Genetic Analyses of Epigenetic Predictors that Estimate Aging, Metabolic Traits, and Lifespan

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Abstract

DNA methylation (DNAm) remodeling is an important aspect of aging. Recently, a ‘pan-mammalian’ microarray was developed that provides a unifying platform to profile conserved CpGs across mammalian clades, and to develop universal biomarkers for aging. We used this array to measure DNAm in 339 predominantly female mice belonging to the BXD family. We computed epigenetic clocks and predicted maximum lifespan (predicted-maxLS), and tested associations with DNAm entropy, diet, weight, metabolic traits, and genetic variation.

Epigenetic age acceleration (EAA) and predicted-maxLS were correlated with genotype-dependent lifespan of the BXD strains. Genetic analyses uncovered quantitative trait loci (QTLs) on chromosome (Chr) 11 that encompasses the Erbb2/Her2 oncogenic region, and on Chr19 that contains a cytochrome P450 cluster. Both loci harbor candidate genes associated with EAA in humans (STXBP4, NKX2-3, CUTC). Transcriptome and proteome analyses revealed enrichment in oxidation-reduction and metabolic genes. The results highlight loci concordant in humans and mice, and demonstrate intimate links between metabolism, body weight, and epigenetic aging.

Keywords: epigenetic clock, lifespan, aging, DNA methylation, QTL, weight, diet
Introduction

Epigenetic clocks are widely used molecular biomarkers of aging. These biological clocks are based on the methylation status across an ensemble of “clock CpGs” that are collectively used to derive a DNA methylation (DNAm) based estimate of age (DNAmAge). This estimate tracks closely, but not perfectly, with an individual’s chronological age. How much the DNAmAge deviates from the known chronological age is a measure of the rate of biological aging. Denoted as epigenetic age acceleration (EAA), a more accelerated measure (positive EAA) suggests an older biological age. While DNAmAge predicts age, its age-adjusted counterpart, EAA, is associated with health, fitness, exposure to stressors, body mass index (BMI), and even life expectancy. These DNAm clocks were initially reported for humans. Since then, the biomarker has been extended to model organisms, and different variants of human clocks have also been developed. Some clocks are tissue specific, others are pan-tissue, and others perform well at predicting health and life expectancy.

A new microarray platform was recently developed to profile CpGs that have high conservation across mammalian clades. This pan-mammalian DNam array (HorvathMammalMethylChip40) provides a common platform to measure DNAm, and has been used to build universal epigenetic clocks that can estimate age across a variety of tissues and mammalian species. Another remarkable development with this array is the novel lifespan predictor that can estimate the maximum lifespan of over 190 mammals at high accuracy.

Here, we test the performance and applicability of these novel DNAm clocks and lifespan predictor in a genetically diverse cohort of mice belonging to the BXD family. The BXDs are a well-established mouse genetic reference panel that were first created as a family of recombinant inbred (RI) strains by crossing two inbred progenitors: C57BL/6J (B6) and DBA/2J (D2). The family has been expanded to ~150 fully sequenced progeny strains. The BXD population incorporates significant genetic variation, and likewise, show considerable phenotypic variability in their metabolic profiles, response to diet, aging rates, and natural life expectancy. The availability of genetic profiles, and accompanying deep multi-omics and phenomics, make the BXDs a powerful experimental population to evaluate the DNAm biomarkers, and to dissect the genetic modulators of epigenetic aging. Previously, we explored the aging methylome in a small number of BXD cases and found that high fat diet (HFD) and higher body weight were associated with higher age-dependent changes in methylation. In the present work, our goals were to (1) test the accuracy of the DNAm measures in predicting age and lifespan of the BXDs, (2) examine associations with diet and metabolic characteristics, and (3) apply quantitative trait locus (QTL) mapping and gene expression analyses to uncover loci and genes associated with these DNAm biomarkers.

Our results are consistent with a faster clock for cases on HFD, and with higher body weight. Both the DNAmAge and lifespan predictors were correlated with the genotype-dependent life expectancy of female BXDs. We report QTLs on chromosomes (Chrs) 11 and 19. A strong candidate gene in the chromosome (Chr) 11 interval (referred to as Eaaq11) is Stxbp4, a gene that has been consistently associated with EAA by human genome-wide association studies (GWAS). The Chr19 QTL (Eaaq1.9) also harbors strong contenders including Cyp26a1, Myof, Cutc, and Nkx2–3, and the conserved genes in humans have been associated with longevity and...
EAA may also have an effect on body weight change with age. We performed gene expression analyses to clarify the molecular pathways associated with the DNAm traits, and this, perhaps unsurprisingly, highlighted metabolic networks as strong expression correlates of epigenetic aging.

