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Authors

Barfield, Richard
Wang, Heming
Liu, Yongmei
et al.

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ORIGINAL ARTICLE

Epigenome-wide association analysis of daytime sleepiness in the Multi-Ethnic Study of Atherosclerosis reveals African-American-specific associations

Richard Barfield^{1,2,✉}, Heming Wang^{3,4,5,✉}, Yongmei Liu⁶, Jennifer A. Brody^{7,✉}, Brenton Swenson^{7,8}, Ruitong Li^{4,10}, Traci M. Bartz^{7,8}, Nona Sotoodehnia⁷, Yii-der I. Chen⁹, Brian E. Cade^{3,4,5,✉}, Han Chen^{10,11,12,✉}, Sanjay R. Patel^{13,✉}, Xiaofeng Zhu¹⁴, Sina A. Gharib^{15,✉}, W. Craig Johnson¹⁶, Jerome I. Rotter⁹, Richa Saxena^{3,4,5,17,✉}, Shaun Purcell^{3,18}, Xihong Lin^{10,19}, Susan Redline^{4,5,20} and Tamar Sofer^{4,5,10,*}

¹Department of Epidemiology, University of Washington, Seattle, WA, ²Public Health Sciences Division, Fred Hutchinson Cancer Research Center, Seattle, WA, ³Program in Medical and Population Genetics, Broad Institute, Cambridge, MA, ⁴Division of Sleep and Circadian Disorders, Department of Medicine, Brigham and Women's Hospital, Boston, MA, ⁵Division of Sleep Medicine, Harvard Medical School, Boston, MA, ⁶Department of Epidemiology and Prevention, Division of Public Health Sciences, Wake Forest University School of Medicine, Winston-Salem, NC, ⁷Cardiovascular Health Research Unit, Department of Medicine, University of Washington, Seattle, WA, ⁸Institute for Public Health Genetics, University of Washington, Seattle, WA, ⁹The Institute for Translational Genomics and Population Sciences, Departments of Pediatrics and Medicine, Los Angeles Biomedical Research Institute at Harbor-UCLA Medical Center, Torrance, CA, ¹⁰Department of Biostatistics, Harvard T. H. Chan School of Public Health, Boston, MA, ¹¹Human Genetics Center, Department of Epidemiology, Human Genetics and Environmental Sciences, School of Public Health, The University of Texas Health Science Center at Houston, Houston, TX, ¹²Center for Precision Health, School of Public Health & School of Biomedical Informatics, The University of Texas Health Science Center at Houston, Houston, TX, ¹³Division of Pulmonary, Allergy and Critical Care Medicine, University of Pittsburgh Medical Center, Pittsburgh, PA, ¹⁴Department of Population and Quantitative Health Sciences, School of Medicine, Case Western Reserve University, Cleveland, OH, ¹⁵Computational Medicine Core, Center for Lung Biology, University of Washington Medicine Sleep Center, Division of Pulmonary, Critical Care, and Sleep Medicine, University of Washington, Seattle, WA, ¹⁶Department of Biostatistics, University of Washington, Seattle, WA, ¹⁷Center for Genomic Medicine and Department of Anesthesia, Pain, and Critical Care Medicine, Massachusetts General Hospital, Boston, MA, ¹⁸Department of Psychiatry, Brigham & Women's Hospital, Harvard Medical School, Boston, MA, ¹⁹Department of Statistics, Harvard University, Cambridge, MA and ²⁰Division of Pulmonary, Critical Care, and Sleep Medicine, Beth Israel Deaconess Medical Center, Boston, MA

*Corresponding author. Tamar Sofer, Brigham and Women's Hospital, Sleep and Circadian Disorders, Room 225C, 221 Longwood Ave, Boston, MA 02115. Email: tsofer@bwh.harvard.edu.

Abstract

Study Objectives: Daytime sleepiness is a consequence of inadequate sleep, sleep-wake control disorder, or other medical conditions. Population variability in prevalence of daytime sleepiness is likely due to genetic and biological factors as well as social and environmental influences. DNA methylation (DNAm) potentially influences multiple health outcomes. Here, we explored the association between DNAm and daytime sleepiness quantified by the Epworth Sleepiness Scale (ESS).

Methods: We performed multi-ethnic and ethnic-specific epigenome-wide association studies for DNAm and ESS in the Multi-Ethnic Study of Atherosclerosis (MESA; $n = 619$) and the Cardiovascular Health Study ($n = 483$), with cross-study replication and meta-analysis. Genetic variants near ESS-associated DNAm were analyzed for methylation quantitative trait loci and followed with replication of genotype-sleepiness associations in the UK Biobank.

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Results: In MESA only, we detected four DNAm-ESS associations: one across all race/ethnic groups; three in African-Americans (AA) only. Two of the MESA AA associations, in genes *KCTD5* and *RXRA*, nominally replicated in CHS (p -value < 0.05). In the AA meta-analysis, we detected 14 DNAm-ESS associations (FDR q -value < 0.05, top association p -value = 4.26×10^{-8}). Three DNAm sites mapped to genes (*CPLX3*, *GFAP*, and *C7orf50*) with biological relevance. We also found evidence for associations with DNAm sites in *RAI1*, a gene associated with sleep and circadian phenotypes. UK Biobank follow-up analyses detected SNPs in *RAI1*, *RXRA*, and *CPLX3* with nominal sleepiness associations.

Conclusions: We identified methylation sites in multiple genes possibly implicated in daytime sleepiness. Most significant DNAm-ESS associations were specific to AA. Future work is needed to identify mechanisms driving ancestry-specific methylation effects.

Statement of Significance

Excessive daytime sleepiness is associated with negative health outcomes such as reduction in quality of life, increased workplace accidents, and cardiovascular mortality. There are race/ethnic disparities in excessive daytime sleepiness. However, the environmental and biological mechanisms for these differences are not yet understood. We performed an association analysis of DNA methylation (DNAm), measured in monocytes, and daytime sleepiness within a racially diverse study population. We detected numerous DNAm markers associated with daytime sleepiness in African-Americans, but only one marker was associated with daytime sleepiness across all race/ethnic groups. Future work is required to elucidate the pathways between DNAm, sleepiness, and related behavioral/environmental exposures.

Key words: diversity; epigenetics; race/ethnic heterogeneity; excessive daytime sleepiness; sleep-wake; methylation; genomics

Introduction

Excessive daytime sleepiness (EDS), is estimated to affect between 10% and 20% of the population [1, 2]. EDS is associated with numerous adverse clinical, behavioral and public health outcomes, including work and vehicular accidents [3–10]; reduced health-related quality of life [11–13]; cognitive and performance deficits [14, 15]; and increased rates of stroke and total and cardiovascular mortality [16, 17]. There are multiple mechanisms for EDS, including insufficient sleep duration due to behavioral, social or work-related factors; sleep disruption due to a sleep disorder (sleep apnea, periodic limb movement disorder); circadian misalignment; the presence of a primary disorder of hypersomnia that affects the central sleep-wake control processes (e.g. narcolepsy); as well as poorly understood mechanisms occurring in a variety of medical (e.g. diabetes) and psychiatric (e.g. depression) disorders [18–20]. The prevalence of EDS has increased over the past decades, likely due to an increase in working hours and availability of electronic devices [3]. A rise in EDS prevalence may also reflect the increased prevalence of obesity, which is associated with sleepiness [21, 22]. This association is hypothesized to be due to obesity-related co-morbidities that reduce sleep quality, including obstructive sleep apnea, as well as to metabolic and neuroendocrine effects of adipokines on wake-promoting neurons [23].

