An New Epigenetic Clock for Aging and Life Expectancy





Morgan Levine

Yale Center for Research on Aging
Department of Pathology
Yale School of Medicine
Department of Chronic Disease
Epidemiology
Yale School of Public Health









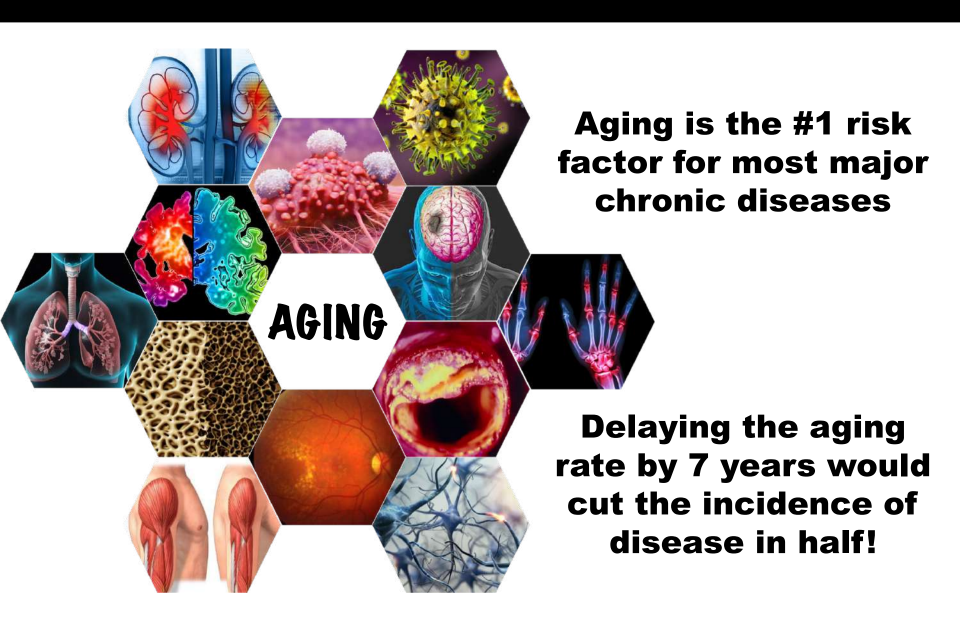








MORE THAN DEATH



AGING HETEROGENEITY

We don't all age in the same way or at the same rate.

Chronological age is an imperfect estimate of the latent concept, "biological aging".

Quantifying "biological age" may:

 Provide an endophenotype from which to identify genetic and environmental contributors to differences in lifespan and healthspan.

2. Facilitate evaluation of interventions aimed at delaying aging.

29

30

31

32

AGING TRAJECTORY

At what level should we estimate "aging?

Proximal to Mechanisms

Proximal to Outcomes

Healthspan (Geroscience Goal)

