations of small non-genetic effects on methylation in blood by showing that while methylation patterns at an estimated 14% of all CpGs in our study were not correlated with ancestry at birth but correlated with ancestry at

age 7, the correlation at individual CpGs at age 7 was too small to be identified as statistically significant. We were also not able to find any statistically significant correlations between methylation at birth or at age 7 and any of the environmental exposure variables measured in this cohort. We note that cord blood cotinine levels, a measure of *in utero* tobacco smoke exposure, were above the level of detection in only 34 of the 196 mothers in our study.

An unsurprising feature of these longitudinal data is that average methylation levels of over 16,000 CpGs changed significantly from birth to age 7. However, what was quite remarkable was that the direction of the change in 97% of those CpGs matched the direction of their corresponding estimated gestational age effect at birth, which included over 14,000 gestational age-associated effects that were not genome-wide significant. Not only does this fit with the above narrative and suggest that methylation levels of the vast majority of the 16,172 age-related CpGs were in fact changing due to age-related mechanisms and not because of differences in environmental exposures at birth and age 7, it also indicates that the “epigenetic clock” present at birth may be the same as that present later in life. While we do not have the data to explore this further, this remains an important avenue of future research.

The results of our study suggest that DNA methylation levels in blood cells are fairly robust to environmental exposures, including those that are correlated with self reported race. A better understanding of tissue-specific methylation responses to environmental exposures could inform the design of future studies and provide insights into the mechanisms through which exposures and gene-environment interactions influence health and disease.

Materials and methods

Sample composition

URECA is a birth cohort study initiated in 2005 in Baltimore, Boston, New York City and St. Louis under the NIAID-funded Inner City Asthma Consortium [29]. Pregnant women were recruited. Either they or the father of their unborn child had a history of asthma, allergic rhinitis, or eczema, and deliveries prior to 34 weeks gestation were excluded (see Gern et al. [29] for full entry criteria). Informed consent was obtained from the women at enrollment and from the parent or legal guardian of the infant after birth.

Maternal questionnaires were administered prenatally and child health questionnaires administered to a parent or caregiver every 3 months through age 7 years. Gestational age at birth and obstetric history were obtained from medical records. Additional details on study design are described in Gern et al. [29] and in the Supplement. Frozen paired cord blood mononuclear cells (CBMCs) and peripheral blood mononuclear cells (PBMCs) at age 7, were available for 196 of the 560 URECA children after completing other studies. After QC (see below), DNA methylation data were available for 194 children at birth, 195 children at age 7, and 193 children at both time points; genotype data were available in 193 children (194 at birth; 195 at age 7) (Supplementary Table 1).

DNA Methylation

DNA for methylation studies was extracted from thawed CBMCs and PBMCs using the Qiagen AllPrep kit (QIAGEN, Valencia, CA). Genome-wide DNA methylation was assessed using the Illumina Infinium MethylationEPIC BeadChip (Illumina, San Diego, CA) at the University of Chicago Functional Genomics Facility (UC-FGF). Birth and 7-year samples from the same child were assayed on the same chip and the data were processed using Minfi [32]; Infinium type I and type II probe bias were corrected using SWAN [33]. Raw probe values were corrected for color

imbalance and background by control normalization. Three out of the 392 samples (two at birth and one at age 7) were removed as outliers following normalization. We removed 82,352 probes that mapped either to the sex chromosomes or to more than one location in a bisulfite-converted genome, had detection P values greater than 0.01% in 25% or more of the samples, or overlapped with known SNPs with minor allele frequency of at least 5% in African, American or European populations. After processing, 784,484 probes were retained and M-values were used for all downstream analyses, which were computed as $\log_2(\text{methylated intensity} + 100) - \log_2(\text{unmethylated intensity} + 100)$. The offset of 100 was recommended in Du et al. [34].

Genotyping

DNA from the 196 URECA children was genotyped with the Illumina Infinium CoreExome+Custom array. Of the 532,992 autosomal SNPs on the array, 531,755 passed Quality control (QC) (excluding SNPs with call rate <95%, Hardy-Weinberg P values <10⁻⁵, and heterozygosity outliers). We conducted all analyses in 293,696 autosomal SNPs with a minor allele frequency $\geq 5\%$. Genotypes for three children failed QC and were excluded from subsequent analysis that involved genotypes, including methylation quantitative locus (meQTL) mapping, inferred genetic ancestry, or used genetic ancestry PC1 as a covariate (see below). These three children were included in all other analyses.

Estimating inferred genetic ancestry

Ancestral principal component analysis (PCA) was performed using a set of 801 ancestry informative markers (AIMs) from Tandon et al. [35] that were genotyped in both the URECA children and in HapMap [36] release 23. Because PC1 captured the majority of variation in

genetic ancestry (Figure 1), we refer to PC1 as inferred genetic ancestry and consider it as a surrogate measure for percent African ancestry.

Statistical analysis

To determine the effect of gestational age on methylation in CBMCs, we used standard linear regression models with the child's gender, sample collection site, inferred genetic ancestry and methylation plate number as covariates in our model. We also estimated cell composition and other unobserved confounding factors using a method described in McKennan et al. [37]. We then computed a gestational age *P* value for each CpG site and used q-values [38] to control the false discovery rate at a nominal level. We took the same approach to determine CpGs whose methylation changed from birth to age 7, except the response was measured as the difference in methylation at birth and age 7. In this analysis, we included the child's gender, gestational age at birth, inferred genetic ancestry and sample collection site as covariates. Because all paired samples were on the same plate, we did not include plate number as a covariate in this analysis. We also estimated unobserved factors that influence differences in methylation at birth and age 7 using McKennan et al. [37] and included these latent factors in our linear model. See models (S1) and (S2) in the Supplement for more detail.

