

## **Title:** An epigenome-wide analysis of DNA methylation, racialized and economic inequities, and air pollution

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## 1 Key points

2 **Question:** Could DNAm be a mechanism by which adversity becomes embodied?

3 **Findings:** Traffic-related air pollution exposure may induce epigenetic changes related to  
4 inflammatory processes; and there are suggestive associations with measures of structural racism.

5 **Meaning:** DNAm may be a biological mechanism through which structural racism and air pollution  
6 become biologically embodied.

## 7 Abstract

8 **Importance:** DNA methylation (DNAm) provides a plausible mechanism by which adverse exposures  
9 become embodied and contribute to health inequities, due to its role in genome regulation and  
10 responsiveness to social and biophysical exposures tied to societal context. However, scant  
11 epigenome-wide association studies (EWAS) have included structural and lifecourse measures of  
12 exposure, especially in relation to structural discrimination.

13 **Objective:** Our study tests the hypothesis that DNAm is a mechanism by which racial discrimination,  
14 economic adversity, and air pollution become biologically embodied.

15 **Design:** A series of cross-sectional EWAS, conducted in My Body My Story (MBMS, biological  
16 specimens collected 2008-2010, DNAm assayed in 2021); and the Multi Ethnic Study of  
17 Atherosclerosis (MESA; biological specimens collected 2010-2012, DNAm assayed in 2012-2013);  
18 using new georeferenced social exposure data for both studies (generated in 2022).

19 **Setting:** MBMS was recruited from four community health centers in Boston; MESA was recruited  
20 from four field sites in: Baltimore, MD; Forsyth County, NC; New York City, NY; and St. Paul, MN.

21 **Participants:** Two population-based samples of US-born Black non-Hispanic (Black NH), white non-  
22 Hispanic (white NH), and Hispanic individuals (MBMS; n=224 Black NH and 69 white NH) and (MESA;  
23 n=229 Black NH, n=555 white NH and n=191 Hispanic).

24 **Exposures:** Eight social exposures encompassing racial discrimination, economic adversity, and air  
25 pollution.

26 **Main outcome:** Genome-wide changes in DNAm, as measured using the Illumina EPIC BeadChip  
27 (MBMS; using frozen blood spots) and Illumina 450k BeadChip (MESA; using purified monocytes).  
28 Our hypothesis was formulated after data collection.

29 **Results:** We observed the strongest associations with traffic-related air pollution (measured via  
30 black carbon and nitrogen oxides exposure), with evidence from both studies suggesting that air  
31 pollution exposure may induce epigenetic changes related to inflammatory processes. We also  
32 found suggestive associations of DNAm variation with measures of structural racial discrimination  
33 (e.g., for Black NH participants, born in a Jim Crow state; adult exposure to racialized economic  
34 residential segregation) situated in genes with plausible links to effects on health.

35 **Conclusions and Relevance:** Overall, this work suggests that DNAm is a biological mechanism  
36 through which structural racism and air pollution become embodied and may lead to health  
37 inequities.

## 38 Introduction

39 Recent advances enabling large population-based epigenetic studies are permitting researchers to  
40 test hypotheses linking socially-patterned exposures, gene regulation, and health inequities<sup>1-3</sup>. DNA  
41 methylation (DNAm) is a plausible biological mechanism by which adverse social exposures may  
42 become embodied<sup>4,5</sup>, because 1) it plays an active role in genome regulation<sup>6-8</sup>, 2) it changes in  
43 response to environmental exposures<sup>1,9</sup> and internal human physiology like ageing<sup>2</sup> and  
44 inflammation<sup>10</sup>, and 3) induced changes can be long-lasting<sup>11-14</sup>. There is a growing literature  
45 reporting associations between DNAm and environmental factors to which social groups are  
46 unequally exposed; a recent review<sup>15</sup> found associations between DNAm and measures of socio-  
47 economic position (SEP), including income, education, occupation, and neighbourhood measures;  
48 and illustrated timing and duration of exposure is important. Exposure to toxins, including air  
49 pollution, is often inequitable between social groups<sup>16,17</sup>. A number of EWAS have identified  
50 associations with particulate matter<sup>18-21</sup> and oxides of nitrogen (NOx)<sup>21,22</sup>; although there is little  
51 replication between studies, and some studies have failed to find effects of particulate matter<sup>22,23</sup>,  
52 NOx<sup>23</sup>, and residential proximity to roadways<sup>24</sup>. Two EWAS have each found two (non-overlapping)  
53 DNAm sites associated with experience of racial discrimination, one in first generation Ghanaian  
54 migrants living in Europe<sup>3</sup>, and one in African American women<sup>25</sup>; given population and migration  
55 differences the lack of replication is perhaps not surprising.

56 However, no EWAS has yet examined associations between DNAm and exposure to racial  
57 discrimination and economic adversity, both at individual and structural levels, and measured at  
58 different points in the lifecourse, in the same group of people. This is important because it is not  
59 clear if the different timing, duration, and levels of these adverse exposures are embodied in  
60 different ways involving differing biological pathways. Supporting attention to these issues is a  
61 growing body of research documenting how exposure to health-affecting factors such as toxins,  
62 quality healthcare, education, fresh food, and green spaces are determined by the way dominant

63 social groups have structured society, which in turn results in health inequities between dominant  
64 social groups and groups they have minoritized <sup>4,26</sup>. Structural racism (the totality of ways in which  
65 society discriminates against racialized groups <sup>27</sup>), for example, results in people of colour often  
66 disproportionately bearing the burden of adverse exposures and economic hardship <sup>4,28</sup>, thus driving  
67 racialized health inequities <sup>29</sup>. Associations between structural racism and cardiovascular health have  
68 been shown for discriminatory housing policies and continuing neighbourhood racial segregation <sup>30,31</sup>;  
69 with the historical legacy of slavery <sup>32</sup>; and with state-level institutional domains <sup>33</sup>. Associations  
70 have also been shown for diabetes outcomes in the US <sup>34</sup> and globally <sup>35</sup>.