Although EDS is prevalent in the population, there appears to be large inter-individual differences in propensity for sleepiness following sleep deprivation, sleep fragmentation, or sleep apnea [24, 25]. Similarly, there is significant population variability in prevalence of EDS [26]. Differences in sleepiness, including vulnerability or resilience to sleep-disrupting influences, have been suggested to be due to genetic and other biological differences, although social and environmental influences also likely play a role [27, 28]. Generally, the bases for differences in sleepiness between populations are not well understood.

Epigenetic modifications are increasingly recognized to mediate the impact of environmental influences on gene expression and on prevalence and severity of a wide range of health outcomes, including neuropsychiatric, metabolic and cardiovascular diseases. The most studied epigenetic marker, DNA methylation (DNAm), occurs when a methyl group is added to a cytosine that is followed by guanine on the genome (a “CpG” site). Changes in DNAm occur in response to a wide range of

exposures, many of which are associated with sleep and sleepiness, such as obesity, diet, and stress [29–35]. Therefore, we postulated that changes in DNAm would associate with variation in sleepiness and these changes may be population-specific, providing insights into underlying susceptibility to sleepiness that may be partially attributed to environmental, and possibly genetic, factors. To the best of our knowledge, there has been no reported study on DNAm and sleepiness.

In this article, we performed an Epigenome-Wide Association Study (EWAS) of daytime sleepiness quantified by the Epworth Sleepiness Scale (ESS) in the Multi-Ethnic Study of Atherosclerosis (MESA) and in the Cardiovascular Health Study (CHS). We leveraged the racial/ethnic diversity of the sample to explore potential differences in EWAS associations by background groups. We further used the MESA dataset to study potential mechanisms underlying the associations. First, using gene-expression analysis in MESA, studying whether DNAm associated with sleepiness by modifying gene expression; and second, by identifying methylation quantitative trait loci (cis-meQTLs) near genes harboring associated methylation sites to suggest whether DNAm affects sleepiness by modifying genetic effects. Candidate genetic associations were carried forward for replication analysis in the UK Biobank.

Methods

MESA study sample

The study population consisted of participants from the MESA, a prospective, longitudinal cohort study established to study factors associated with the development of cardiovascular disease. MESA clinic visits were first performed between 2000 and 2002 in six field centers across the United States when participants were free of known cardiovascular disease [36]. The subset of MESA individuals in this study is composed of those who participated in a sleep examination conducted in conjunction with MESA Exam 5 (described in detail previously [26, 36]), and who also participated in the MESA DNAm study [37]. The blood draws for the methylation study were obtained during MESA Exam 5 (2010–2012) [38] on a random subset of MESA participants at four of the six field centers: John Hopkins University, University of Minnesota, Columbia University, and Wake Forest University.

A total of 623 individuals were available with both sleep data and DNAm data. After excluding four individuals due to missing ESS scores, the final study sample included 619 individuals: 132 African-Americans (AAs), 202 Hispanic Americans (HAs), and 285 European Americans (EAs), where race/ethnicities were self-reported. The study was approved by the Institutional Review Boards of each participating site, and participants provided written informed consent, including the use of genetic data.

DNAm collection and processing in MESA

Methods for collection and assays for DNAm have been described previously [38]. In brief, peripheral blood was separated into mononuclear cells (CD14+) within two hours of collection. DNAm in monocytes was measured using the Illumina HumanMethylation450 BeadChip. Residual cell contamination in the monocyte data was assessed using Gene Set Enrichment Analysis, providing enrichment scores for neutrophils, B cells, T cells, and Natural Killer cells from Gene Expression data collected using the Illumina HumanHT-12 v4 Expression BeadChip and Illumina Bead Array Reader [38, 39]. The DNAm data underwent quality control tests prior to analysis using the lumi Bioconductor package [40]. These included color bias adjustments via smooth quantile normalization, median background adjustment, standard quantile adjustment, checks for potential sex or race mismatches, and outlier detection via multi-dimensional plots. Additional details on preprocessing of the DNAm data can be found in Liu et al. [37]. Following DNAm preprocessing, each of 484882 methylation probes for each person has a Beta-value, representing the proportion of methylated monocytes at that site and person.

We further prepared the methylation data for analysis based on meta information and sample-specific characteristics, as follows. First, we excluded 61219 probes that were within 10bp of a single nucleotide polymorphism (SNP) with a minor allele frequency greater than 1% in European, African, or American populations from the 1000 genomes reference data [41]. This was done to exclude loci where variation is solely due to genetic polymorphisms. We also removed any non-cg methylation probes and probes on sex chromosomes. We then removed any cross-reactive probes as defined by Chen et al. [42], leaving us with 399526 DNAm probes to analyze. We used the software program ComBat [43] on these CpG sites to remove any signal due to technical artifacts of chip and position on chip effects while maintaining correlation with our primary set of covariates (self-reported ancestry, recruitment site, sex, age, and residual cell type enrichment). ComBat was run on the M-values (logit transformed Beta-values [44], see [Supplementary Methods](#)). After ComBat, the data was transformed back to the Beta-Value scale for analysis.

SNP data

For consenting individuals, DNA was extracted from whole blood and genotyped on Affymetrix 6.0 GWAS array. Standard quality control methods for SNP- and sample-level quality were applied, including the exclusion of participants and SNPs with over 5% missing call rates. This resulted in 895289 genotyped variants in 615 individuals with DNAm data. For downstream analysis, we excluded SNPs with minor allele frequency (MAF) less than 1%. Further details on the genotype and quality control can be found in Vargas et al. [45]. We calculated the top 5

principal components (PCs) from the Linkage-Disequilibrium pruned set SNPs with $MAF \geq 5\%$. PCs were calculated in the combined sample and also in each ethnic-specific group. There were four AA individuals that had missing SNP information; therefore, the PCs were imputed with the mean value of the respective PC in the AA subset that had SNP information.