We used data from the self-reported Hispanic and Black individuals with methylation measured at both time points to analyze the effect of ancestry (either inferred genetic ancestry or self-reported race) on methylation at birth and age 7 jointly using a Bayesian model. We did not include the 10 individuals of other reported races in this analysis because we did not want our estimates to be influenced by the groups with small samples sizes. We included age (birth or age 7), sample collection site, gestational age at birth, gender and methylation plate number as

covariates in our model, and estimated additional unobserved covariates (including cell composition) using a method specifically designed for correlated data [39]. Once we estimated the relevant hyper-parameters, we extended the sign rate paradigm developed in Stephens [40] to perform inference in longitudinal data. This is discussed in more detail in the context of the specific questions we present in the results section. We encourage the reader to review the Supplement for a more detailed presentation of this model and previously discussed models.

We performed meQTL mapping in the 145 genotyped, self-reported Black children using the set of 269,622 SNPs with 100% genotype call rate in this subset. We restricted ourselves to this subset of samples to minimize heterogeneity in effect sizes. To identify CpG-SNP pairs, we considered SNPs within 5kb of each CpG, as this region has been previously shown to contain the majority of genetic variability in DNA methylation [1] and is small enough to mitigate the multiple testing burden, and computed a P value for the effect of the genotype at a single SNP on methylation at the corresponding CpG with ordinary least squares. We then defined the meQTL for each CpG site as the SNP with the lowest P value. In addition to genotype, we included inferred genetic ancestry (i.e., ancestry PC1), gestational age at birth, gender, sample collection site and methylation plate number in the linear model, along with the first nine principal components of the residual methylation data matrix after regressing out the intercept and the five additional covariates. We then tested the null hypothesis that a CpG did not have an meQTL in the 10kb region by using the minimum marginal P value in the region as the test statistic and computed its significance via bootstrap. Lastly, we used q-values to control the false discovery rate.

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References

1. Bell JT, Pai AA, Pickrell JK, Gaffney DJ, Pique-Regi R, Degner JF, et al. DNA methylation patterns associate with genetic and gene expression variation in HapMap cell lines. *Genome Biol.* 2011;12(1):R10. Epub 2011/01/22. doi: 10.1186/gb-2011-12-1-r10. PubMed PMID: 21251332; PubMed Central PMCID: PMC3091299.
2. Smith AK, Kilaru V, Kocak M, Almli LM, Mercer KB, Ressler KJ, et al. Methylation quantitative trait loci (meQTLs) are consistently detected across ancestry, developmental stage, and tissue type. *BMC Genomics.* 2014;15:145. Epub 2014/02/22. doi: 10.1186/1471-2164-15-145. PubMed PMID: 24555763; PubMed Central PMCID: PMC3091299.
3. Chatterton Z, Hartley BJ, Seok MH, Mendeleev N, Chen S, Milekic M, et al. In utero exposure to maternal smoking is associated with DNA methylation alterations and reduced neuronal content in the developing fetal brain. *Epigenetics Chromatin.* 2017;10:4. Epub 2017/02/06. doi: 10.1186/s13072-017-0111-y. PubMed PMID: 28149327; PubMed Central PMCID: PMC5270321.
4. Goodrich JM, Dolinoy DC, Sanchez BN, Zhang Z, Meeker JD, Mercado-Garcia A, et al. Adolescent epigenetic profiles and environmental exposures from early life through peri-adolescence. *Environ Epigenet.* 2016;2(3):dvw018. Epub 2016/08/14. doi: 10.1093/eep/dvw018. PubMed PMID: 29492298; PubMed Central PMCID: PMC5804533.
5. Joubert BR, Felix JF, Yousefi P, Bakulski KM, Just AC, Breton C, et al. DNA Methylation in Newborns and Maternal Smoking in Pregnancy: Genome-wide Consortium Meta-analysis. *Am J Hum Genet.* 2016;98(4):680-96. Epub 2016/04/05. doi: 10.1016/j.ajhg.2016.02.019. PubMed PMID: 27040690; PubMed Central PMCID: PMC4833289.
6. Kippler M, Engstrom K, Mlakar SJ, Bottai M, Ahmed S, Hossain MB, et al. Sex-specific effects of early life cadmium exposure on DNA methylation and implications for birth weight. *Epigenetics.* 2013;8(5):494-503. Epub 2013/05/07. doi: 10.4161/epi.24401. PubMed PMID: 23644563; PubMed Central PMCID: PMC3741219.
7. Koestler DC, Avissar-Whiting M, Houseman EA, Karagas MR, Marsit CJ. Differential DNA methylation in umbilical cord blood of infants exposed to low levels of arsenic in utero. *Environ Health Perspect.* 2013;121(8):971-7. Epub 2013/06/13. doi: 10.1289/ehp.1205925. PubMed PMID: 23757598; PubMed Central PMCID: PMC3733676.
8. Rzehak P, Saffery R, Reischl E, Covic M, Wahl S, Grote V, et al. Maternal Smoking during Pregnancy and DNA-Methylation in Children at Age 5.5 Years: Epigenome-Wide-Analysis in the European Childhood Obesity Project (CHOP)-Study. *PLoS One.* 2016;11(5):e0155554. Epub 2016/05/14. doi: 10.1371/journal.pone.0155554. PubMed PMID: 27171005; PubMed Central PMCID: PMC4865176.