Demographic Aging

Molecular Alterations



Physiological Dysregulation



Disease/ Disability



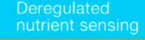
DEATH



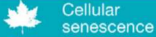






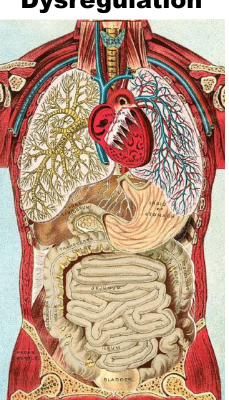






Stem cell exhaustion

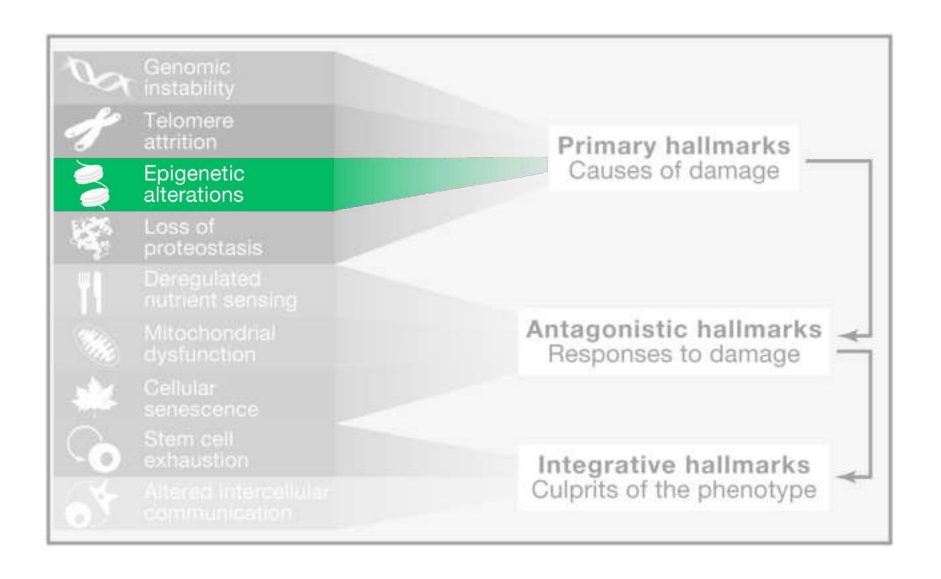
Altered intercellular communication





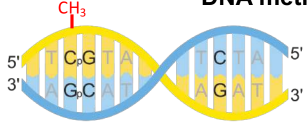


WHAT IS AGING?



EPIGENETIC CLOCKS

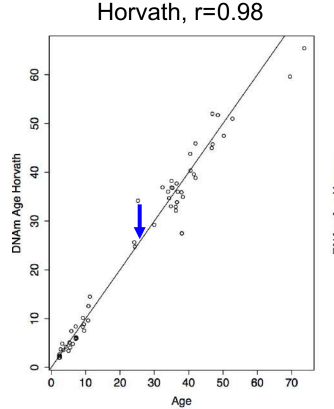
Chronological age has been shown correspond with distinct changes in DNA methylation (DNAm) at specific CpG sites.

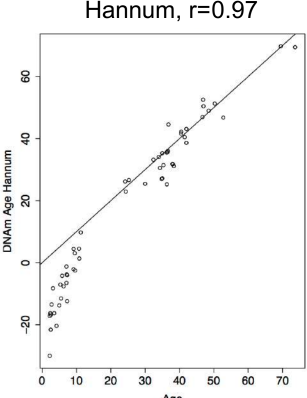


Very accurate epigenetic age predictors have been developed

Instead of minimizing the residual, the goal should be to capture the "true residual".

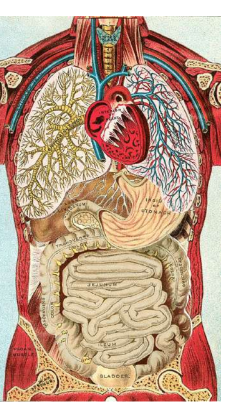
(i.e. decouple chronological time from biological aging)





AIM: Train a clock to predict a variable that already captures differences in physiological dysregulation; susceptibility to disease/disability; and risk of death among same aged individuals.









Develop a multi-system estimate of "Phenotypic Age". Predictor of agingrelated mortality based on clinical measures.

Validate Associations with:

All-Cause Mortality
Cause Specific Mortality
Coexisting Disease Count
Physical Functioning
Mortality Ages 20-64
Mortality Ages 65-79
Mortality Ages 85+



Train a composite epigenetic predictor of phenotypic age, called "DNAm PhenoAge".

Based on DNAm at 513 CpGs.

Validate Associations with:				
All-Cause Mortality	Familial Longevity	Socioeconomic Status		
Coronary Heart Disease Risk	Dementia	Race/ethnicity		
Coexisting Disease Count	Down Syndrome	Diet		
Physical Functioning	Parkinson's Disease	Physical activity		
Disease Free Status	HIV positive	Metabolic Syndrome		
Age at Menopause	Chronological age in 35 tissues/cells	Smoking Status		
Cancer (Lung, Breast)	Neuropathology (Brain DNAm)	Obesity (Liver DNAm)		



Identify <u>underlying</u>
<u>biology</u> of the 513
CpGs in the DNAm
PhenoAge Score

Test for:			
GO Enrichment	Immune Cell Associations		
Pathway Enrichment	CpG Overlap with Hannum/Horvath		
LTL Correlation	Loci-Specific DNAm vs. Transcription		
Differential Expression	Polycomb Group Protein Targets		
CpG Island Enrichment	Chromosomal Locations		
Heritability Analysis	Change in DNAm PhenoAge over Time		
DNAm Network Analysis	Transcriptional Analysis in Monocytes		

Develop a Multisystem Phenotypic Age Estimate and Validate Predictions Develop a New Epigenetic Age Estimate and Validate Predictions/Associations Underlying Biology of the Clock and the 513 CpGs

Training Sample: (N=9,926), Ages 20+, up to 23 years of mortality follow-up

Input Variables: 42 clinical biomarkers and age.