Sleep assessments

As part of the MESA Sleep Exam (2010–2013), participants completed standardized questionnaires and underwent single night in-home polysomnography (Compumedics Somte Systems, Abbotsville, Australia, AU0) and 7-day wrist actigraphy (Philips-Respironics Spectrum, Murrysville, PA), as described before [26]. The primary sleep measure was daytime sleepiness as quantified by the ESS, an 8-item validated instrument that asks the individual to assess likelihood of dozing off in a variety of daily activities using a 4 point (0–3) scale [46]. ESS scores vary from 0 to 24, with higher scores denoting more sleepiness [46]. ESS was assessed within one year of blood draw for DNAm. We identified additional sleep phenotypes/exposures that might be associated with both DNAm and ESS, potentially acting as mediators or confounders of any DNAm and ESS association. These measures were: insomnia/insomnia symptoms (report of doctor-diagnosed insomnia or the Women's Health Initiative (WHI) Insomnia Rating Scale [47], a validated scale varying from 0 to 20); Apnea-Hypopnea Index [AHI] (sum of all apneas and hypopneas associated with 3% or more oxygen desaturation divided by total sleep time by polysomnography); overnight hypoxemia (percentage of sleep time t with oxyhemoglobin saturation less than 90% [Per90]); and sleep duration (average sleep duration over the 7 monitoring nights by actigraphy as described before [26]).

Covariates

Age, sex and race/ethnicity were self-reported. Other behavioral, socioeconomic and lifestyle exposures that may confound sleepiness-methylation associations were also assessed, as follows. Alcohol use was self-reported (yes/no). Smoking status was classified by ever/former/never. Depressive symptoms were based on responses to the Center for Epidemiological Studies (CES) depression scale [48]. Socioeconomic variables were defined by having less than a college education and mother having less than a college education; these were added because they were previously shown in MESA to be associated with methylation in inflammatory genes [49]. Body mass index (BMI) was calculated from weight and height measured at MESA Exam 5. Dietary variables were derived from a food frequency questionnaire: long chain score (sum of Omega fatty acids mg/d), total fats, carbohydrates, and Total Alternative Healthy Eating Index - 2010 (Total AHEI-2010) [50]. Antidepressant and anti-psychotic medications usage, potentially influencing sleepiness, were identified from medication inventories.

The CHS

The CHS is a population-based cohort study of risk factors for coronary heart disease and stroke in adults ≥ 65 years conducted across four field centers [51]. The original predominantly

European ancestry cohort of 5201 persons was recruited in 1989–1990 from random samples of the Medicare eligibility lists; subsequently, an additional predominantly AA cohort of 687 persons was enrolled for a total sample of 5888. CHS was approved by institutional review committees at each field center and individuals in the present analysis had available DNA and provided informed consent for use of genetic information. DNAm was measured on 336 self-reported European American and 329 AA participants at study year 5. The samples were randomly selected among participants without coronary heart disease, congestive heart failure, peripheral vascular disease, valvular heart disease, stroke or transient ischemic attack at study baseline or lack of available DNA. Sleepiness was assessed at year 6 using the ESS.

Methylation measurements were performed at the Institute for Translational Genomics and Population Sciences at the Harbor-UCLA Medical Center—Los Angeles Biomedical Research Institute using the Infinium HumanMethylation450 BeadChip (Illumina Inc, San Diego, CA). Quality control was performed in the minfi R package [52–54] (version 1.12.0, <http://www.bioconductor.org/packages/release/bioc/html/minfi.html>). Samples were removed for any of the following: low median intensities of below 10.5 (log₂) across the methylated and unmethylated channels; a proportion of probes falling detection of greater than 0.5%; QC probes falling greater than 3SD from the mean; sex-check mismatches; or failed concordance with prior genotyping or >0.5% of probes with a detection *p*-value > .01. In total, 11 samples were removed for sample QC resulting in a sample of 323 European-ancestry and 326 AA samples. Methylation values were normalized using the SWAN quantile normalization method [54]. Since white blood cell proportions were not directly measured in CHS they were estimated from the methylation data using the Houseman method [55].

Statistical analysis

Our complete analysis approach is depicted in [Figure 1](#). We first describe our discovery analyses, which includes a meta-analysis, followed by sensitivity, gene expression, local methylation expressive quantitative trait loci (cis-meQTL), and replication analyses.

Discovery analysis for methylation-ESS association

The primary analysis outcome was ESS score, square-root transformed to achieve approximate normality. It was analyzed in a linear regression on normalized DNAm Beta-values, treated as exposures. The primary analyses were adjusted for sex, age, residual cell type enrichment (in MESA) or estimated white blood cell counts (in CHS), recruitment site, and the top 5 genetic PCs. CHS analyses were further adjusted for current smoking status (in MESA, smoking status was adjusted in sensitivity analyses, as described below). We performed discovery analysis via two different approaches. The first approach focused exclusively on each cohort (MESA, CHS), followed by cross-replication, if associations were detected. We conducted analyses in both the combined multi-ethnic sample, and in self-reported race/ethnic group-specific analyses. In MESA, combined analyses adjusted for PCs calculated in the overall group, as well as for a covariate for self-reported ethnicity (EA, AA, or HA). In group-specific analyses, we adjusted for their specific PCs. In CHS, group-specific

analyses were combined in inverse-variance fixed-effects meta-analysis. We evaluated heterogeneity across race/ethnic-specific results by fitting a linear model in MESA using the overall sample with self-reported race/ethnicity interaction terms with DNAm and testing these interaction terms using an *F*-test via analysis of variance (ANOVA). For analyses reported in MESA only, we report replication *p*-values in the appropriate race/ethnic group (or combined) in CHS. Replication *p*-values are computed as the one-sided *p*-value, guided by the direction of association in MESA. This approach has been shown to be more powerful and have better control of type 1 error compared with two-sided *p*-values for replication [56].

The second discovery approach was a meta-analysis combining both CHS AA and EA with their respective similar MESA groups, as well as a meta-analysis across all groups from both studies. The meta-analysis was done using inverse variance weighted, fixed-effects approach.

To account for multiple testing, we controlled the false discovery rate (FDR) at the 5% level in each of the multi-ethnic and race/ethnic group-specific analyses. An association was deemed as significant if the *q*-value was less than 0.05 [57].

Sensitivity analysis for methylation-ESS associations

We next identified behavioral, socioeconomic and lifestyle exposures, including variation in sleep and sleep disturbances, that may influence the sleepiness-methylation associations and further adjusted for these factors in sensitivity analyses conducted in significant probe-trait primary associations. Covariates considered for the sensitivity analysis were: BMI, sleep duration (actigraphy-based), insomnia (both doctor diagnosed and the WHI insomnia rating scale), AHI, nocturnal hypoxemia [Per90], alcohol use, smoking status, CES depression scale, having less than a college education, mother having less than a college education, and dietary variables, as defined earlier. We also examined moderate sleep apnea (AHI > 15), short sleep (<5 hours), and overnight hypoxemia (Per90 > 5%). Complete details for this sensitivity analysis are provided in the [Supplementary Material](#).

In another sensitivity analysis, we repeated analyses in MESA for the top association, after excluding individuals who reported use of antidepressant or antipsychotic medications.

Screening for cis-meQTLs in MESA

We hypothesized that genetic associations underlie some of the DNAm-ESS associations. Due to our limited sample size, we implemented a screening process on genetic variants in the CpG region, followed by a replication analysis of the SNP-sleepiness association in an independent cohort (UK Biobank).