- 468 9. Bocklandt S, Lin W, Sehl ME, Sanchez FJ, Sinsheimer JS, Horvath S, et al. Epigenetic
469 predictor of age. PLoS One. 2011;6(6):e14821. Epub 2011/07/07. doi:
470 10.1371/journal.pone.0014821. PubMed PMID: 21731603; PubMed Central PMCID:
471 PMCPMC3120753.
- 472 10. Horvath S. DNA methylation age of human tissues and cell types. Genome Biol.
473 2013;14(10):R115. Epub 2013/10/22. doi: 10.1186/gb-2013-14-10-r115. PubMed PMID:
474 24138928; PubMed Central PMCID: PMCPMC4015143.
- 475 11. Horvath S, Erhart W, Brosch M, Ammerpohl O, von Schonfels W, Ahrens M, et al.
476 Obesity accelerates epigenetic aging of human liver. Proc Natl Acad Sci U S A.
477 2014;111(43):15538-43. Epub 2014/10/15. doi: 10.1073/pnas.1412759111. PubMed PMID:
478 25313081; PubMed Central PMCID: PMCPMC4217403.
- 479 12. Johnson AA, Akman K, Calimport SR, Wuttke D, Stolzing A, de Magalhaes JP. The role
480 of DNA methylation in aging, rejuvenation, and age-related disease. Rejuvenation Res.
481 2012;15(5):483-94. Epub 2012/10/27. doi: 10.1089/rej.2012.1324. PubMed PMID: 23098078;
482 PubMed Central PMCID: PMCPMC3482848.
- 483 13. Knight AK, Craig JM, Theda C, Baekvad-Hansen M, Bybjerg-Grauholm J, Hansen CS,
484 et al. An epigenetic clock for gestational age at birth based on blood methylation data. Genome
485 Biol. 2016;17(1):206. Epub 2016/10/09. doi: 10.1186/s13059-016-1068-z. PubMed PMID:
486 27717399; PubMed Central PMCID: PMCPMC5054584.
- 487 14. Levine ME, Crimmins EM. Evidence of accelerated aging among African Americans and
488 its implications for mortality. Soc Sci Med. 2014;118:27-32. Epub 2014/08/03. doi:
489 10.1016/j.socscimed.2014.07.022. PubMed PMID: 25086423; PubMed Central PMCID:
490 PMCPMC4197001.
- 491 15. Marioni RE, Shah S, McRae AF, Chen BH, Colicino E, Harris SE, et al. DNA
492 methylation age of blood predicts all-cause mortality in later life. Genome Biol. 2015;16:25.
493 Epub 2015/01/31. doi: 10.1186/s13059-015-0584-6. PubMed PMID: 25633388; PubMed Central
494 PMCID: PMCPMC4350614.
- 495 16. Parets SE, Conneely KN, Kilaru V, Fortunato SJ, Syed TA, Saade G, et al. Fetal DNA
496 Methylation Associates with Early Spontaneous Preterm Birth and Gestational Age. PLoS One.
497 2013;8(6):e67489. Epub 2013/07/05. doi: 10.1371/journal.pone.0067489. PubMed PMID:
498 23826308; PubMed Central PMCID: PMCPMC3694903.
- 499 17. Schroeder JW, Conneely KN, Cubells JC, Kilaru V, Newport DJ, Knight BT, et al.
500 Neonatal DNA methylation patterns associate with gestational age. Epigenetics.
501 2011;6(12):1498-504. Epub 2011/12/06. doi: 10.4161/epi.6.12.18296. PubMed PMID:
502 22139580; PubMed Central PMCID: PMCPMC3256334.