71 Guided by the ecosocial theory of disease distribution <sup>4,5</sup>, we tested the hypothesis that DNAm is a  
72 biological mechanism by which embodiment of structural racial discrimination, economic hardship,  
73 and air pollution may occur. We tested our study hypothesis using data from US-born participants in  
74 two US population based studies with similar exposure data: our primary study, the My Body My  
75 Story study (MBMS), and the Multi-Ethnic Study of Atherosclerosis study (MESA), which we use for  
76 evidence triangulation <sup>36</sup> due to differences between the two study populations.

## 77 Methods

### 78 Participants

79 This study utilises biological specimens obtained in 2010-2012 from MBMS and MESA, two US  
80 population-based studies that contain similar data on the study exposures. In 2021-2022, the study  
81 team newly conducted epigenetic assays for MBMS and added new georeferenced social exposure  
82 data. Full study descriptions are in the [supplementary materials](#). Our analyses comprised 293  
83 participants (224 Black and 69 white) from MBMS; and 975 participants from MESA who were US-  
84 born (229 Black, 555 white NH, additionally including 191 Hispanic).

### 85 Social exposures

86 We tested the relationship between DNAm and eight variables relating to exposure to racial  
87 discrimination (both structural and self-reported), economic hardship, and air pollution; these are  
88 described in detail in [Supplementary Table 2](#).

### 89 DNA methylation

90 For detailed description of DNA extraction and DNAm data generation, please see the  
91 [Supplementary materials](#). Briefly, for MBMS DNA was extracted from frozen blood spots in 2021,  
92 and data were generated using the Illumina Infinium MethylationEPIC Beadchip. For MESA, DNA was  
93 extracted from purified monocytes in 2012-2013 and data were generated using the Illumina  
94 Infinium HumanMethylation450 BeadChip. We used DNAm beta values for both studies, which  
95 measure DNAm on a scale of 0 (0% methylation) to 1 (100% methylation).

### 96 Participant stratification

97 EWAS were stratified by self-reported membership of racialized groups, for two reasons. Firstly, for  
98 most of our exposures, different constructs are represented between the racialized groups; for  
99 example, being born in a Jim Crow state means something very different for individuals who identify  
100 as Black versus white. Secondly, stratification prevents potential confounding by racialized group

101 due to exposure and a degree of genetic differences between groups. Racialized groups are social  
102 constructs that are changeable and dependent on local context <sup>37</sup>; they are important to our  
103 research question because group membership is pertinent to the experience of social inequities  
104 perpetuated by structural racial discrimination.

## 105 EWAS

106 All EWAS were conducted using linear regression models implemented using the R package *meffil* <sup>38</sup>.  
107 Many exposures had low levels of missing data, complete case numbers for each EWAS can be found  
108 in Table 3. EWAS details can be found in the [Supplementary materials](#); briefly, we adjusted for age,  
109 reported gender (MBMS)/sex (MESA), smoking status, blood cell count proportions, and batch  
110 effects.

## 111 Sensitivity analysis

112 In MESA we conducted a sensitivity analysis to test whether our results were influenced by  
113 population stratification; details are in the [Supplementary materials](#). Additional sensitivity analysis  
114 restricted the MESA analysis to participants recruited from the Baltimore and New York sites,  
115 because these cities bear the greatest similarity to the Boston area in terms of geographical location,  
116 city environment, and social histories.

## 117 Meta-analysis

118 We meta-analysed associations with air pollution within MBMS and within MESA because air  
119 pollution is the only exposure we tested that we would hypothesize to have the same meaning, and  
120 therefore biological effect, for all individuals. We used *METAL* <sup>39</sup> to meta-analyse effect sizes and  
121 standard errors of the EWAS summary statistics of each racialized group, for black carbon/LAC and  
122 NO<sub>x</sub>.

## 123 Functional relevance of sites passing the genome-wide threshold

124 For DNAm sites associated with an exposure, we used the UCSC genome browser to identify  
125 genomic regions. For sites within known genes, we used GeneCards (<https://www.genecards.org/>)  
126 and literature searches to identify putative gene functions. We used the EWAS Catalog to determine  
127 if associations between DNAm sites and other traits had been reported in previous studies. Where  
128 multiple DNAm sites were associated with an exposure we performed gene set enrichment analysis  
129 using the *missMethyl* R package<sup>40</sup>.

## 130 Biological enrichments of top sites

131 Following each EWAS we performed analyses to ascertain whether DNAm sites associated with our  
132 exposures indicate effects on particular biological pathways, processes or functions. Details are in  
133 the [Supplementary materials](#); briefly, we conducted gene set enrichment analyses, and for  
134 enrichments of tissue-specific chromatin states, genomic regions and transcription factor binding  
135 sites (TFBS).

## 136 Lookup of associations in *a priori* specified genomic locations

137 We hypothesised *a priori* that our EWAS would detect DNAm sites that have been robustly  
138 associated with our study exposures, or factors that might relate to our exposures, in previous  
139 studies. See [Supplementary materials](#) for details.