<u>Model:</u> Proportional Hazard Elastic Net (Outcome=Mortality from major agerelated diseases)

Variables

Albumin Creatinine Glucose

C-reactive protein
Lymphocyte percent
Mean cell volume
Red cell distribution width
Alkaline phosphatase
White blood cell count
Age

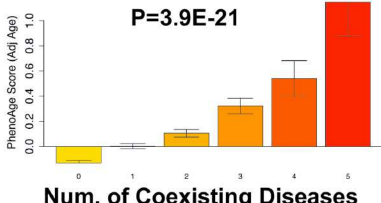
 $Linear\ Prediction = Albumin \times \beta_{Albumin} + CRP \times \beta_{CRP} + \cdots Age \times \beta_{Age} + constant$

Converted to an age (units of years) using parameters from a Gompertz proportional hazard model.

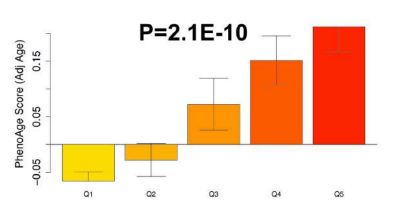
Develop a Multisystem Phenotypic Age **Estimate and Validate Predictions**

Mortality Prediction in Independent Sample

Cause	Cases	HR	P-Value
All-Cause	1052	1.09	3.8E-49
Aging-Related	661	1.09	4.5E-34
CVD	272	1.10	5.1E-17
Cancer	265	1.07	7.9E-10
Alzheimer's	30	1.04	2.6E-01
Diabetes	41	1.20	1.9E-11
Lung	53	1.09	6.3E-04



Num. of Coexisting Diseases



Physical Functioning Quintile

Develop a multi-system estimate of "Phenotypic Age". Predictor of agingrelated mortality based on clinical measures.

Validate Associations with:

All-Cause Mortality
Cause Specific Mortality
Coexisting Disease Count
Physical Functioning
Mortality Ages 20-64
Mortality Ages 65-79
Mortality Ages 85+



Train a composite epigenetic predictor of phenotypic age, called "DNAm PhenoAge".

Based on DNAm at 513 CpGs.

All-Cause Mortality
Coronary Heart Disease Risk
Coexisting Disease Count
Physical Functioning
Disease Free Status
Age at Menopause
Cancer (Lung, Breast)

Validate Associations with: Familial Longevity Dementia Down Syndrome Parkinson's Disease HIV positive Chronological age in 35 tissues/cells Neuropathology (Brain DNAm)

Socioeconomic Status
Race/ethnicity
Diet
Physical activity
Metabolic Syndrome
Smoking Status
Obesity (Liver DNAm)



Identify underlying
biology of the 513
CpGs in the DNAm
PhenoAge Score

for:	Test for:	
	GO Enrichment	
CpG Ove	Pathway Enrichment	
Loci-Spec	LTL Correlation	
Polyc	Differential Expression	
	CpG Island Enrichment	
Change in D	Heritability Analysis	
Transcripti	DNAm Network Analysis	

Immune Cell Associations
CpG Overlap with Hannum/Horvath
Loci-Specific DNAm vs. Transcription
Polycomb Group Protein Targets
Chromosomal Locations
Change in DNAm PhenoAge over Time
Transcriptional Analysis in Monocytes

Develop a Multisystem Phenotypic Age Estimate and Validate Predictions Develop a New Epigenetic Age Estimate and Validate Predictions/Associations

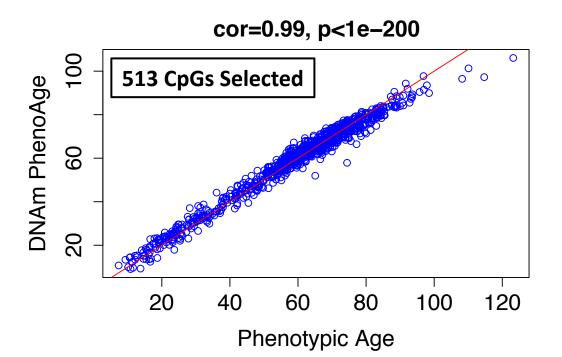
Underlying Biology of the Clock and the 513 CpGs

<u>Training Sample:</u> InCHIANTI—N=456 at two time-points (1998 & 2007).