**Results**

**Description of samples**

The present study used liver DNAm data from 339 predominantly female mice (18 males only) belonging to 45 isogenic members of the BXD family, including F1 hybrids, and parental strains. Age ranged from 5.6 to 33.4 months. Mice were all weaned onto a normal chow (control diet; CD) and a balanced subset of cases were then randomly assigned to HFD (see Roy et al. for details). Tissues were collected at approximately six months intervals (see Williams et al.). Individual-level data of cases used in this study are in Data S1.

**DNAm clocks and chronological age prediction**

The first epigenetic clock we tested was the universal relative age clock described in Lu et al. We refer to the age estimates as univ.DNAmAge, and this had a correlation of $r = 0.92$ with the chronological age of mice (Table 1; Fig 1a).

We also built three different mouse clock, and each was developed as a pair depending on whether the training set used all tissues (pan-tissue) or only liver (liver specific). These are: (1) a general DNAm clock (referred to simply as DNAmAge): clock trained without pre-selecting for any specific CpG subsets; (2) developmental clock (dev.DNAmAge): built from CpGs that change during development; and (3) interventional clock (int.DNAmAge): built from CpGs that change in response to aging related interventions such as caloric restriction and growth hormone knockout. These clocks were trained in a large mouse dataset that excluded the BXDs (previously reported in 16,17), and are therefore unbiased to BXD characteristics. The specific CpGs and coefficients for clock computation are in Data S2. All the mouse clocks performed well in age estimation but the interventional clocks had higher deviation from chronological age.

| Clock type         | DNAmAge name | Tissue | $r$ with age (n=339) | $r$ with entropy (n=339) |
|--------------------|--------------|--------|---------------------|-------------------------|
| Universal clock    | univ.DNAmAge | pan    | 0.92                | 0.59                    |
| Standard clocks    | DNAmAge      | pan    | 0.89                | 0.50                    |
|                    |              | liver  | 0.92                | 0.51                    |
| Developmental      | dev.DNAmAge  | pan    | 0.87                | 0.46                    |
| clocks             |              | liver  | 0.91                | 0.45                    |
| Interventional     | int.DNAmAge  | pan    | 0.85                | 0.38                    |
| clocks             |              | liver  | 0.86                | 0.49                    |
| Lifespan           | Predicted-maxLS | pan | -0.07 (ns) | -0.13 (p=0.02) |
| predictor          |              |        |                     |                         |
(Table 1). The age-adjusted EAA derived from these clocks showed wide individual variation (Fig 1b).

We also used the universal maximum lifespan predictor\(^\text{17}\) to estimate the potential maximum lifespan (predicted-maxLS) of mice (Data S1). Predicted-maxLS was uncorrelated with chronological age (Table 1), and this is expected since the chronological age represents the time when the biospecimens were collected; not the time of natural demise. Instead, the predicted-maxLS showed an overall inverse correlation with EAA from the different clocks, and this suggests higher age-acceleration for mice with lower predicted-maxLS (Data S3).

**Association with methylome entropy**

The methylome-wide entropy provides a measure of randomness and information loss, and this increased with chronological age (Fig 1c).\(^7\) As direct correlates of chronological age, all the DNAmAge estimates had significant positive correlations with entropy (Table 1).

We hypothesized that higher entropy levels will be associated with (a) higher EAA, and (b) lower predicted-maxLS. Indeed, the univ.EAA had a positive correlation with entropy that was significant regardless of diet (Fig 1d). However, the EAA from the mouse clocks showed only weak correlations with entropy (Data S3). Entropy had a modest negative correlation with predicted-maxLS primarily in the CD group (Fig 1e). Taken together, our results indicate that discordance in the methylome increases with age, and is higher with higher univ.EAA. Mice with shorter predicted-maxLS may also had slightly higher entropy.

Fig 1. DNA methylation readouts and inter-correlations

(a) Correlation between predicted age (universal clock) and chronological age. Mice on control diet depicted by black circles (CD; normal lab chow; n = 210); mice on high fat diet (HFD; n = 129) by red crosses. (b) Violin plots of epigenetic age acceleration (EAA) derived from different DNAmAge clocks. (c) Shannon entropy, calculated from the full set of high quality CpGs, increases with age. (d) DNA methylation entropy has a direct correlation with EAA derived from the universal clock. (e) There is a slight inverse correlation between the entropy and predicted maximum lifespan (predicted-maxLS).
How the epigenetic readouts relate to diet, body weight, and sex

**Diet.** EAA from most of the clocks, including the universal clock, were significantly higher in the HFD group (Table 2). Entropy was also significantly higher in the HFD group. The maxLS did not differentiate between diets (Table 2).