- 503 18. Simpkin AJ, Suderman M, Gaunt TR, Lyttleton O, McArdle WL, Ring SM, et al.
504 Longitudinal analysis of DNA methylation associated with birth weight and gestational age.
505 Hum Mol Genet. 2015;24(13):3752-63. Epub 2015/04/15. doi: 10.1093/hmg/ddv119. PubMed
506 PMID: 25869828; PubMed Central PMCID: PMC4459393.
- 507 19. Adkins RM, Krushkal J, Tylavsky FA, Thomas F. Racial differences in gene-specific
508 DNA methylation levels are present at birth. Birth Defects Res A Clin Mol Teratol.
509 2011;91(8):728-36. Epub 2011/02/11. doi: 10.1002/bdra.20770. PubMed PMID: 21308978;
510 PubMed Central PMCID: PMC3429933.
- 511 20. Galanter JM, Gignoux CR, Oh SS, Torgerson D, Pino-Yanes M, Thakur N, et al.
512 Differential methylation between ethnic sub-groups reflects the effect of genetic ancestry and
513 environmental exposures. Elife. 2017;6. Epub 2017/01/04. doi: 10.7554/eLife.20532. PubMed
514 PMID: 28044981; PubMed Central PMCID: PMC5207770.
- 515 21. Heyn H, Moran S, Hernando-Herraez I, Sayols S, Gomez A, Sandoval J, et al. DNA
516 methylation contributes to natural human variation. Genome Res. 2013;23(9):1363-72. Epub
517 2013/08/03. doi: 10.1101/gr.154187.112. PubMed PMID: 23908385; PubMed Central PMCID:
518 PMC3759714.
- 519 22. Moen EL, Zhang X, Mu W, Delaney SM, Wing C, McQuade J, et al. Genome-wide
520 variation of cytosine modifications between European and African populations and the
521 implications for complex traits. Genetics. 2013;194(4):987-96. Epub 2013/06/25. doi:
522 10.1534/genetics.113.151381. PubMed PMID: 23792949; PubMed Central PMCID:
523 PMC3730924.
- 524 23. Mozhui K, Smith AK, Tylavsky FA. Ancestry dependent DNA methylation and influence
525 of maternal nutrition. PLoS One. 2015;10(3):e0118466. Epub 2015/03/06. doi:
526 10.1371/journal.pone.0118466. PubMed PMID: 25742137; PubMed Central PMCID:
527 PMC4350920.
- 528 24. Rahmani E, Shenhav L, Schweiger R, Yousefi P, Huen K, Eskenazi B, et al. Genome-
529 wide methylation data mirror ancestry information. Epigenetics Chromatin. 2017;10:1. Epub
530 2017/02/06. doi: 10.1186/s13072-016-0108-y. PubMed PMID: 28149326; PubMed Central
531 PMCID: PMC5267476.
- 532 25. Chan MA, Ciaccio CE, Gigliotti NM, Rezaiekhaliq M, Siedlik JA, Kennedy K, et al.
533 DNA methylation levels associated with race and childhood asthma severity. J Asthma.
534 2017;54(8):825-32. Epub 2016/12/09. doi: 10.1080/02770903.2016.1265126. PubMed PMID:
535 27929694.
- 536 26. Ladd-Acosta C, Hansen KD, Briem E, Fallin MD, Kaufmann WE, Feinberg AP.
537 Common DNA methylation alterations in multiple brain regions in autism. Mol Psychiatry.

- 2014;19(8):862-71. Epub 2013/09/04. doi: 10.1038/mp.2013.114. PubMed PMID: 23999529; PubMed Central PMCID: PMC4184909.
27. Nicodemus-Johnson J, Myers RA, Sakabe NJ, Sobreira DR, Hogarth DK, Naureckas ET, et al. DNA methylation in lung cells is associated with asthma endotypes and genetic risk. *JCI Insight*. 2016;1(20):e90151. Epub 2016/12/13. doi: 10.1172/jci.insight.90151. PubMed PMID: 27942592; PubMed Central PMCID: PMC4513990 PulmOne Advanced Medical Devices Ltd., Israel, and received reimbursement for expenses. He served on the Respiratory Therapy Clinical Advisory Board for Hollister Inc., and for this received honoraria and was reimbursed for travel and meal expenses incurred during meetings. He has received a research grant from AstraZeneca Inc., from 2006 to 2014, that was administered through the University of Chicago. He has multiple patents concerning a smooth muscle gene promoter and one pending concerning a method to determine respiratory physiological parameters (6090618; 6114311; 6284743; 6291211; 6297221; 6331527; 7169764). He has consulted for Novartis Institute for Biomedical Research, for which he received an honorarium and travel reimbursement. He was a member of the scientific advisory board for Cytokinetics Inc., for which he received honoraria and travel reimbursement.
28. Yokoyama AS, Rutledge JC, Medici V. DNA methylation alterations in Alzheimer's disease. *Environ Epigenet*. 2017;3(2):dvx008. Epub 2018/03/02. doi: 10.1093/eep/dvx008. PubMed PMID: 29492310; PubMed Central PMCID: PMC5804548.
29. Gern JE, Visness CM, Gergen PJ, Wood RA, Bloomberg GR, O'Connor GT, et al. The Urban Environment and Childhood Asthma (URECA) birth cohort study: design, methods, and study population. *BMC Pulm Med*. 2009;9:17. Epub 2009/05/12. doi: 10.1186/1471-2466-9-17. PubMed PMID: 19426496; PubMed Central PMCID: PMC2689166.
30. O'Connor GT, Lynch SV, Bloomberg GR, Kattan M, Wood RA, Gergen PJ, et al. Early-life home environment and risk of asthma among inner-city children. *J Allergy Clin Immunol*. 2017. Epub 2017/09/25. doi: 10.1016/j.jaci.2017.06.040. PubMed PMID: 28939248.
31. Breton CV, Marsit CJ, Faustman E, Nadeau K, Goodrich JM, Dolinoy DC, et al. Small-Magnitude Effect Sizes in Epigenetic End Points are Important in Children's Environmental Health Studies: The Children's Environmental Health and Disease Prevention Research Center's Epigenetics Working Group. *Environ Health Perspect*. 2017;125(4):511-26. Epub 2017/04/01. doi: 10.1289/EHP595. PubMed PMID: 28362264; PubMed Central PMCID: PMC5382002.
32. Aryee MJ, Jaffe AE, Corrada-Bravo H, Ladd-Acosta C, Feinberg AP, Hansen KD, et al. Minfi: a flexible and comprehensive Bioconductor package for the analysis of Infinium DNA methylation microarrays. *Bioinformatics*. 2014;30(10):1363-9. Epub 2014/01/31. doi: 10.1093/bioinformatics/btu049. PubMed PMID: 24478339; PubMed Central PMCID: PMC4016708.