<u>Input Variables:</u> DNAm from whole blood for about 20,000 CpGs (those on the 27k, 450k, and EPIC chips)

Model: Elastic Net (Outcome=Phenotypic Age)

$$DNAmPhenoAge = CpG1 \times \beta_{CpG1} + \cdots + CpG513 \times \beta_{CpG513} + constant$$



Develop a Multisystem Phenotypic Age
Estimate and Validate Predictions

Develop a New Epigenetic Age Estimate and Validate Predictions/Associations

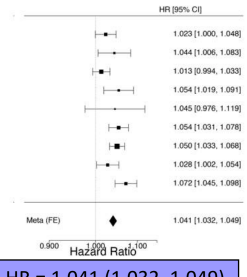
Underlying Biology of the Clock and the 513 CpGs

Levine

1.033 [1.016, 1.050]
1.044 [1.014, 1.075]
1.026 [1.010, 1.043]
1.049 [1.024, 1.075]
•— 1.078 [1.029, 1.129]
1.050 [1.033, 1.068]
1.052 [1.040, 1.065]
1.031 [1.012, 1.050]
1.062 [1.045, 1.080]
1.045 [1.039, 1.051]

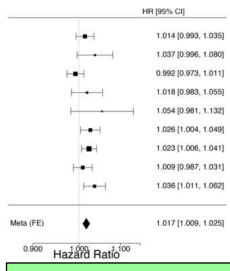
HR = 1.045 (1.039, 1.051) *Meta-p* = 7.9E-47

Hannum



HR = 1.041 (1.032, 1.049) *Meta-p* = 1.7E-21

Horvath

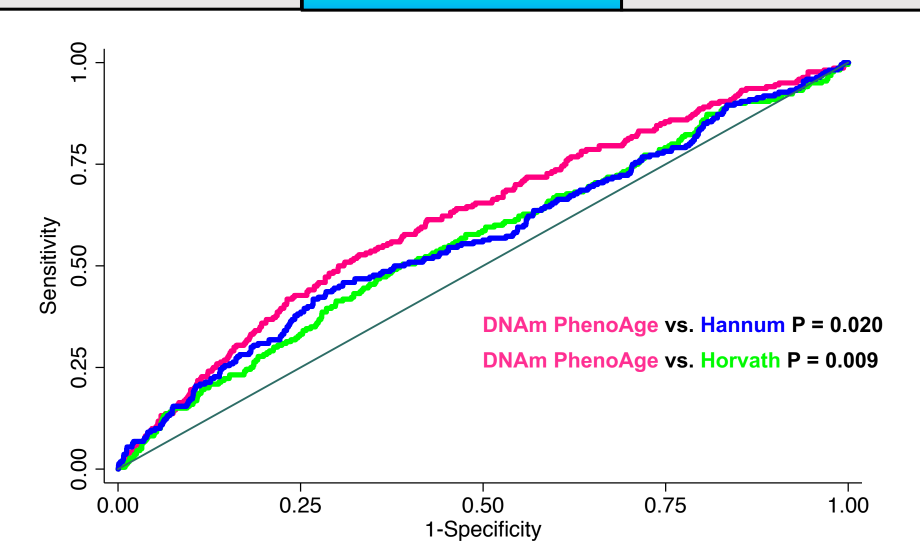


HR = 1.017 (1.009, 1.025) *Meta-p* = 4.5E-05

	Disease Count	Disease Free	CHD Risk	Physical Functioning
Levine	4.56E-15	1.06E-07	2.43E-10	2.05E-13
Horvath	6.76E-06	2.03E-03	1.10E-03	2.03E-05
Hannum	4.54E-02	1.31E-03	7.51E-01	4.66E-04

Develop a Multisystem Phenotypic Age Estimate and Validate Predictions Develop a New Epigenetic Age Estimate and Validate Predictions/Associations