# 140 Results

## 141 Participant characteristics

142 Both cohorts include racialized groups that are underrepresented in epigenetic studies. Beyond this,  
143 substantial differences existed between the racialized groups within and across MBMS and MESA.  
144 Overall, MBMS participants were on average 21 years younger than MESA participants, had less  
145 variability in exposure to air pollution, and far more were current smokers. In both studies, Black NH  
146 compared to white NH participants had higher BMI, rates of smoking, impoverishment, lower



147 education, rates of self-reported exposure to racial discrimination, and were more likely to be born  
148 in a Jim Crow state and live in a neighbourhood with extreme concentrations of low-income persons  
149 of colour. In MESA, Hispanic participants reported the lowest levels of personal and parental  
150 education.

Variable	MBMS: Black NH	MBMS: white NH	MESA: Black NH	MESA: white NH	MESA: Hispanic	
<b>Total N</b>	<b>224</b>	<b>69</b>	<b>229</b>	<b>555</b>	<b>191</b>	
<b>Sociodemographic characteristics</b>						
Age: mean (SD)	49.02 (7.8)	48.7 (8.3)	71 (8.9)	70.1 (9.5)	68.5 (8.9)	
Gender: N (%) women	135 (60.3%)	49 (71%)	133 (58.1%)	264 (47.6%)	86 (45%)	
BMI: mean (SD)	32.1 (7.7)	29.7 (7.2)	30.6 (5.7)	28.7 (5.3)	30.8 (5.5)	
Smoking: N (%)	Current	115 (51.3%)	24 (34.8%)	31 (13.7%)	44 (8%)	16 (8.6%)
	Former	31 (13.8%)	23 (33.3%)	101 (44.9%)	262 (47.7%)	80 (43%)
	Never	78 (34.8%)	22 (31.9%)	93 (41.5%)	243 (44.3%)	90 (48.4%)
	<i>Missing</i>	0	0	4 (1.7%)	6 (1.1%)	5 (2.6%)
<b>Childhood exposure to racialized and economic adversity:</b>						
Born in a Jim Crow state <sup>1</sup> : N (%) yes	71 (31.7%)	2 (3%)	165 (72.1%)	166 (29.9%)	19 (9.9%)	
Parent's highest education: N (%)	<High school	29 (18.4%)	8 (14%)	95 (42.2%)	161 (29.3%)	129 (69.7%)
	>= High school and <4yr college	94 (59.5%)	24 (42.1%)	106 (47.3%)	258 (47%)	51 (27.6%)
	4+ years college	35 (22.2%)	25 (43.9%)	24 (10.7%)	130 (23.7%)	5 (2.7%)
	<i>Missing</i>	66 (29.5%)	12 (17.4%)	4 (1.7%)	6 (1.1%)	6 (3.1%)
Participant's education: N (%)	<High school	34 (15.2%)	8 (11.6%)	23 (10%)	21 (3.8%)	35 (18.3%)
	>= High school and <4yr college	161 (71.9%)	33 (47.8%)	175 (76.4%)	413 (74.4%)	140 (73.3%)
	4+ years college	29 (12.9%)	28 (40.6%)	31 (13.5%)	121 (21.8%)	16 (8.4%)
	<i>Missing:</i>	0	0	0	0	0
<b>Adult exposure to racialized and economic adversity:</b>						
Household income to poverty ratio <sup>2</sup> : mean (SD)	2.2 (2.2)	2.9 (2.3)	3.9 (2.3)	4.8 (2.9)	3.3 (2.1)	
	<i>Missing</i>	34 (15.2%)	3 (4.3%)	9 (3.9%)	24 (4.3%)	7 (3.7%)
Index of Concentration at the Extremes for racialized economic segregation <sup>3</sup> : mean(SD)	-0.07 (0.2)	0.19 (0.2)	-0.11 (0.2)	0.16 (0.2)	0.09 (0.2)	
	<i>Missing</i>	0	0	2 (0.9%)	4 (0.7%)	12 (6.3%)
Black carbon (µg/m <sup>3</sup> ): mean (SD)	0.64 (0.1)	0.63 (0.17)				
	<i>Missing</i>	0	0			
Light absorption coefficient (10 <sup>-5</sup> /m): mean (SD)			0.89 (0.35)	0.6 (0.3)	0.7 (0.4)	
	<i>Missing</i>		14 (6.1%)	23 (4.1%)	11 (5.6%)	
Pollution Proximity Index <sup>4</sup> (scale of 0-5): mean (SD)	4.3 (1.1)	3.9 (1.4)				

	<i>Missing</i>	5 (2.2%)	0			
Oxides of nitrogen (NO <sub>x</sub> , parts per billion): mean (SD)				31.9 (16.2)	21.55 (12.2)	27 (16.4)
	<i>Missing</i>			14 (6.1%)	23 (4.1%)	11 (5.6%)
Experiences of Discrimination (EOD, N of domains) <sup>5</sup> : N (%)	0	30 (13.4%)	35 (50.7%)			
	1-2	52 (23.2%)	24 (34.8%)			
	3+	140 (62.5%)	10 (14.5%)			
	<i>Missing</i>	2 (0.9%)	0			
Major Discrimination Scale (MDS, N of domains) <sup>6</sup> : N (%)	0			129 (56.6%)	534 (96.4%)	131 (68.6%)
	1-2			79 (34.6%)	20 (3.6%)	53 (27.7%)
	3+			20 (8.8%)	0	7 (3.7%)
	<i>Missing</i>			1 (0.4%)	1 (0.2%)	0
<b>Predicted cell count proportions</b>						
B Cell		0.08 (0.02)	0.06 (0.01)	0.04 (0.03)	0.03 (0.02)	0.03 (0.02)
CD4+T cells		0.17 (0.05)	0.15 (0.04)	0.04 (0.02)	0.03 (0.03)	0.03 (0.01)
CD8+T cells		0.005 (0.02)	0.002 (0.01)	0.002 (0.006)	0.0007 (0.003)	0.0006 (0.003)
Monocytes		0.124 (0.02)	0.116 (0.02)	0.9 (0.05)	0.91 (0.04)	0.92 (0.04)
Neutrophils		0.55 (0.1)	0.62 (0.08)	0.0003 (0.002)	0.0008 (0.005)	0.0009 (0.007)
Natural Killer		0.1 (0.04)	0.09 (0.04)	0.015 (0.01)	0.012 (0.01)	0.01 (0.01)
Eosinophils		0.007 (0.02)	0.003 (0.009)	0.01 (0.01)	0.01 (0.01)	0.01 (0.01)