First, we mapped all DNAm sites that passed the significance threshold (*q*-value < 0.05) to their respective genes to assess if genetic variants in that gene were associated with both ESS and DNAm. Specifically, for each gene that a top DNAm site mapped to, we performed association analysis between genotyped SNPs within 100kb of the gene (as exposures) and the associated DNAm site (as an outcome) adjusting for age, sex, top five PCs, study site, and residual cell type. This identified local (cis) meQTLs (screening step 1). SNPs with associations at *p*-value < 0.01 were next assessed for their association with sleepiness, with and without additional adjustment for the DNAm (screening step 2). These two screening steps were performed on the MESA subsample of

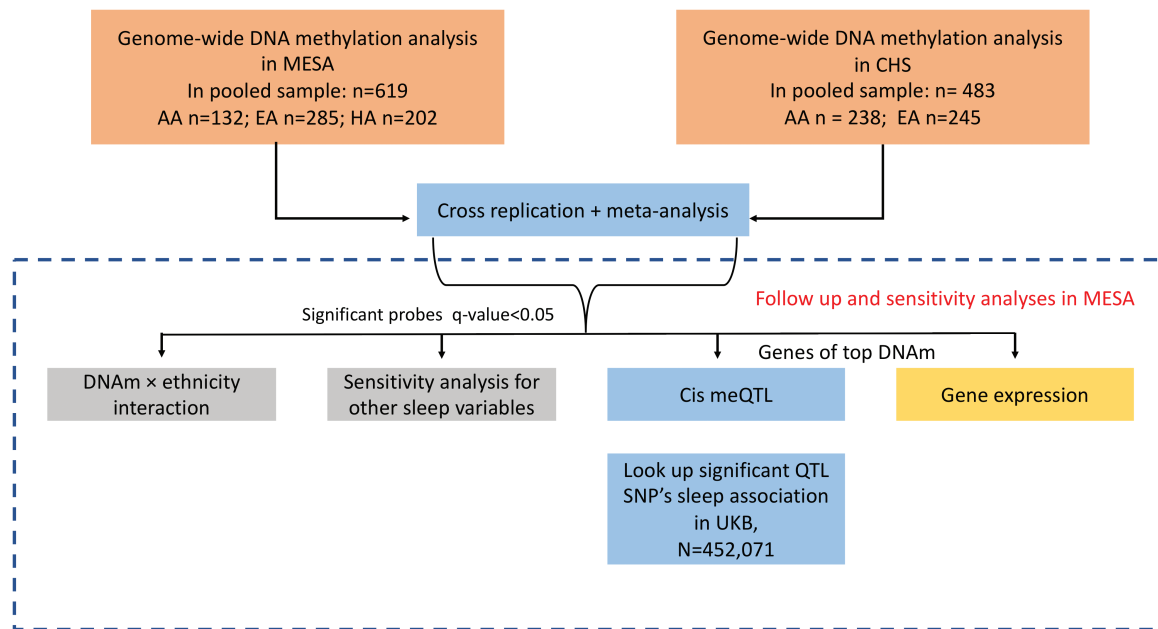


Figure 1. Analysis workflow for assessing associations between DNAm and ESS.

the race/ethnic group in which the DNAm-ESS association was detected, and in all groups if the association was detected in the combined analysis. SNPs that were associated with ESS with p -value < 0.05 were carried forward to independent replication analysis (see below) of the SNP-sleepiness association in the UK Biobank [58]. We also report SNPs-ESS associations in MESA race/ethnic groups that did not identify the original DNAm-ESS association, while noting that such associations may not be detected due to lower sample size and power.

Replication analysis of cis-meQTL-sleepiness association in the UK Biobank

The UK Biobank is a large prospective study for a wide range of genetics and health outcomes in over 500 000 participants aged 40–69 years recruited from 2006 to 2010 in the United Kingdom [59]. Genotyping and imputation data are described in Bycroft et al. (2017) [60]. Sleepiness was assessed by self-reported responses to the question “How likely are you to dose off or fall asleep during the daytime when you don’t mean to? (e.g. when working, reading or driving)” with the options of “Never/rarely”, “sometimes”, “often”, “all of the time”, “do not know”, and “prefer not to answer.” For SNPs passing the cis-meQTL screening process in MESA, association analysis was performed in 452 071 individuals adjusted for age, sex, genotyping array, 10 PCs and genetic relatedness matrix.

Gene expression analysis

For genes corresponding to the DNAm sites with FDR q -value < 0.05 , we used gene expression data in MESA to study the evidence for association between DNAm and gene expression (of the same gene), and gene expression association with ESS. Gene expression profiling and processing are described in Liu et al. [37]. In brief, gene expression was measured using Illumina HumanHT-12 v4 Expression BeadChip array in mononuclear

cells, on the same participants with DNAm measures. We performed (1) association analysis of DNAm and gene expression, using all individuals with available data (234 AA, 386 HA, and 582 EA individuals), with ESS as the outcome and DNAm as exposure; and (2) association analysis of gene expression (as exposure) and squared-root ESS (as the outcome) in the set of individuals that had both gene expression and ESS measured. Complete details are provided in the [Supplementary Material](#).

Results

Study characteristics

The MESA sample consisted of 619 individuals (53% female) with a mean age of 68 years, with 132 AA, 202 HA, and 285 EA individuals (Table 1). The mean ESS was 6.0 and 14% were classified as having EDS (ESS > 10), with the highest prevalence of EDS among AAs (18%; $p = 0.07$). Sleep duration was significantly shorter in the AA group compared to the other groups (p -value < 0.001). Mean AHI was 19.6, and 46% of the sample had moderate or more severe sleep apnea (AHI > 15). Across groups, sleep apnea was more prevalent and severe in the HA group (Table 1). Significant differences were observed for BMI, having a college education, smoking status, sleep apnea, sleep duration, dietary long-chain score, and alcohol consumption by race/ethnic group (Table 1). We provide information on the median of these variables as well as the number of missing values in [Supplementary Table 1](#).

The CHS sample consisted of 483 participants (63% females) with 238 AA and 245 EA (Table 2). The mean age was 74 years and the mean ESS was 6.0 with 15% being classified as having EDS, with similar trends for a higher prevalence of EDS among AAs (Table 2).