Underlying Biology of the Clock and the 513 CpGs



MORTALITY & MORBIDTY PREDICTIONS

Breast Cancer Incidence (4% increased risk)





Lung Cancer Incidence (10% increased risk)



Centenarian
Offspring
(2.4 years
younger)

Down Syndrome (5-12 years older)



HIV infection (8 years older)





MCI (2.4 years older)

Develop a Multisystem Phenotypic Age Estimate and Validate Predictions Develop a New Epigenetic Age Estimate and Validate Predictions/Associations

Underlying Biology of the Clock and the 513 CpGs



PRECIPITATING FACTORS

- **Exercise**
- **Females**
- **1** Meat Consumption
 - Income

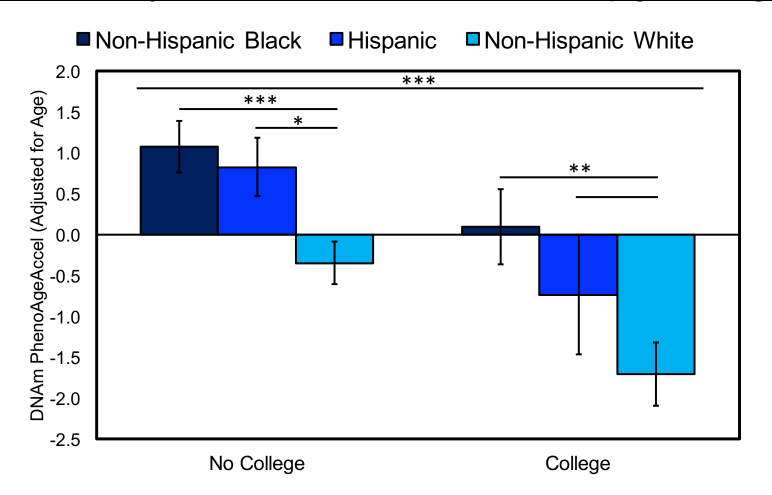




Develop a Multisystem Phenotypic Age Estimate and Validate Predictions Develop a New Epigenetic Age Estimate and Validate Predictions/Associations

Underlying Biology of the Clock and the 513 CpGs

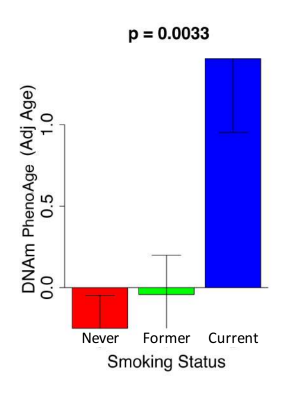
Race/Ethnicity and SES Relate to Differences in Epigenetic Age

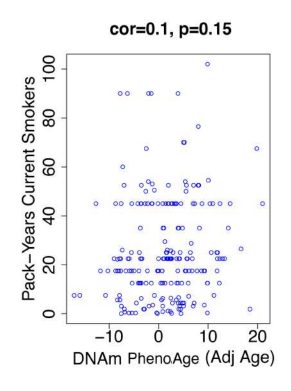


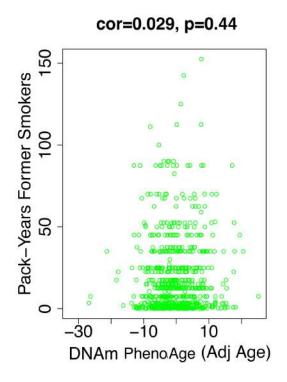
Develop a Multisystem Phenotypic Age Estimate and Validate Predictions Develop a New Epigenetic Age Estimate and Validate Predictions/Associations

Underlying Biology of the Clock and the 513 CpGs

Smoking, but not pack-years is associated with higher DNAm PhenoAge.