- 574 33. Maksimovic J, Gordon L, Oshlack A. SWAN: Subset-quantile within array normalization
575 for illumina infinium HumanMethylation450 BeadChips. *Genome Biol.* 2012;13(6):R44. Epub
576 2012/06/19. doi: 10.1186/gb-2012-13-6-r44. PubMed PMID: 22703947; PubMed Central
577 PMCID: PMCPMC3446316.
- 578 34. Du P, Zhang X, Huang CC, Jafari N, Kibbe WA, Hou L, et al. Comparison of Beta-value
579 and M-value methods for quantifying methylation levels by microarray analysis. *BMC*
580 *Bioinformatics.* 2010;11:587. Epub 2010/12/02. doi: 10.1186/1471-2105-11-587. PubMed
581 PMID: 21118553; PubMed Central PMCID: PMCPMC3012676.
- 582 35. Tandon A, Patterson N, Reich D. Ancestry informative marker panels for African
583 Americans based on subsets of commercially available SNP arrays. *Genet Epidemiol.*
584 2011;35(1):80-3. Epub 2010/12/25. doi: 10.1002/gepi.20550. PubMed PMID: 21181899;
585 PubMed Central PMCID: PMCPMC4386999.
- 586 36. International HapMap C. The International HapMap Project. *Nature.*
587 2003;426(6968):789-96. Epub 2003/12/20. doi: 10.1038/nature02168. PubMed PMID:
588 14685227.
- 589 37. McKennan C, Nicolae DL. Accounting for unobserved covariates with varying degrees
590 of estimability in high dimensional biological data; 2018. Preprint. Available from: arXiv:
591 arXiv:1801.00865v2. Cited 22 August 2018
- 592 38. Storey JD. A direct approach to false discovery rates. *J Royal Stat Soc B.* 2002;64:479-
593 98.
- 594 39. McKennan C, Nicolae DL. Estimating and accounting for unobserved covariates in high
595 dimensional correlated data; 2018. Preprint. Available from: arXiv:1808.05895v1. Cited 22
596 August 2018
- 597 40. Stephens M. False discovery rates: a new deal. *Biostatistics.* 2017;18(2):275-94. Epub
598 2016/10/21. doi: 10.1093/biostatistics/kxw041. PubMed PMID: 27756721; PubMed Central
599 PMCID: PMCPMC5379932.
- 600 41. Nguyen AB, Moser R, Chou WY. Race and health profiles in the United States: an
601 examination of the social gradient through the 2009 CHIS adult survey. *Public Health.*
602 2014;128(12):1076-86. Epub 2014/12/03. doi: 10.1016/j.puhe.2014.10.003. PubMed PMID:
603 25457801.
- 604 42. Fu J, Wolfs MG, Deelen P, Westra HJ, Fehrmann RS, Te Meerman GJ, et al. Unraveling
605 the regulatory mechanisms underlying tissue-dependent genetic variation of gene expression.
606 *PLoS Genet.* 2012;8(1):e1002431. Epub 2012/01/26. doi: 10.1371/journal.pgen.1002431.
607 PubMed PMID: 22275870; PubMed Central PMCID: PMCPMC3261927.

- 608 43. Fritz MS, Mackinnon DP. Required sample size to detect the mediated effect. Psychol
609 Sci. 2007;18(3):233-9. Epub 2007/04/21. doi: 10.1111/j.1467-9280.2007.01882.x. PubMed
610 PMID: 17444920; PubMed Central PMCID: PMCPMC2843527.
- 611 44. Gaunt TR, Shihab HA, Hemani G, Min JL, Woodward G, Lyttleton O, et al. Systematic
612 identification of genetic influences on methylation across the human life course. Genome Biol.
613 2016;17:61. Epub 2016/04/03. doi: 10.1186/s13059-016-0926-z. PubMed PMID: 27036880;
614 PubMed Central PMCID: PMCPMC4818469.
- 615 45. Martin EM, Fry RC. Environmental Influences on the Epigenome: Exposure-Associated
616 DNA Methylation in Human Populations. Annu Rev Public Health. 2018. Epub 2018/01/13. doi:
617 10.1146/annurev-publhealth-040617-014629. PubMed PMID: 29328878.
- 618 46. Peixoto L, Risso D, Poplawski SG, Wimmer ME, Speed TP, Wood MA, et al. How data
619 analysis affects power, reproducibility and biological insight of RNA-seq studies in complex
620 datasets. Nucleic Acids Res. 2015;43(16):7664-74. Epub 2015/07/24. doi: 10.1093/nar/gkv736.
621 PubMed PMID: 26202970; PubMed Central PMCID: PMCPMC4652761.
- 622 47. Yao C, Li H, Shen X, He Z, He L, Guo Z. Reproducibility and concordance of
623 differential DNA methylation and gene expression in cancer. PLoS One. 2012;7(1):e29686.
624 Epub 2012/01/12. doi: 10.1371/journal.pone.0029686. PubMed PMID: 22235325; PubMed
625 Central PMCID: PMCPMC3250460.
- 626 48. Joubert BR, Haberg SE, Nilsen RM, Wang X, Vollset SE, Murphy SK, et al. 450K
627 epigenome-wide scan identifies differential DNA methylation in newborns related to maternal
628 smoking during pregnancy. Environ Health Perspect. 2012;120(10):1425-31. Epub 2012/08/02.
629 doi: 10.1289/ehp.1205412. PubMed PMID: 22851337; PubMed Central PMCID:
630 PMCPMC3491949.
- 631 49. Lee KW, Richmond R, Hu P, French L, Shin J, Bourdon C, et al. Prenatal exposure to
632 maternal cigarette smoking and DNA methylation: epigenome-wide association in a discovery
633 sample of adolescents and replication in an independent cohort at birth through 17 years of age.
634 Environ Health Perspect. 2015;123(2):193-9. Epub 2014/10/18. doi: 10.1289/ehp.1408614.
635 PubMed PMID: 25325234; PubMed Central PMCID: PMCPMC4314251.
- 636 50. Markunas CA, Xu Z, Harlid S, Wade PA, Lie RT, Taylor JA, et al. Identification of DNA
637 methylation changes in newborns related to maternal smoking during pregnancy. Environ Health
638 Perspect. 2014;122(10):1147-53. Epub 2014/06/07. doi: 10.1289/ehp.1307892. PubMed PMID:
639 24906187; PubMed Central PMCID: PMCPMC4181928.
- 640 51. Richmond RC, Simpkin AJ, Woodward G, Gaunt TR, Lyttleton O, McArdle WL, et al.
641 Prenatal exposure to maternal smoking and offspring DNA methylation across the lifecourse:
642 findings from the Avon Longitudinal Study of Parents and Children (ALSPAC). Hum Mol