Associations between sleepiness and DNAm

We report histograms of p -values from the various analyses as well as inflation factors in [Supplementary Figure 1](#). The genomic

Table 1. MESA study participants characteristics

	Overall	EA	AA	HA	p-value
Total	619	285	132	202	
Age (years)	68.72 (9.24)	69.58 (9.55)	69.05 (8.90)	67.29 (8.88)	0.024
Sex, males (%)	290 (46.8%)	138 (48.4%)	53 (40.2%)	99 (49.0%)	0.219
Recruitment site (%)					<0.001
Columbia University	193 (31.2%)	38 (13.3%)	51 (38.6%)	104 (51.5%)	
John Hopkins University	165 (26.7%)	86 (30.2%)	79 (59.8%)	0 (0.0%)	
University of Minnesota	236 (38.1%)	138 (48.4%)	0 (0.0%)	98 (48.5%)	
Wake Forest University	25 (4.0%)	23 (8.1%)	2 (1.5%)	0 (0.0%)	
ESS Score	6.05 (4.04)	5.93 (3.78)	6.74 (4.45)	5.76 (4.10)	0.075
Excessive Daytime Sleepiness (ESS > 10)	85 (13.7%)	38 (13.3%)	24 (18.2%)	23 (11.4%)	0.204
BMI (kg/m ²)	29.79 (5.55)	28.96 (5.55)	30.73 (5.51)	30.35 (5.43)	0.002
AHI (events/hour)	19.59 (18.79)	19.12 (19.60)	17.68 (17.77)	21.45 (18.25)	0.184
Moderate to Severe Apnea (AHI > 15)	267 (46.05%)	109 (42.2%)	53 (41.7%)	105 (53.8%)	0.027
Nocturnal Hypoxemia (%time < 90% saturation)	4.22 (10.28)	5.25 (12.48)	3.34 (9.44)	3.43 (6.97)	0.097
Nocturnal Hypoxemia ≥ 5%	108 (18.6%)	56 (21.7%)	18 (14.2%)	34 (17.4%)	0.177
Sleep Duration (minutes)	391.42 (79.83)	410.20 (71.77)	360.48 (87.11)	385.13 (78.76)	<0.001
Less than 5 hours of sleep a night	73 (11.9%)	19 (6.8%)	32 (24.6%)	22 (11.0%)	<0.001
Less than a college education (%)	204 (33.0%)	54 (18.9%)	42 (32.1%)	108 (53.5%)	<0.001
Mother having less than a college education (%)	485 (80.8%)	197 (69.9%)	107 (82.9%)	181 (95.8%)	<0.001
Doctor Diagnosed Insomnia	34 (5.5%)	11 (3.9%)	9 (6.8%)	14 (6.9%)	0.262
WHI Insomnia Rating Scale	7.65 (4.68)	7.44 (4.63)	7.67 (4.51)	7.95 (4.86)	0.494
CES Depression Scale	8.33 (7.34)	7.82 (6.93)	8.20 (7.21)	9.15 (7.94)	0.144
Currently Drink Alcohol	286 (46.3%)	167 (58.6%)	53 (40.2%)	66 (32.8%)	<0.001
Smoking Status					0.005
Never	242 (39.2%)	102 (35.8%)	50 (37.9%)	90 (44.8%)	
Former	325 (52.6%)	163 (57.2%)	62 (47.0%)	100 (49.8%)	
Current	51 (8.3%)	20 (7.0%)	20 (15.2%)	11 (5.5%)	
Nutrients: Total Fat (g)	72.97 (44.14)	73.06 (38.90)	79.11 (53.50)	68.68 (44.13)	0.113
Nutrients: Total Carbohydrates (g)	215.13 (115.39)	207.70 (90.12)	226.91 (144.91)	218.06 (125.33)	0.264
Long chain fatty acid score (mg/day)	2.56 (2.36)	2.23 (2.03)	3.32 (2.80)	2.53 (2.36)	<0.001
Total AHEI-2010	54.00 (10.48)	54.85 (10.70)	52.88 (11.43)	53.51 (9.38)	0.148
Antipsychotic medications	4 (0.6%)	1 (0.4%)	1 (0.8%)	2 (1.0%)	0.675
Non-tricyclic Antidepressants (other than MAOI)	79 (12.8%)	57 (20.1%)	4 (3.0%)	18 (9.0%)	<0.001
Tricyclic antidepressants	12 (1.9%)	7 (2.5%)	2 (1.5%)	3 (1.5%)	0.689

Data presented as mean (SD) or n (%). CES depression scale originated from the Center for Epidemiological Studies. AHEI is the Alternative Healthy Eating Index. WHI is the Women's Health Initiative, and the scale ranges from 0 to 20. Test of difference were either done via regression (continuous variables) or chi-squared test (categorical variable).

Table 2. CHS study participants characteristics

	Overall	AA	EA	p-value
Total	483	238	245	
Age, years	74.03 (5.3)	72.79 (5.4)	75.23(4.9)	<0.001
Sex, males (%)	177 (37%)	82 (34%)	95 (38%)	0.370
Epworth Sleepiness Scale Score	5.98 (4.1)	6.38 (4.4)	5.58 (3.8)	0.037
Excessive Daytime Sleepiness (ESS > 10)	72 (15%)	43 (18%)	29 (12%)	0.073
BMI (kg/m ²)	27.8 (5)	29.0 (4.9)	26.7 (4.8)	<0.001
Currently drink alcohol	209 (43%)	97 (41%)	112 (46%)	0.310
Smoking status				0.071
Never	216 (45%)	108 (45%)	108 (44%)	
Former	207 (43%)	93 (39%)	114 (47%)	
Current	60 (12%)	37 (16%)	23 (9%)	

Data is presented as mean (SD) or n (%).

inflation ranged from 0.97 (CHS combined) to 1.09 (MESA AA). Results from the discovery analysis of each cohort with cross-replication in the other cohort are reported in Table 3. This analysis identified four significant associations (FDR q -value < 0.05), all observed in MESA. A single DNAm-ESS association was detected in the combined analysis across MESA race/ethnic

groups at cg25472882 located near gene *C7orf50* (p -value = 6.59×10^{-8}). There were three DNAm-ESS associations observed in the AA group: cg12833508 (p -value = 1.38×10^{-7}), cg26609398 (p -value = 2.33×10^{-7}), and cg13687497 (p -value = 3.66×10^{-7}) located near genes *AP1S3*, *KCTD5*, and *RXRA*, respectively (Table 3). For each of these AA-specific associations, tests of interaction

between race/ethnic group and DNAm on the transformed ESS scores conducted in the overall MESA sample were significant (Table 3). The race/ethnic associations between DNAm in these four sites are visualized in Figure 2. Replication (one-sided) p -values in CHS are reported in Table 3. Two DNAm sites were nominally associated with ESS in CHS: cg26609398 (replication p -value = 0.028), and cg13687497 (replication p -value = 0.045). Considering DNAm sites with FDR q -value < 0.1, there were

two significant associations in DNAm sites within *RAI1*, a gene associated with several sleep and circadian phenotypes [61, 62]. Therefore, we further investigated these DNAm sites in follow-up analyses.