151 *Table 1: Characteristics of MBMS and MESA participants.*

152 <sup>1</sup> *Jim Crow states are the 21 US states (plus the District of Columbia) which permitted legal racial discrimination prior to the*  
153 *1964 US Civil Rights Act.*

154 <sup>2</sup> *Participants' ratio of household income in 2010 dollars to the US 2010 poverty line given household composition.*

155 <sup>3</sup> *Census tract measure of economic and racialized segregation, scored from -1 to 1*

156 <sup>4</sup> *NO<sub>x</sub> measurements were used to construct a weighted score of roadway pollution*

157 <sup>5</sup> *Validated self-report questionnaire measuring the number of domains of exposure to racial discrimination. Score range 0-*  
158 *9, categorised into 0, 1-2, 3*

159 <sup>6</sup> *Validated self-report questionnaire measuring the number of domains of exposure to racial discrimination; combined with*  
160 *the attribution aspect from EDS (everyday discrimination scale) to enable comparability between EOD and MDS. Score*  
161 *range 0-5, categorised into 0, 1-2, 3+*

162

## 163 EWAS results and biological interpretation

164 In MBMS, among the Black NH participants one DNAm site, in ZNF286B, was associated with being  
165 born in a Jim Crow state. Another DNAm site, PLXND1, was associated with participants having less  
166 than high school education. Among white NH participants, no associations passed the genome-wide  
167 threshold. See Table 2 for details of gene functions; Table 3 for numbers of associated EWAS sites;  
168 and [Supplementary figures 1 and 2](#) for Miami plots. In MESA, two DNAm sites were associated with  
169 racialized economic segregation – one in Black NH participants (in FUT6) and one in white NH  
170 participants (a CpG previously associated with BMI); and in Black NH participants one DNAm site (in  
171 PDE4D) was associated with an MDS score of 0. The majority of associations in MESA were related to  
172 air pollution exposure – among Black NH participants, 12 sites with LAC and 22 sites with NO<sub>x</sub>.  
173 Notably, many of these sites are clustered in genes with putative roles in immune responses and are  
174 known to interact with one another, including KLF6, MIR23A, FOS, FOSB, ZFP36 and DUSP1. Among  
175 the MESA white NH participants, four DNAm sites were associated with both LAC and NO<sub>x</sub>, and an  
176 additional 3 uniquely associated with LAC. Associations of 53 DNAm sites with birth in a Jim Crow  
177 state were the result of confounding by air pollution (see [Supplementary Materials](#)). Among Hispanic  
178 participants, one site was associated with LAC (NPNT) and one with NO<sub>x</sub> (ADPRHL1). See Table 3 for  
179 numbers of associations for all EWAS performed and [Supplementary Figures 3-5](#) for corresponding  
180 Miami plots.

Gene	Chr	Functional relevance	MBMS		MESA		
			Black NH	White NH	Black NH	White NH	Hispanic
ZNF286B	17	A pseudogene, which is predicted to be involved in regulation of RNA polymerase 2 (Pol II)-mediated transcription (Pol II transcribes protein-coding genes into mRNA <sup>41</sup> ).	Born in a Jim Crow state: 1				
PLXND1	3	Encodes a cell receptor involved in axonal guidance, migration of endothelial cells, and regulates atherosclerotic plaque deposition <sup>42</sup> .	<HS education: 1				
FUT6	19	A Golgi stack membrane protein that is involved in basophil-mediated allergic inflammation <sup>43</sup> .			residential racialized economic segregation: 1		
KLF6	10	A transcriptional activator and tumour suppressor, which regulates macrophage inflammatory responses <sup>44</sup> .			LAC: 2 NOx: 3	LAC: 1 NOx: 1	
FOS	14	FOS is an early-response gene, and is a subunit of the AP-1 transcription factor complex, which regulates gene expression involved in lung injury, repair and transformation <sup>45</sup> , as well as regulating many cytokine genes and T-cell differentiation <sup>46,47</sup> .			LAC: 1 NOx: 7		
FOSB	19	FOSB is another subunit of AP-1.			LAC: 1 NOx: 1		
ZFP36	19	ZFP36 encodes a protein (TTP) that is a key regulator of post-transcriptional regulation, which has roles in immune and inflammatory responses <sup>48</sup> .			NOx: 2		
DUSP1	5	DUSP1 is a gene that regulates airway inflammation; DUSP1's key mechanism of inflammation modulation may be via modulating the actions of the protein TTP encoded by the ZFP36 gene <sup>49</sup> .			LAC: 1 NOx: 1		
VIM	10	Encodes a filament protein responsible for integrity of cell shape and cytoplasm. Pathogens can attach to this protein on the cell surface. Putative involvement regulating innate immune response to lung injury and irritation <sup>50</sup>			NOx: 1		
PDE4D	5	PDE4s, including PDE4D, have roles in cell signalling, as well as			MDS: 1		