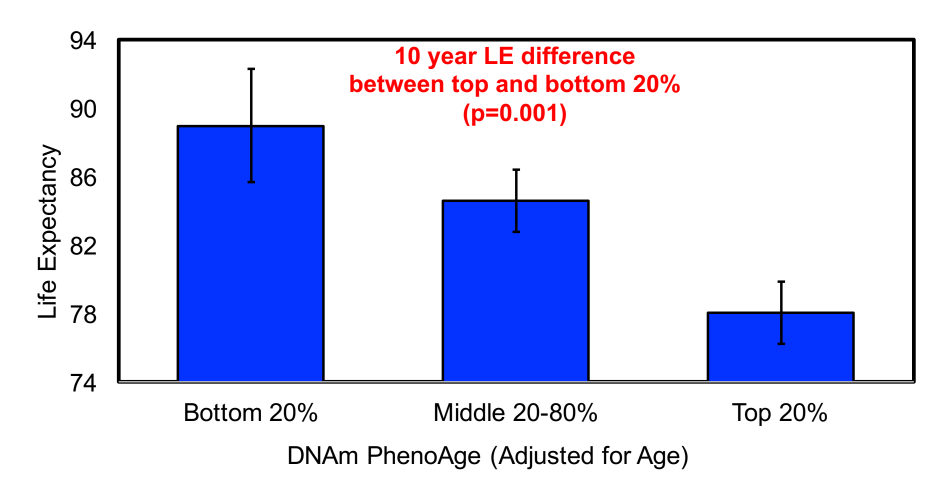




Develop a Multisystem Phenotypic Age Estimate and Validate Predictions Develop a New Epigenetic Age Estimate and Validate Predictions/Associations

Underlying Biology of the Clock and the 513 CpGs

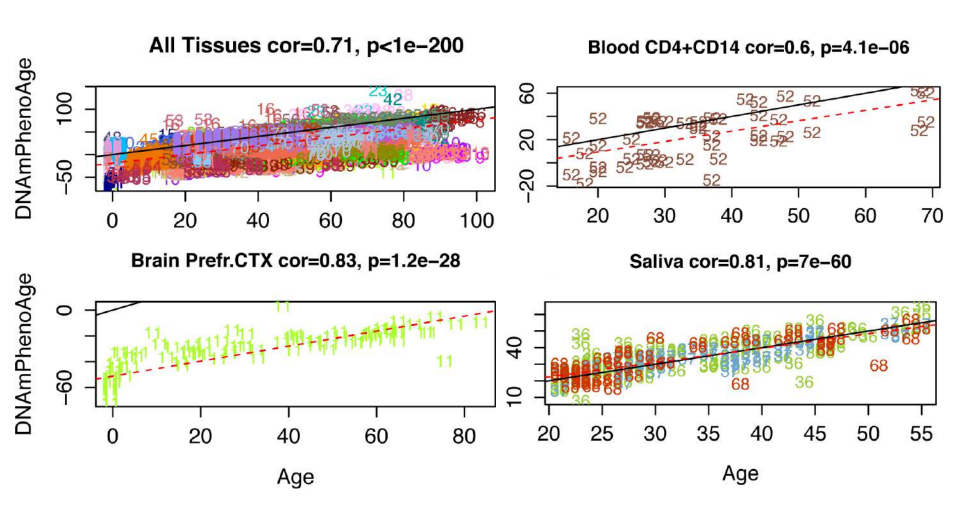
Does DNAm PhenoAge Capture Resilience?



Develop a Multisystem Phenotypic Age
Estimate and Validate Predictions

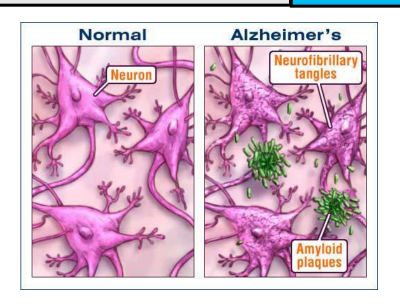
Develop a New Epigenetic Age Estimate and Validate Predictions/Associations

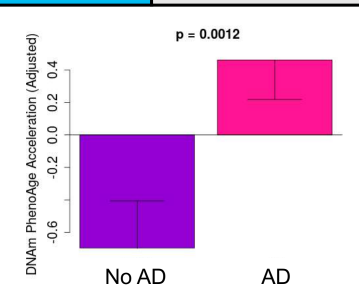
Underlying Biology of the Clock and the 513 CpGs



Develop a Multisystem Phenotypic Age Estimate and Validate Predictions Develop a New Epigenetic Age Estimate and Validate Predictions/Associations

Underlying Biology of the Clock and the 513 CpGs





Multivariate Associations with DNAm PhenoAge			
Beta (P-Value)			
Amyloid Load	d 0.451 (0.004)		
Neuritic Plaques	0.468 (0.004)		
Diffuse Plaques	0.377 (0.021)		
Neurofibrillary Tangles 0.100 (0.006)			
Results are from independent multivariate models that adjust for age at death, study, and sex			

Develop a multi-system estimate of "Phenotypic Age". Predictor of agingrelated mortality based on clinical measures.