- 643 Genet. 2015;24(8):2201-17. Epub 2015/01/02. doi: 10.1093/hmg/ddu739. PubMed PMID:
644 25552657; PubMed Central PMCID: PMC4380069.
- 645 52. Wu D, Yang H, Winham SJ, Natanzon Y, Koestler DC, Luo T, et al. Mediation analysis
646 of alcohol consumption, DNA methylation, and epithelial ovarian cancer. J Hum Genet.
647 2018;63(3):339-48. Epub 2018/01/13. doi: 10.1038/s10038-017-0385-8. PubMed PMID:
648 29321518.
649

Figure Legends

Figure 1: Estimated ancestry principal components (PCs) 1 and 2. Nearly all the variation in ancestry separates along PC1 in the URECA sample. Filled triangles represent the 196 URECA children in this study, with their self-reported race shown in different colors. Open circles are reference control samples from HapMap; red = Utah residents from northern and western Europe (CEU), yellow = east Asian (Chinese and Japanese); dark blue = Africans from Nigeria (Yoruban).

Figure 2: Overlapping ancestry CpGs at birth and at age 7. (a): IGA-CpGs in CBMCs at birth ($\hat{c}o\hat{r}^{(0)} \geq 0.95$) and PBMCs at age 7 ($\hat{c}o\hat{r}^{(7)} \geq 0.95$). Overlapping CpGs (violet) have $\hat{c}o\hat{n}^{(0,7)} \geq 0.95$. (b): RR-CpGs at in CBMCs at birth ($\hat{c}o\hat{r}^{(0)} \geq 0.95$) and PBMCs at age 7 ($\hat{c}o\hat{r}^{(7)} \geq 0.95$). Overlapping CpGs (violet) have $\hat{c}o\hat{n}^{(0,7)} \geq 0.95$. (c): IGA-CpGs and RR-CpGs ($\hat{c}o\hat{r}^{(0)} \geq 0.95$) in CBMCs at birth. (d): IGA-CpGs and RR-CpGs ($\hat{c}o\hat{r}^{(0)} \geq 0.95$) in PBMCs at age 7.

Figure 3: IGA-CpGs are enriched for CpGs with meQTLs. (a) Illustration of the causal relationship between the methylation (M) at a CpG site, the genotype (G) at the SNP within ± 5 kb of the CpG that had the smallest meQTL P value and inferred genetic ancestry (IGA). Each graph corresponds to a unique CpG. (b) Plots of the meQTL P value for edge a in CBMCs at birth, where CpGs were stratified by whether or not it was an IGA-CpG at birth or age 7 ($\max(\hat{c}o\hat{r}_g^{(0)}, \hat{c}o\hat{r}_g^{(7)}) \geq 0.95$). The ten enlarged red circles are just for visual aid. (c) Plots of the logistic regression P value for edge c (Genotype $\sim$ IGA + Ancestry PC2 (see Figure 1)), stratified by whether or not the SNP was an IGA-meQTL.

Figure 4: Inferred genetic ancestry effects on methylation are primarily genetic in origin.

Histograms of $\max(\hat{c}o\hat{r}^{(0)}, \hat{c}o\hat{r}^{(7)})$ in the IGA (a) and the RR (b) analyses. If

674 $\max(\hat{c}or_g^{(0)}, \hat{c}or_g^{(7)})$ is close to 1 in the IGA or RR analysis, CpG g is an IGA-CpG or RR-
675 CpG, respectively, in CBMCs at birth or PBMCs at 7. The red histogram is created with
676 the set of 726 ethnicity-associated CpGs identified in Galanter et al. [20] that were also
677 among the 784,484 CpGs in our study and the blue are the remaining 783,758 CpGs.