		regulating inflammatory responses <sup>51</sup> .					
MALAT1	11	Metastasis associated lung adenocarcinoma transcript 1, lncRNA that acts as transcriptional regulator; upregulation linked to cancerous tissues and proliferation and metastasis of tumour cells				LAC: 1	
CYTIP	2	Modulates activation of ARF (ADP-ribosylation factor) genes, which regulate vesicle budding, tethering and cytoskeleton organization. Dysregulation of ARFs may be involved in cancer cell migration and invasion.				LAC: 1 NOx: 1	
ZEB2	2	DNA-binding transcriptional repressor involved in the transforming growth factor- $\beta$ (TGF- $\beta$ ) signalling pathway that interacts with activated SMADs. May be related to small cell lung cancer <sup>52</sup> .				LAC: 1 NOx: 1	
PTPRC	1	A receptor-type PTP that is an essential regulator of T- and B-cell antigen receptor signalling.				LAC: 1 NOx: 1	
NPNT	4	An extracellular matrix protein that has roles in kidney development and carcinogenesis <sup>53</sup> .					LAC: 1
ADPRHL1	13	a protein encoding a pseudoenzyme involved in cardiogenesis <sup>54</sup> .					NOx: 1

Table 2: Putative functions of genes in which the top exposure-associated DNAm sites sit; with details of how many sites within that gene were identified, and in which main analysis EWAS they were identified.

181  
182

	MBMS				MESA					
	Black NH		white NH		Black NH		white NH		Hispanic	
	N	N sites	N	N sites	N	N sites	N	N sites	N	N sites
Birth in a Jim Crow state	224	1	NA <sup>1</sup>	NA <sup>1</sup>	225	0	549	53 <sup>2</sup>	186	0
Parent's highest education (high vs low)	64	0	33	0	117	0	288	0	NA <sup>3</sup>	NA <sup>3</sup>
Parent's highest education (high vs mid)	129	0	49	0	128	0	384	0	NA <sup>3</sup>	NA <sup>3</sup>
Participant's education (high vs low)	63	1	36	0	54	0	142	0	51	0
Participant's education (high vs mid)	190	0	61	0	202	0	534	0	156	0
Household poverty to income ratio	190	0	66	0	218	0	528	0	180	0
Racialized economic segregation	224	0	69	0	223	1	545	1	174	0
Black carbon	224	0	69	0	211	12	526	7	175	1
Nitrogen oxides	219	0	69	0	211	22	526	4	175	1
EOD <sup>5</sup> (1-2 vs 0)	82	0	59	0						
EOD <sup>5</sup> (1-2 vs 3+)	192	0	34	0						
MDS <sup>6</sup> (1-2 vs 0)					204	1	554	0	184	0
MDS <sup>6</sup> (1-2 vs 3+)					97	0	NA <sup>4</sup>	NA <sup>4</sup>	60	0

183 Table 3: Summary of the number of DNAm sites passing the genome-wide threshold in each individual EWAS in MBMS (threshold  $2.4e-7$ ) and MESA (threshold  $9e-8$ ). The list of specific DNAm  
 184 sites passing the genome-wide threshold can be found in [supplementary table 4](#). <sup>1</sup> The EWAS was not run for Jim Crow birth state for white NH participants in MBMS, due to small cell numbers.  
 185 <sup>2</sup> See text; these 53 sites were driven by air pollution differences between individuals born and not born in a Jim Crow state. <sup>3</sup> The two EWAS for parental education were not run for Hispanic  
 186 participants in MESA, due to small cell numbers. <sup>4</sup> The EWAS was not run for MDS (score of 1-2 vs 3+) for white NH participants in MESA, as no participants had a score of 3 or more. <sup>5</sup> EOD –  
 187 Experiences of Discrimination scale. <sup>6</sup> MDS – Major Discrimination Scale

## 188 MESA subgroup analysis

189 The main impact of removing the Minnesota and Forsyth County sites (which both had very low  
190 levels of air pollution) was to remove the confounding structure between air pollution and Jim Crow  
191 birth state among white NH participants. It also increased the similarity of air pollution associations  
192 between the Black NH and white NH participants; for example, of the 19 DNAm sites associated with  
193 NOx among white NH participants, 12 passed the genome-wide threshold in the Black NH participant  
194 EWAS. Numbers of associated sites are in Table 4. Miami plots for this MESA subgroup can be found  
195 in [Supplementary figures 6, 7 and 8](#).