In a second discovery step, we performed meta-analysis of all DNAm for the two cohorts, by race/ethnic groups and across all groups (Table 4). Fourteen associations (FDR q -value \leq 0.05) were detected, all in the meta-analysis of MESA AA and

Table 3. Methylation sites associated (FDR q -value \leq 0.05) with the Epworth Sleepiness Scale (square-root transformed) in MESA either in AA ($n = 132$) or in the combined analysis ($n = 619$) along with their association in CHS AA ($n = 238$) or CHS combined ($n = 483$)

Probe	CHR	Location	Analyses race/ethnic group	Gene	Effect estimate in MESA	MESA			Effect estimate in CHS ^c	p -value in CHS ^c	p -value meta analysis ^d
						p -value ^a	q -value ^a	p -value ^b			
cg25472882	7	1112040	Combined	<i>C7orf50</i>	15.26	6.59e-08	0.0237	0.312	-1.77	0.49	0.00125
cg12833508	2	224701802	AA	<i>AP1S3</i>	-17.31	1.38e-07	0.0393	0.00021	1.19	0.54	1.42e-02
cg26609398	16	2752235	AA	<i>KCTD5</i>	-14.88	2.33e-07	0.0412	0.00067	-3.30	0.028	4.95e-06
cg13687497	9	137249839	AA	<i>RXRA</i>	-20.98	3.66e-07	0.0393	0.00024	-3.90	0.045	2.63e-05

^aModels adjusted for age, sex, residual cell type enrichment, site of recruitment, and top five principal components calculated within the group listed in the analysis column. If Analyses column indicates 'Combined', adjustment was also made for self-reported race/ethnicity.

^bModels adjusted for age, sex, residual cell type enrichment, recruitment site, top five principal components in the overall sample, and self-reported race/ethnicity when testing for interaction between DNAm Beta-value and self-reported race/ethnicity in overall sample.

^cAssociation results in respective CHS analysis. p -value is based on one-sided p -values informed by the direction of effect.

^d p -value from meta-analysis of MESA and CHS. Genomic locations are provided in genome build 37.

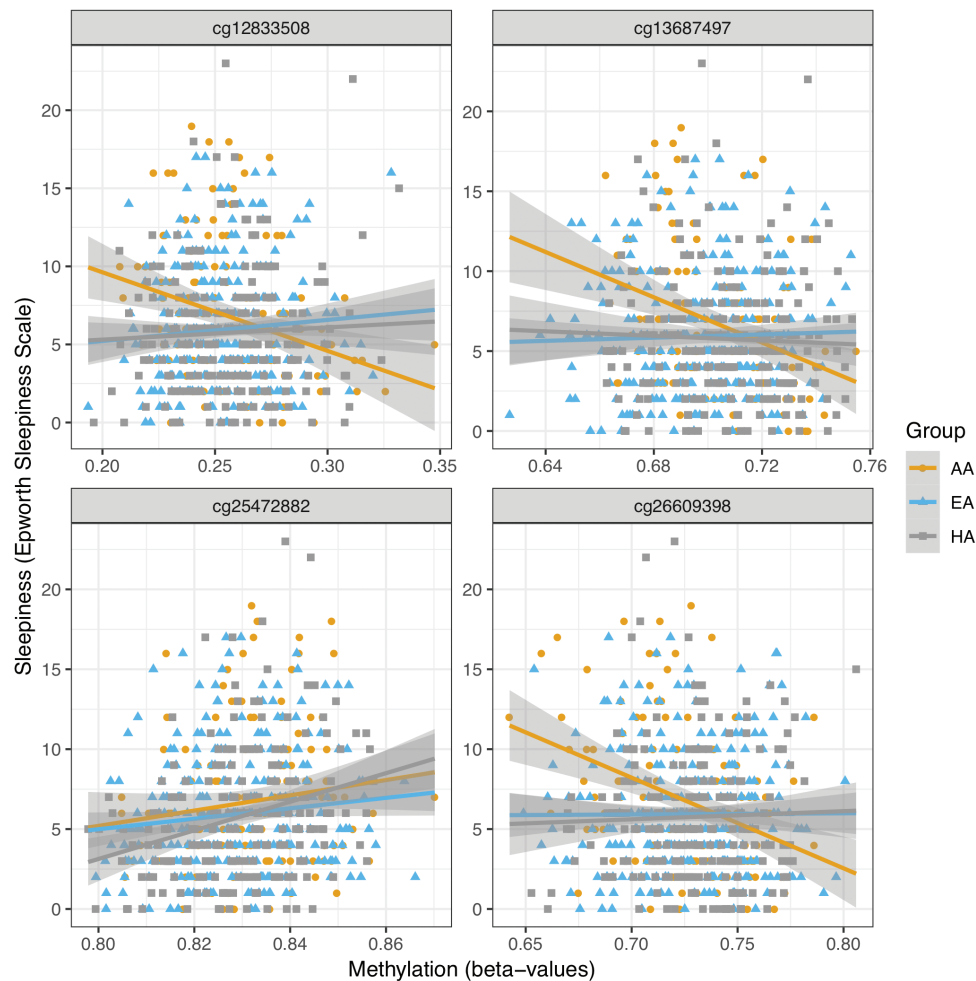


Figure 2. Four ESS-DNAm associations detected in MESA and reported in Table 3. The bottom left panel displays an association that was detected in the (combined) MESA sample, while the other panels display associations that were detected only in the AA group.

CHS AA. Of these, eight DNAm sites had p -value < 0.05 in a test for interaction of DNAm and self-reported race/ethnicity (Supplementary Table 2-Sheet 2). The top associated DNAm site was cg18371750 (meta p -value = 4.26×10^{-8}) located in the HRH2 gene. In 13 of the 14 DNAm sites, decreased methylation was correlated with an increase in ESS (Table 4).

Supplementary Table 2-Sheet 1 reports association results from all MESA and CHS race/ethnic groups for all associations reported in Tables 3 and 4. Supplementary Tables 3 and 4 report results from sensitivity analysis. Overall, the analysis did not highlight any phenotype as driving any of the observed DNAm-ESS associations, with the exception that the ESS association with cg03737424 significantly changed in the model with the sensitivity variables (permutation p -value = 0.03). Removing medication users from the analyses resulted in little difference in results.

Cis-meQTL analysis and UK Biobank replication

We performed cis-meQTL analysis for the DNAm reported in Tables 3 and 4, which mapped to 14 genes. Across the genes, we tested 877 SNPs. Of these, 20 SNPs passed screening step 1 (p -value < 0.01 ; Supplementary Table 5), three of which passed screening step 2. Rs9896285 of RAI1 was associated with ESS both in the model with and without adjustment to DNAm at the site of interest. In contrast, rs10785870 of RXRA and rs7495739 of LMAN1L/CPLX3 were associated with ESS only in the model that was unadjusted for DNAm at the relevant site, suggesting that DNAm mediated an association between these SNPs and sleepiness.

These SNPs were carried forward for testing the SNP-sleepiness association in the UK Biobank data set of European Ancestry individuals (Supplementary Table 6). An association was deemed replicated if it had a replication p -value threshold $< 0.05/3 = 0.017$, where replication p -value is a one-sided p -value guided by the direction of association in the discovery sample. Only rs9896285 in RAI1 replicated with p -value = 6.5×10^{-5} . Notably, the two other associations were nominally significant, with replication p -values 0.024 and 0.038.