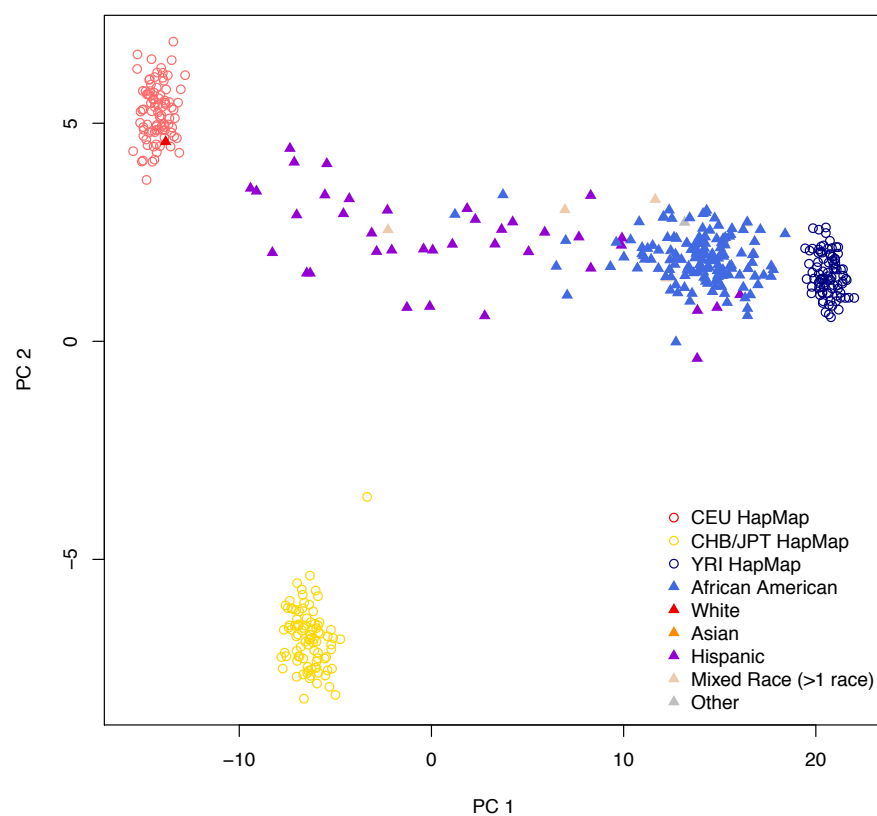


Figure 1

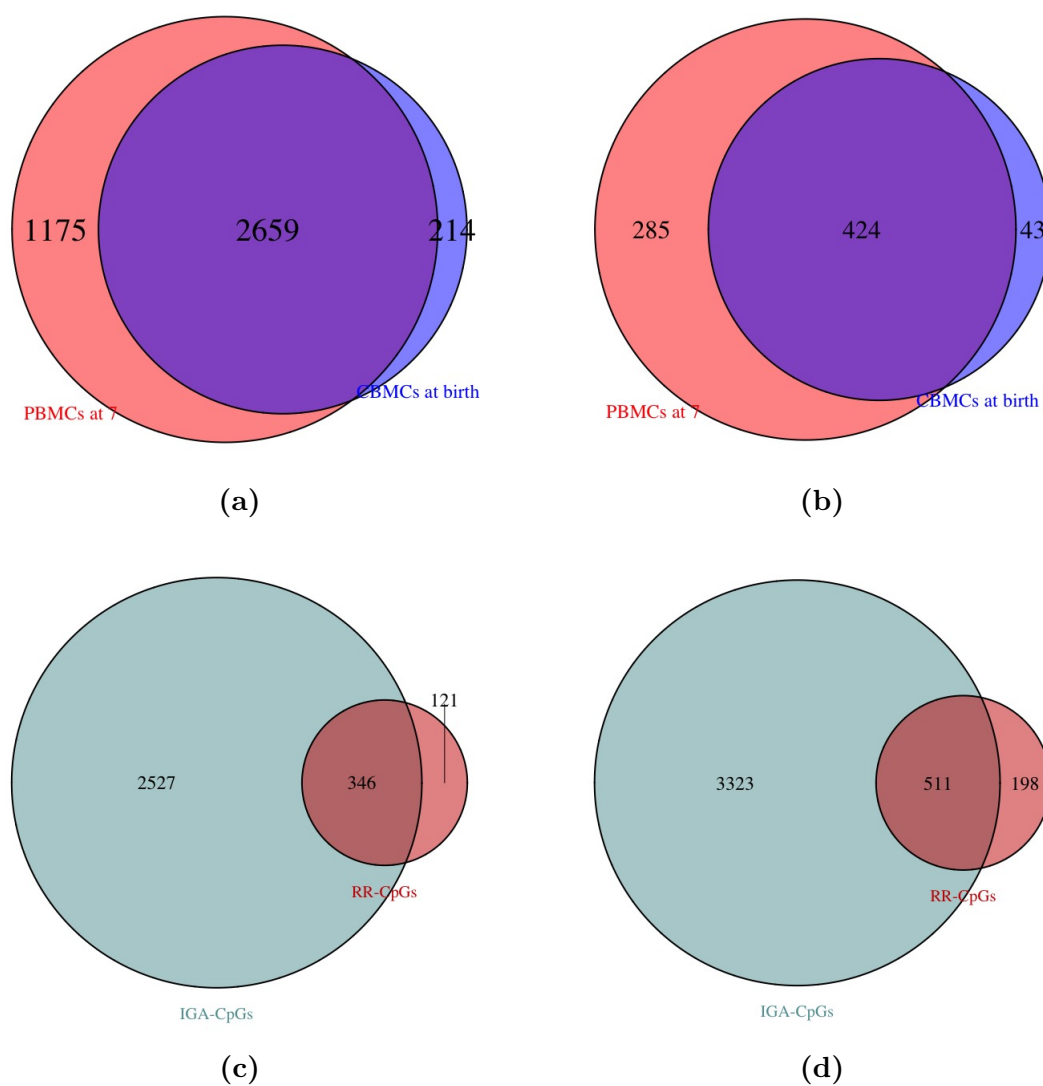


Figure 2