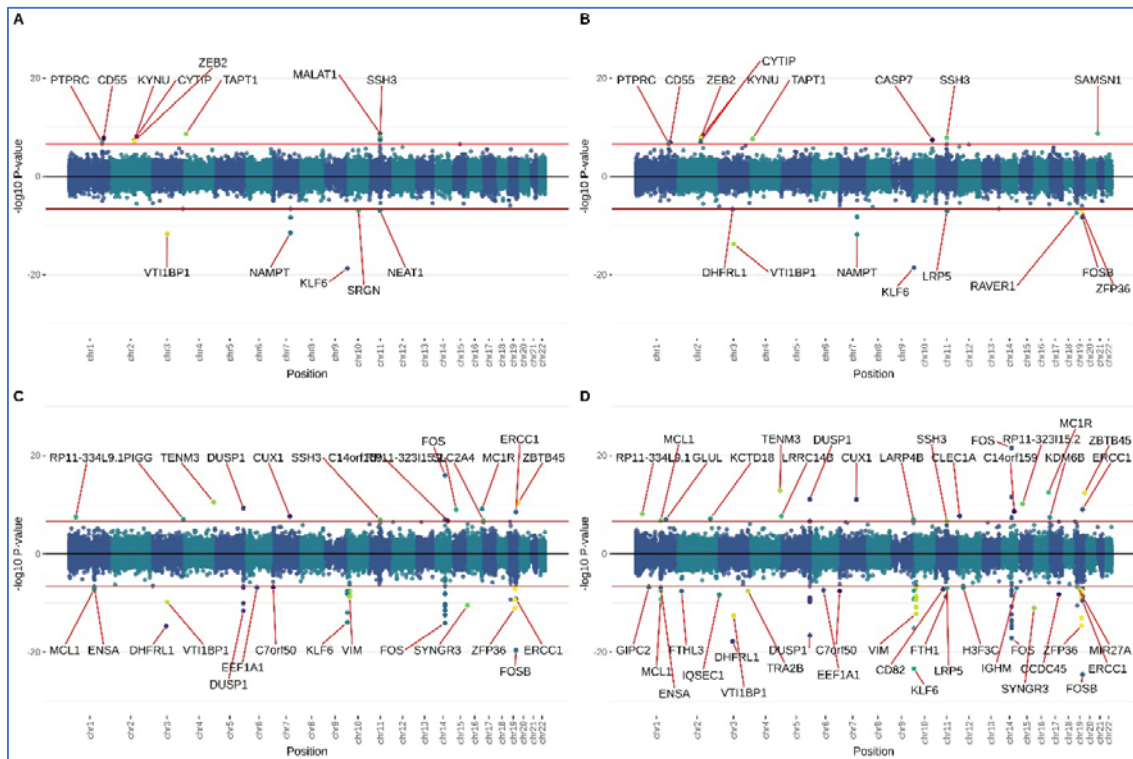
	Black NH		white NH		Hispanic	
	N	N sites	N	N sites	N	N sites
<b>Birth in a Jim Crow state</b>	221	0	237	0	NA <sup>1</sup>	NA <sup>1</sup>
<b>Parent's highest education (high vs low)</b>	115	0	134	0	NA <sup>2</sup>	NA <sup>2</sup>
<b>Parent's highest education (high vs mid)</b>	125	0	164	0	NA <sup>2</sup>	NA <sup>2</sup>
<b>Participant's education (high vs low)</b>	54	0	55	0	NA <sup>2</sup>	NA <sup>2</sup>
<b>Participant's education (high vs mid)</b>	198	0	227	0	NA <sup>2</sup>	NA <sup>2</sup>
<b>Household poverty:income ratio</b>	214	0	227	1	67	0
<b>Racialized economic segregation</b>	219	1	233	0	59	0
<b>Light Absorption Coefficient</b>	208	10	231	6	57	0
<b>Nitrogen oxides</b>	208	20	231	19	57	0
<b>Major Discrimination Scale (1-2 vs 0)</b>	200	1	236	0	67	0
<b>Major Discrimination Scale (1-2 vs 3+)</b>	94	0	NA <sup>3</sup>	NA <sup>3</sup>	NA <sup>3</sup>	NA <sup>3</sup>

196 *Table 4: Summary of EWAS results for MESA subgroup analysis.* <sup>1</sup> The EWAS for Jim Crow birth state was not run for  
197 Hispanic participants due to small cell numbers. <sup>2</sup> The EWAS for parental and participant education were not run for  
198 Hispanic participants, due to small cell numbers. <sup>3</sup> The EWAS was not run for MDS (score of 1-2 vs 3+) for white NH and  
199 Hispanic participants in MESA, as no participants had a score of 3 or more.

## 200 Meta-analysis

201 Meta-analysis in MBMS did not yield any sites passing the genome-wide threshold. In MESA we see  
202 approximately similar numbers of associations as with the Black NH subgroup (17 for LAC and 18 for  
203 NOx); see Supplementary Table 3. When we restricted to participants recruited at the Baltimore and  
204 New York sites, a much larger number of DNAm sites passed the genome-wide threshold (51 for LAC  
205 and 79 for NOx); this may be because Minnesota and Forsyth County sites had very low variance in  
206 pollution levels. The MESA sensitivity meta-analysis identified multiple associations linked to DUSP1,  
207 FOS, KLF6, MCL1, and VIM; genes that have putative roles in inflammation and immunity.





208

209 *Figure 1: MESA air pollution meta-analysis Manhattan plots. A: MESA full cohort LAC meta-analysis. B: MESA full cohort NOx*  
 210 *meta-analysis. C: MESA subgroup LAC meta-analysis. D: MESA subgroup NOx meta-analysis.*

## 211 Biological enrichments of exposure associations

### 212 Gene ontology

213 We observed no evidence for gene set enrichments for any Gene Ontology terms among the top 100  
 214 sites of the main EWAS we conducted. However, we did observe that the 22 sites associated with  
 215 NOx above the genome-wide threshold among MESA Black NH participants were enriched for the  
 216 gene ontology terms ‘response to glucocorticoid’ and ‘response to corticosteroid’ (FDR>0.05). We  
 217 also observed that the MESA meta-analysis of NOx among all participants was associated with 13  
 218 Gene Ontology terms (FDR>0.05) related to blood-based immune response.

