Supplementary Info

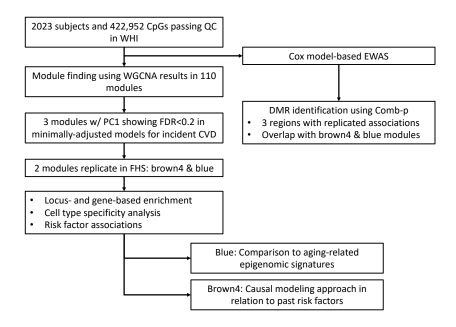


Figure S1: Study overview, including module- and region-based analyses as well as follow-up.

Table S1: P-values for module associations with incident CVD in discovery and replication.

	WHI (disc	covery)	FHS (replication)			
Module	Partially adjusted	Fully adjusted	Partially adj.	Partially adj. (mixed)	Fully adjusted	
blue	0.0002736	0.0500018	0.0000085	0.0000085	0.8189348	
brown4	0.0045462	0.0872688	0.0000962	0.0000963	0.0997390	
lavenderblush3	0.0050028	0.0210976	0.0202819	0.0202804	0.1580861	

¹ Partially-adjusted models are adjusted for technical covariates (DNA pull batch in WHI and study center + 7 control probe PCs in FHS) and estimated cell counts. Fully-adjusted models are additionally adjusted for age, sex, smoking status and smoking pack-years.

² Mixed model contains a random intercept for each family.

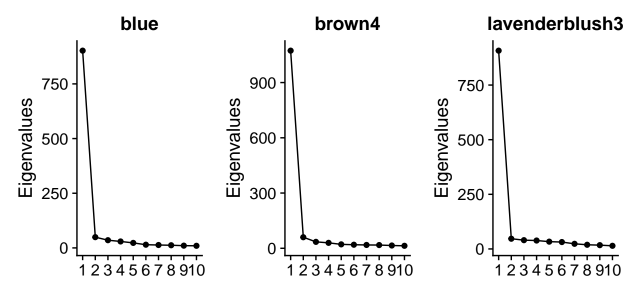


Figure S2: Scree plots for PCA on the set of CpGs corresponding to each of the top modules.

Table S2: CpGs with FDR < 0.05 in the discovery set (Bonferroni threshold = 1.18e-7)

CpG	Chromosome	Dir. of Assoc.	P-value	Location	Annotated Gene	Replication P-value
cg09155044	chr16	+	6.63e-09	TSS1500	VKORC1	0.107
cg24434800	chr1	+	5.04e-08			0.629
cg11691298	chr2	+	1.1e-07	Body	FAM59B	0.525
cg02379107	chr20	+	4.72e-07	TSS1500	KIAA1755	0.930

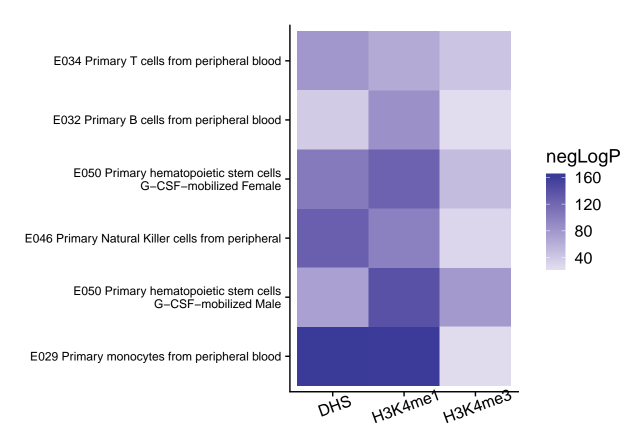


Figure S3: eFORGE cell type-specificity plot for the brown4 module.

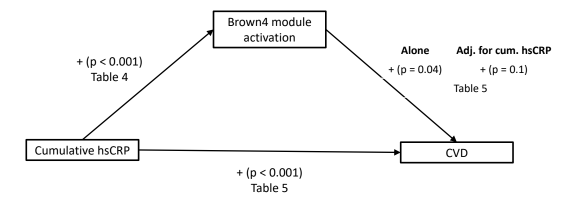


Figure S4: Example diagram of cumulative risk factor mediation by brown4 methylation module activation. Results from 4 regressions are shown: cumulative risk factor exposure to brown4 activation, cumulative risk factor exposure to incident CVD, and brown4 activation to incident CVD with and without adjustment for cumulative risk factor exposure. Regression terms represented as: sign of coefficient (p-value).