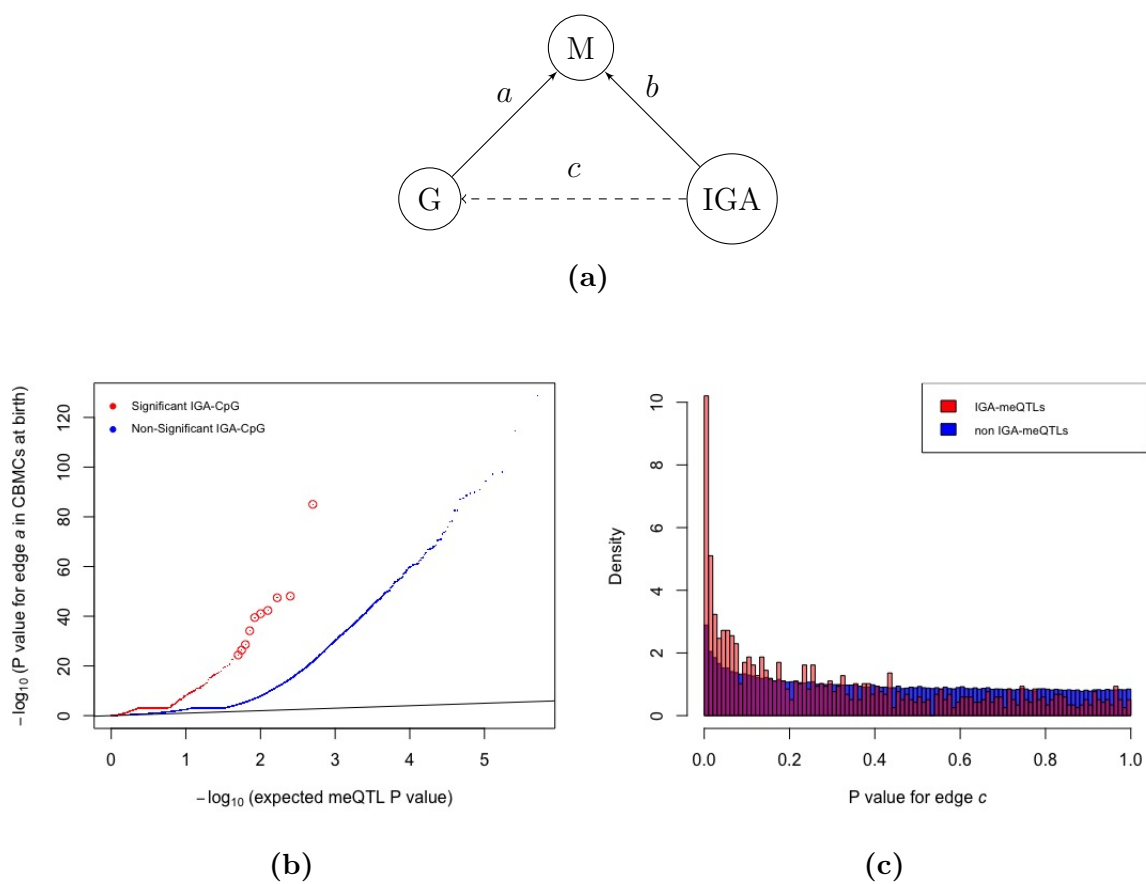


Figure 3

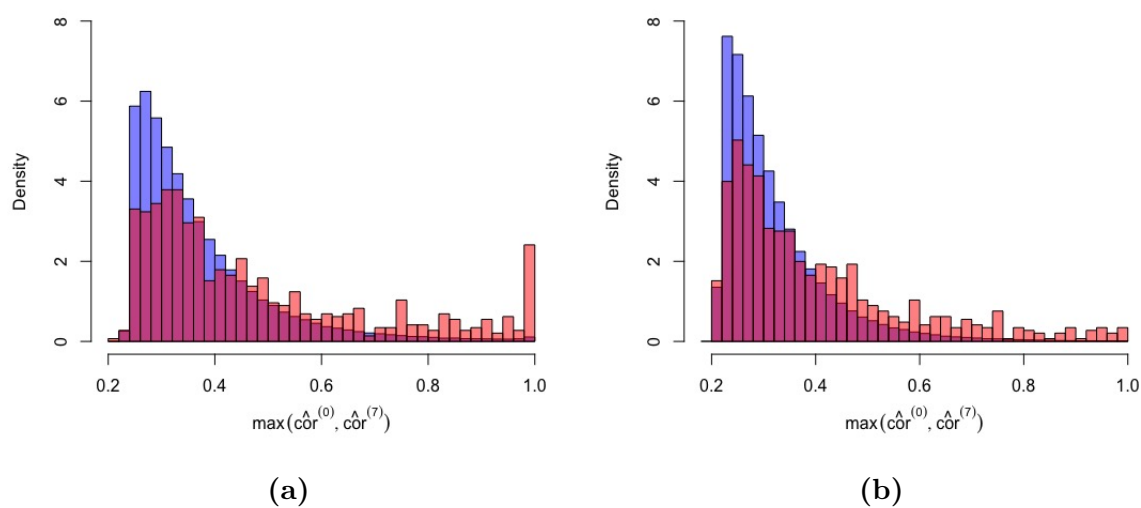


Figure 4