### 219 EWAS catalog

220 We observed a number of relevant enrichments among sites identified in our EWAS. Details of the  
 221 associations ( $p < 0.05$ , Fisher’s exact test) can be found in the Supplementary Materials and  
 222 **Supplementary figures 6-16**. Briefly, in MBMS, we see enrichment for inflammation for both NOx  
 223 and LAC EWAS among Black NH participants, and in the NOx meta-analysis. In MESA, we observed

224 consistent enrichment for infection and cancer among Black NH and white NH participants, and also  
225 in the meta-analyses. We observed enrichment for inflammation among Hispanic participants. We  
226 also found among both MESA Black NH and Hispanic participants, the racialized economic  
227 segregation EWAS was enriched for neurological traits. Among both the white NH and Hispanic  
228 participants, household poverty to income ratio EWAS was enriched for SEP and education. In the  
229 MESA subgroup analysis, enrichment for prenatal exposures was observed for the Jim Crow birth  
230 state EWAS among the Black NH and Hispanic participants.

### 231 **Enrichment for genomic features**

232 When we looked at enrichment of genomic locations of the top 100 sites ( $p < 0.05$ , Fisher's exact  
233 test), we found that among MBMS Black NH participants, NO<sub>x</sub> was the only exposure with  
234 associated CpGs being located in active genomic regions (please see [Supplementary materials](#) for  
235 details). In the MBMS meta-analyses, NO<sub>x</sub> was enriched for regions related to gene promoters.  
236 Among MESA Black NH participants, we observe enrichment for regions related to transcription and  
237 genome regulation in the LAC and NO<sub>x</sub> EWAS. We also observed enrichment relating to transcription  
238 regulation for the birth in a Jim Crow state EWAS. Among MESA white NH participants, we observed  
239 enrichment for transcription regulation for both measures of air pollution. Among MESA Hispanic  
240 participants, LAC exposure shows some associations with active genomic regions. When we restrict  
241 MESA to the New York and Baltimore sites, we see a similar set of enrichments; and in the MESA  
242 meta-analyses we see consistent enrichment related to transcription regulation and promoters.  
243 Notably, genomic feature enrichments for NO<sub>x</sub> among both MBMS and MESA Black NH participants  
244 involved similar genomic locations (CpG islands and shores) and chromatin states (related to  
245 promoters), as well as 6 of a possible 9 TFBS.

### 246 **Lookup of associations in a priori specified genomic locations**

247 We did not observe any associations in our EWAS results for sites identified in previous EWAS of  
248 related exposures.

## 249 Discussion

250 The series of EWAS we conducted on a range of adverse exposures at different levels and at  
251 different points in the lifecourse, drawing on two different population-based studies with similar  
252 exposure data, provide evidence that DNAm may be a biological pathway by which societal context  
253 shapes health inequities. This work has shown for the first time associations between DNAm and  
254 multiple levels of structural discrimination, in genes that are biologically plausible routes of  
255 embodiment involving gene regulation, including inflammation. Additionally, our EWAS and meta-  
256 analyses of air pollution showed clear association between two road traffic-related measures of air  
257 pollution, and DNAm of multiple CpGs in multiple genes that have been consistently associated with  
258 inflammation and infection, suggesting that the material environment people live may induce  
259 inflammatory changes. Our study has added to the existing literature on air pollution; there are few  
260 EWAS studies looking at NO<sub>x</sub> (n=3), and none so far looking at black carbon. In total, this work  
261 highlights the need for researchers to consider multiple levels of discrimination and adversity across  
262 the lifecourse, especially structural inequities in the material world in which people live, to fully  
263 elucidate drivers and biological mechanisms of inequitable health.

264 Associations detected at the genome-wide level in MBMS related more closely to early-life  
265 exposures (being born in a Jim Crow state, and low educational attainment); in MESA they related  
266 more to current experiences and exposures (air pollution, racialized economic segregation, and  
267 experiences of discrimination), possibly reflecting the relatively older age of the MESA participants.  
268 The much stronger associations with air pollution in MESA compared to MBMS could potentially be  
269 due to: (1) the use of purified monocytes in MESA, with a single cell type making associations easier  
270 to detect; (2) less variation in exposure to air pollution in MBMS compared MESA; (3) longer  
271 duration of air pollution exposure in MESA (due to older age of the participants); or (4) reduced  
272 statistical power in MBMS, due to lower quantities of DNA.

273 Notably, inflammation was the predominant pathway indicated in the air pollution analyses, both via  
274 putative gene functions and enrichment analyses. These findings underscore that while there is a  
275 large psychosocial literature on inflammation being a mechanism by which discrimination harms  
276 health<sup>28,55,56</sup>, it is also critical to consider inequities in biophysical exposures in the material world as  
277 an important driver of this inflammation. Overall, air pollution sites tend to be enriched for  
278 inflammation in MBMS and infection in MESA; this could represent different mechanisms of the  
279 same process due to the different blood cell types sampled in the two cohorts; with monocytes  
280 being specialised in infection prevention, and neutrophils (the highest proportion cell in whole blood)  
281 being specialised in inflammatory responses.

282 Our study identified a greater number of associations with air pollution measures than previous  
283 work in MESA<sup>20,57</sup>; this is likely due to the fact that we do not adjust for recruitment site (which  
284 would reduce variation in the exposure because exposure is location-dependent); and previous  
285 analyses have adjusted for racialized group membership, which is also associated with air pollution  
286 exposure; this may have masked the effects that we have detected. This joins other research that  
287 has demonstrated the importance of considering spatial effects of air pollution<sup>58</sup>.