Table 4. Methylation sites associated (FDR q -value ≤ 0.05) with the Epworth Sleepiness Scale (square-root transformed) in the meta-analysis of MESA AA ($n = 132$) and CHS AA ($n = 238$)

Probe	CHR.	Location	Gene	Effect estimate in MESA AA ^a	MESA p -value ^a	Effect estimate in CHS AA	p -value in CHS AA	p -value in meta-analysis	q -value in meta-analysis
cg18371750	5	175112799	HRH2	-22.94	1.02E-02	-22.47	2.65E-06	4.26E-08	0.0138
cg26130090	12	54615724		-7.85	2.82E-05	-6.16	7.89E-04	1.26E-07	0.0164
cg23574298	2	55382186		-17.11	1.21E-03	-16.16	6.05E-05	1.53E-07	0.0164
cg01904985	15	75117658	CPLX3/ LMAN1L	-18.19	1.24E-04	-14.90	2.92E-04	2.55E-07	0.0195
cg02374944	15	91463093	MAN2A2	-12.36	1.27E-05	-9.06	1.83E-03	3.02E-07	0.0195
cg13588265	17	1382042	MYO1C	-14.66	4.37E-03	-13.70	5.96E-05	6.21E-07	0.0335
cg01769243	17	42992917	GFAP	-13.19	3.06E-04	-10.96	5.69E-04	8.62E-07	0.0398
cg11335203	1	6398594	ACOT7	-14.23	2.65E-02	-15.19	3.89E-05	2.14E-06	0.0494
cg03737424	16	67550511		-11.84	1.53E-04	-9.42	3.71E-03	2.12E-06	0.0494
cg00181327	13	31439278		-17.86	7.61E-04	-15.58	4.74E-04	1.16E-06	0.0425
cg00583046	6	31024990	HCG22	-11.07	5.24E-05	-8.36	1.92E-03	1.18E-06	0.0425
cg00331852	16	88723987	MVD	-12.34	4.69E-05	-9.03	4.11E-03	1.98E-06	0.0494
cg13776718	9	139617752	FAM69B	-9.97	1.97E-03	-8.74	2.06E-04	1.83E-06	0.0494
cg21883754	19	54926437	TTYH1	21.19	1.09E-01	21.88	1.12E-05	1.89E-06	0.0494

^aModels adjusted for age, sex, residual cell type enrichment, site of recruitment, and top five principal components calculated in AA. Genomic location is provided in genome build 37.

Gene expression analysis

Comprehensive results are provided in Supplementary Tables 7 and 8. Briefly, we considered six genes corresponding to some of DNAm sites reported in Tables 3 and 4, and that also had expressed transcripts in the gene expression data set. Of the two DNAm sites associated with RAI1, cg14720773 had p -value = 0.05 in the combined cohort in association with expression of one of the RAI1 transcripts, but the expression of this transcript was not associated with ESS. In contrast, cg27208169 had p -value = 0.04 in the MESA AA analysis with a second RAI1 transcript, and increased expression of this transcript was associated with lower ESS (p -value = 0.002 in the combined cohort). None of the other associations were nominally significant.

Summary statistics and code availability

Code and results have been deposited on GitHub (https://github.com/tamartsi/sleepiness_methylation).

Discussion

This study reports for the first time a DNAm association analysis in a multi-ethnic population designed to detect DNAm markers associated with daytime sleepiness. We performed association analyses in two cohorts, MESA and CHS, using both a cross-replication approach and a meta-analysis, examining analyzing data for each race/ethnicity group (AA, EA, and HA) individually as well as by pooling groups. In the cross-replication analysis, two associated DNA sites discovered in MESA AAs (cg26609398, KCTD5 gene, and cg13687497, RXRA gene) replicated in CHS AAs. In the meta-analysis, we discovered additional 14 DNAm associations in 11 genes in CHS and MESA AAs. At a higher statistical threshold, we also identified DNAm sites in the RAI1 gene (FDR adjusted p -value < 0.1). Some of the genes that we identified have strong biological relevance and were previously reported as associated with several sleep traits [61–66].

Our study focused on the multi-ethnic epigenetic bases of daytime sleepiness. Several studies have shown that even after accounting for sleep duration and sleep disorders, sleepiness is lower in EA compared to other ethnic groups [20, 26, 67–69]. Compared to EA groups, AAs also have a higher prevalence of both short and long sleep durations and sleep apnea syndrome (defined by an elevated AHI plus daytime sleepiness [26]) and in the larger MESA cohort, AA had higher ESS than EA [26]. Race/ethnicity differences in sleep patterns and susceptibility to EDS appear early in life. AA children have been shown to have shorter sleep duration than white children as early as 1 year of age [70]. In children with sleep apnea treated with adenotonsillectomy, AA children have higher sleepiness scores than EA children both at baseline and following treatment [71]. Here, in our study of a middle-aged to older community multi-racial sample, we found significant epigenetic findings mostly in the AA samples of two independent cohorts. The specificity of the findings in the AA sample could reflect differences in underlying genetic architecture that influence susceptibility to methylation or differences in environmental exposures, such as air pollution, or stress that differentially affect different groups [72–75]. We address this hypothesis using sensitivity analyses that explored potential explanatory effects due to sleep variation and other lifestyle exposures. These analyses mostly did not find any phenotype explaining the association, when comparing the results to those obtained via permutation, with the exception of one DNAm-ESS association. However, these measures do not characterize lifetime exposures nor included precise measurements of the social or physical environments, emphasizing the need for additional research that links key aspects of an individual's environmental and sociological exposures with epigenetic markers and health conditions such as sleepiness as important routes for understanding health disparities.

A few of our sleepiness-associated DNAm sites are within genes that have strong evidence for associations with sleep-related processes. In the MESA discovery analysis, the most significant DNAm association was in the combined MESA cohort (cg25472882, p -value = 6.59×2^{-8} , CHS replication p -value = 0.75), in gene *C7orf50*. Among sub-Saharan Africans [76] different DNAm sites in this gene were associated with type 2 Diabetes, a condition commonly associated with sleep problems. Epigenetic analysis of cord blood samples showed that epigenetic changes in this gene also were associated with serum immunoglobulin E in children [77].

Three DNAm associations were discovered in MESA AA only, and two of these has replication p -value < 0.05 in CHS. One of these associations was in the *KCTD5* gene (cg26609398 p -value 2.33×10^{-7} , CHS AA replication p -value = 0.028). *KCTD5* is a mammalian ortholog of the *Drosophila* gene *Inc*, a gene expressed in multiple sleep associated regions in the central nervous system and shown to play a fundamental role in sleep in *Drosophila* by interacting with Cullin-3 ubiquitin ligase (*Cul3*). When mutated, both genes (*KCTD5* and *Cul3*) produce similar phenotypes characterized by markedly shortened and fragmented sleep together with altered synaptic structure and impaired synaptic transmission in relevant neuronal circuits [63]. This suggests that alterations in protein dynamics (e.g. ubiquitination) might disrupt synaptic function and impact sleep. Future research is needed to examine how epigenetic changes in *KCTD5* and similar genes influence gene expression and core elements of the sleep homeostat, with initial work indicating potential protein interactions with the circadian genes *FBXW11* and *BTRC* [78].