Validate Associations with:

All-Cause Mortality
Cause Specific Mortality
Coexisting Disease Count
Physical Functioning
Mortality Ages 20-64
Mortality Ages 65-79
Mortality Ages 85+



Train a composite epigenetic predictor of phenotypic age, called "DNAm PhenoAge".

Based on DNAm at 513 CpGs.

Validate Associations with:				
All-Cause Mortality	Familial Longevity	Socioeconomic Status		
Coronary Heart Disease Risk	Dementia	Race/ethnicity		
Coexisting Disease Count	Down Syndrome	Diet		
Physical Functioning	Parkinson's Disease	Physical activity		
Disease Free Status	HIV positive	Metabolic Syndrome		
Age at Menopause	Chronological age in 35 tissues/cells	Smoking Status		
Cancer (Lung, Breast)	Neuropathology (Brain DNAm)	Obesity (Liver DNAm)		



Identify <u>underlying</u>
<u>biology</u> of the 513
CpGs in the DNAm
PhenoAge Score

lest	ior:
GO Enrichment	Immune Cell Associations
Pathway Enrichment	CpG Overlap with Hannum/Horvath
LTL Correlation	Loci-Specific DNAm vs. Transcription
Differential Expression	Polycomb Group Protein Targets
CpG Island Enrichment	Chromosomal Locations
Heritability Analysis	Change in DNAm PhenoAge over Time
DNAm Network Analysis	Transcriptional Analysis in Monocytes

Develop a Multisystem Phenotypic Age Estimate and Validate Predictions Develop a New Epigenetic Age Estimate and Validate Predictions/Associations

Identify the Underlying Biology of the Clock and the 513 CpGs

SNP HERITABILITY (h^2)

Defined as the total proportion of phenotypic variance attributable to genetic variation h^2 =0.38 to 0.54













CONCLUSIONS

- Developed an aging biomarker that is predictive/relates to numerous multifactorial aging conditions and outcomes.
 - Better predictor than the Horvath & Hannum clocks
 - Predicts after adjusting for confounders (smoking, cell counts).
- 2) Variation in the residual relates to genetic, social, behavioral, and demographic factors.
- 3) Reliable age correlations in 35 different tissues.
- 4) Variations is non-blood tissues predict outcomes that are pathologically/physiologically related to that tissues.

NEXT STEPS

- Tissue Consensus WGCNA (group CpGs)
- Identify genetic determinants

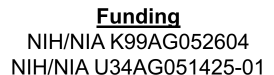




ACKNOWLEDGEMENTS

Collaborators

Steve Horvath, UCLA
Ake Lu, UCLA
Austin Quach, UCLA
Luigi Ferrucci, NIA
Brian Chen, NIA
Themistocles Assimes, Stanford
Lifang Hou, Northwestern
Andrea Baccarelli, Columbia
Eric Whitsel, UNC-Chapel Hill









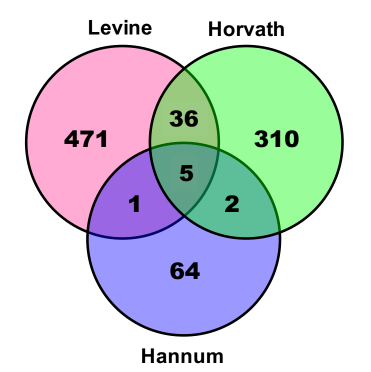




Develop a Multisystem Phenotypic Age Estimate and Validate Predictions Develop a New Epigenetic Age Estimate and Validate Predictions/Associations

Underlying Biology of the Clock and the 513 CpGs

	Levine DNAm Age	Horvath DNAm Age	Hannum DNAm Age
Levine DNAm Age	1	0.460	0.482
Horvath DNAm Age	0.460	1	0.511
Hannum DNAM Age	0.482	0.511	1



Only moderate correlations between the three clocks after adjusting for chronological age.

The clocks are not using the same CpGs.

They appear to be capture different phenomena.