288 A limitation of our study is that we cannot infer causality. Although it would be possible to conduct  
289 Mendelian randomization instrumenting *cis*-mQTLs, we did not conduct this analysis because we  
290 think the results would be highly speculative. Additionally, the MESA sample we used may have been  
291 subject to selection bias, because (1) individuals who had experienced prior cardiovascular events  
292 were excluded from recruitment, and (2) a number of participants died between Exam 1 and Exam 5.  
293 If adversity and discrimination are associated with these cardiovascular events and mortality,  
294 associations could be biased in MESA.

## 295 Conclusions

296 We think this work provides direction for future epigenetic studies to consider the role of  
297 inequitable adverse social and biophysical exposures across the lifecourse, including but not limited

298 to structural discrimination. Our results suggest inflammation may be a key biological pathway by  
299 which inequities become embodied, in our case driven primarily by exposure to air pollution, and  
300 not self-reported racial discrimination. These findings accordingly suggest that attention to how  
301 social inequities shape biophysical as well as social exposures is crucial for understanding how  
302 societal inequities can become embodied, via pathways involving DNAm.

## 303 Declaration statements

### 304 Ethics approval

305 The study protocol, involving use of both the MBMS and MESA data, was approved by the Harvard  
306 T.H. Chan School of Public Health Office of Human Research Administration (Protocol # IRB19-0524;  
307 June 10, 2019).

308 The original MBMS study protocol, implemented in accordance with the Helsinki Declaration of 1975,  
309 as revised in 2000, was approved by the Harvard School of Public Health Office of Human Research  
310 Administration (protocol #11950–127, which covered 3 of the 4 health centers through reciprocal  
311 IRB agreements), and was also separately approved by the fourth community health center’s  
312 Institutional Review Board. All participants provided written informed consent.

313 Information regarding the MESA protocols and their IRB approvals and other information, is  
314 available at: [www.mesa-nhlbi.org](http://www.mesa-nhlbi.org).

### 315 Data sharing

316 This study ([NIH Grant number R01MD014304](#)) relied on three sources of data, each of which is  
317 subject to distinct data sharing stipulations: (1) the non-public data from the “My Body, My Story”  
318 (MBMS) study; (2) the non-public data from the Multi-Ethnic Study of Atherosclerosis (MESA; data  
319 use agreement G638); and (3) the public de-identified data from the US Census, the American  
320 Community Survey, and the State Policy Liberalism Index. We provide descriptions of these data

321 sharing stipulations and access to these data below; this information is also available at:

322 <https://www.hsph.harvard.edu/nancy-krieger/data-sharing-resources/>

- 323 • ICE metrics relating to racial composition, income distribution, and housing tenure that were  
324 derived from sources in the public domain i.e. the US Census and the American Community  
325 Survey are available at the census tract level now on [GitHub](#).
- 326 • Code used to construct the variables is available on [GitHub](#) and [here](#).
- 327 • The State Policy Liberalism Index data used in our study is also publicly available and can be  
328 obtained from the [Harvard Dataverse](#). Reference: Caughey, Devin; Warshaw, Christopher,  
329 2014, "The Dynamics of State Policy Liberalism, 1936-  
330 2014", <http://dx.doi.org/10.7910/DVN/ZXZMJB> Dataverse [Distributor] V1 [Version].
- 331 • De-identified data from the *My Body My Story* study used for this project will be made  
332 available only for purposes approved by the study PI, as stipulated by the study's informed  
333 consent protocol. The application form to obtain these data will be made available via this  
334 website after completion of this project in late Fall 2024.
- 335 • Data from the [Multi-Ethnic Study of Atherosclerosis \(MESA\)](#) must be obtained directly from  
336 the MESA website via their application protocol.
- 337 • The scripts to run the EWAS and downstream analyses are available on [GitHub](#).
- 338 • EWAS summary statistics **will be uploaded** to the [EWAS catalog](#) website **upon publication**.

### 339 [Author contributions](#)

- 340 • SHW performed quality checks and normalisation of MBMS DNAm data, co-designed and  
341 conducted the analyses, wrote the first manuscript draft, produced tables and figures, and  
342 prepared study materials to be shared via the data repository (software code).
- 343 • MS provided advice on data QC, co-designed the analyses, and contributed to interpreting the  
344 results.

- 345 • CR co-led obtaining funds for the research project, co-designed the analyses, and contributed to  
346 interpreting the results.
- 347 • NK led conceptualization of the study, contributed to designing the analyses, and co-led  
348 obtaining funds for the research project.
- 349 • CT accessed the electronic public use data and generated the study variables derived from these  
350 data, contributed to designing the analyses and interpreting the results, and prepared the study  
351 materials to be shared via the data repository (data dictionary; software code).
- 352 • JTC contributed to designing the analyses and interpreting results.
- 353 • PDW facilitated finalizing all human subject approvals and data use agreements and also the  
354 data transfer of the MBMS epigenetic data from HSPH to Bristol, geocoded the place of birth  
355 data, extracted the historical census data from PDF files, and contributed to interpreting results.
- 356 • AS, BC, KT, and GDS contributed to designing the analyses and interpreting the results.
- 357 • IDV led and supervised the assays to extract epigenetic data from the MBMS blood spots and  
358 contributed to designing the analyses and interpreting the results.
- 359 • ADR facilitated interpretation of the MESA data and contributed to designing the analyses and  
360 interpreting the study results.
- 361 • All co-authors provided critical intellectual content to and approved the submitted manuscript.

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374 Agreement (G638) was approved by MESA (on 11/22/2019) to use pre-existing MESA data.

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## 407 Conflict of interest

408 The authors declare no conflict of interest.

409

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