In MESA AA and replicated in CHS AA, DNA methylation association was also observed for cg13687497, within the *RXRA* gene, a retinoic acid receptor involved in lipid metabolism and cellular senescence [79–81]. Genes involved in retinoic acid pathways may be particularly relevant to sleepiness phenotypes given the central role of retinoic acid in central nervous system signaling pathways. Interestingly, two of our associated methylation sites (at a less significant statistical threshold) were within *RAI1*, which is induced by retinoic acid. *RAI1* encodes a protein that is highly expressed in neuronal tissues and is involved in early neural differentiation and transcriptional regulation of circadian clock components [62, 64–66]. Variants in this gene have been associated with circadian rhythm disruption, several abnormal sleep traits, neurobehavioral problems and obesity [64, 66]. In addition, SNPs in *RAI1* have been found to be associated with sleep apnea [61]. In addition to our DNAm-sleepiness association, we identified additional evidence supporting a role of *RAI1* in sleepiness: (1) expression of *RAI1* in mononuclear cells was associated with ESS in MESA; (2) our cis-meQTL analysis detected a SNP in *RAI1* influencing sleepiness in an EA population from the UK Biobank. Moreover, our analysis in MESA suggests that the SNP rs9896285 has both direct and indirect (through DNAm) effects on sleepiness in MESA AAs. The nonsignificant associations in MESA HAs and EAs could relate to differences in genetic architecture, environmental exposures or to lower power, which is a result of both low sample size and lower allele frequency in EAs and HAs compared to AAs (0.04 and 0.11 in MESA EAs and HAs, respectively, compared to 0.28 in MESA AAs). More work is needed to further dissect the association of retinoic acid pathway genes with sleepiness.

In the meta-analysis of MESA and CHS AAs, we discovered 14 DNAm associations with ESS, revealing relevant biological mechanism associated with sleepiness. The most significant association was in the Histamine Receptor H2 (*HRH2*) gene (cg18371750, AA meta p -value = 4.3×10^{-8}). The histaminergic system plays a central role in modulating brain activity, including alertness. H2 receptors are found in a wide variety of tissues, including neurons, where cross-talk with H1 receptors modulate complex signaling pathways [82]. Genetic variability in this gene is associated with some neurological diseases (e.g. Parkinson, autism [82–84]) and in pain reception in knockout mice [85]. Another association was in the *CPLX3/LMAN1L* genes region (cg01904985, p -value = 2.5×10^{-7}). *CPLX3* is involved in light sensitization and in thalamo-cortical integration, which are key for sleep regulation [86, 87]. Both genes are highly expressed in brain tissue (see Supplementary Figure 2). Our cis-meQTL analysis identified rs7495739 associated with these genes which predicted methylation, was associated with ESS in MESA AAs (p -value = 0.034), and was also nominally associated with sleepiness in UK Biobank whites (replication p -value = 0.038). Another gene highlighted by the meta-analysis is the Glial Fibrillary Acidic Protein (*GFAP*, cg01769243, p -value = 8.6×10^{-7}), which is also highly expressed in brain tissues (Supplementary Figure 3), and is associated with nervous system disease [88].

Cedernaes et al. [89] recently reported numerous associations between experimental sleep loss in 15 individuals and DNAm in adipose tissue. Of the 148 DMRs reported (Supplementary Table 2 in Cedernaes et al.), seven DNAm sites (cg25265360, cg05481452, cg01771651, cg25265360, cg23781467, cg20815339, and cg01168201) were within 5000 bp of a DNAm site associated with ESS in our analysis of MESA AA at the FDR < 0.1 level. Interestingly, Cedernaes et al. reported that most of

the differentially methylated regions following sleep loss were hypermethylated, while in our analysis, sleepiness was mostly associated with hypomethylation in the top associated DNAm sites. We studied whether this could be due to a global pattern of hypomethylation among participants with EDS in MESA. We computed mean global methylation for all participants and compared the distribution of these means between individuals with EDS, and others. However, the mean and median global methylation averages were nearly identical between the groups (Supplementary Figure 4). There are multiple differences in the study by Cedernaes et al. [89] and our, including differences in sleep exposures, the age of study participants and tissue assayed (adipose tissue vs blood).

A unique feature of this study is its multi-ethnic sample. In MESA alone, we identified three DNAm-sleepiness associations that are only present in AAs, and one association in the combined population. Two of the AA-specific associations replicated in CHS. We further identified 14 AA-specific associations in the meta-analysis of MESA and CHS. DNAm differences in some of these sites may explain some of the differences in sleepiness between AAs and EAs, highlighting potential mechanisms to explore in studies of health disparities. A limitation is that we had small sample sizes in the group-specific analyses. Despite this, we could directly replicate two of the associations discovered in MESA AAs. Lack of further replication may relate to many factors, including the older sample of CHS participants, biases and misclassification in self-reported sleepiness assessment [90], and potential difference in their exposure history compared to MESA participants. Another limitation is the time difference in measurement of DNAm, and sleep study. Participants filled the ESS questionnaire within a year following the blood draw for DNAm, and both DNAm and ESS may vary over this time frame. In addition, we examined DNAm within monocytes, while sleep is organized in the brain. Despite this, we identified associated DNAm sites near genes with potential neurological impact. A limitation which is shared by many epigenome-wide association studies, is the lack of identified exposure explaining the change in methylation and the group-specificity of the associations. We did study potential mechanism by considering cis-meQTLs analysis and a sensitivity analysis adjusting for various, plausible, exposure variable (sleep, lifestyle, and nutrition variables) which may be independently associated with either DNAm, ESS, or both, but these analyses did not lead to strong hypotheses about causal pathways.

In conclusion, we identified multiple DNAm sites associated with sleepiness. For the majority of these sites, sleepiness was associated with decreased methylation. Some of our top DNAm sites are near genes known to be associated with critical neuronal processes that influence sleep. Taken together, our findings suggest that these sleepiness-associated DNAm sites are related to genes that have a biologically meaningful function with regards to sleep. The significance of our findings, which were mostly in AAs, identifies biological areas that may contribute to the observed differences in sleepiness between AAs and EAs, providing an avenue for further investigation, both genetic and environmental, of health disparities. As mentioned in the introduction, poor sleep outcomes are associated with negative life outcomes and a recent review presented extensive evidence that obstructive sleep apnea and aging share various biological pathways [91]. Future studies with larger sample sizes may elucidate these genomic pathways and how they relate to potential

biological or sociological patterns that are being reflected in excessive sleepiness.

Supplementary material

Supplementary material is available at SLEEP online.

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