**Body weight.** Body weight was first measured when mice were at an average age of 4.5 ± 2.7 months. We refer to this initial weight as baseline body weight (BW0). For mice on HFD, this was usually before introduction to the diet, with the exception of 48 cases that were first weighed 1 or 3 days after HFD (Data S1). In the CD group, only the EAA from the pan-tissue standard clock and liver developmental clock showed significant positive correlations with BW0 (Table 2). In the HFD group, the positive correlation with BW0 was more robust and consistent across all the clocks, and this may have been due to the inclusion of the 48 cases that had been on HFD for 1 or 3 days. Taking only these 48 cases, we found that higher weight even after 1 day of HFD had an age-accelerating effect (Data S3). This was particularly strong for the interventional clocks (r = 0.45, p = 0.001 for int.EAA, pan-tissue; r = 0.58, p < 0.0001 for int.EAA, liver), and for the universal clock (Fig 2a). Second weight was measured 7.4 ± 5.2 weeks after BW0 (mean age 6.3 ± 2.8 months). We refer to this as BW1 and we estimated the weight change as deltaBW = BW1 – BW0. DeltaBW was a positive correlate of EAA on both diets, albeit more pronounce in the HFD group (Fig 2b; Data S3). The final body weight (BWF) was measured.

### Table 2. Association with diet and weight, and heritability of the epigenetic readouts

| Type         | EAA            | Diet  | Mean (SD) | Diet (p) | r  | p  | r  | p  | H² | Strain (p) |
|--------------|----------------|-------|-----------|----------|----|----|----|----|----|------------|
| Universal    | univ.EAA       | CD    | -0.08 ± 0.22 | <0.001   | 0  | ns | 0.11 | ns | 0.37 | 0.67       |
| clock        |                | HFD   | 0.13 ± 0.27 |          |    |    | 0.50 | <0.0001 |    |            |
| EAA, pan     | CD             | HFD   | -0.05 ± 0.21 | <0.001   | 0.19 | 0.006 | 0.29 | <0.0001 | 0.49 | 0.54       |
|              |                |       | 0.07 ± 0.21 |          |    |    | 0.42 | <0.0001 |    |            |
| EAA, liver   | CD             | HFD   | 0 ± 0.17    |        | 0.09 | ns | 0.20 | 0.003 | 0.40 | 0.73       |
|              |                |       | 0.03 ± 0.14 |          |    |    | 0.49 | <0.0001 |    |            |
| dev.EAA, pan | CD             | HFD   | -0.04 ± 0.23 | 0.004   | 0.09 | ns | 0.22 | 0.001 | 0.53 | 0.76       |
|              |                |       | 0.03 ± 0.22 |          |    |    | 0.45 | <0.0001 |    |            |
| dev.EAA, liver | CD           | HFD   | 0 ± 0.2    |         | 0.19 | 0.002 | 0.29 | <0.0001 | 0.46 | 0.78       |
|              |                |       | 0 ± 0.16   |          |    |    | 0.47 | <0.0001 |    |            |
| int.EAA, pan | CD             | HFD   | -0.05 ± 0.25 | 0.0003  | 0.03 | ns | 0.21 | 0.002 | 0.27 | 0.66       |
|              |                |       | 0.06 ± 0.33 |          |    |    | 0.46 | <0.0001 |    |            |
| int.EAA, liver | CD           | HFD   | -0.04 ± 0.22 | <0.001  | 0.05 | ns | 0.18 | 0.01  | 0.59 | 0.80       |
|              |                |       | 0.11 ± 0.25 |          |    |    | 0.58 | <0.0001 |    |            |
| Entropy      | CD             | HFD   | 10.034 ± 0.007 | 0.004   | -0.15 | 0.03 | -0.34 | <0.0001 | 0.39 | 0.20 (ns)  |
|              |                |       | 10.036 ± 0.007 |        | -0.11 | 0.21 | -0.34 | <0.0001 |    |            |
| Pred-maxLS   | CD             | HFD   | 4.14 ± 0.37 | ns      | 0.03 | ns | 0.05 | ns    | 0.66 | 0.89       |
|              |                |       | 4.15 ± 0.32 |        | -0.06 | ns | -0.11 | 0.20 |    |            |

**a** BW0 is body weight at about 4.5 months of age (n = 339; 210 CD and 129 HFD); BWF is final weight at tissue collection (1 HFD case missing data; n = 338; 210 CD and 128 HFD)

**b** Pearson correlation between strain means for n = 29 BXA genotypes kept on CD and HFD
at the time of tissue harvest, and EAA from all the mouse clocks were significant correlates of BWF on both diets (Table 2).

Somewhat unexpected, entropy had an inverse correlation with body weight. This effect was primarily in the CD mice (Table 2). We found no association between predicted-maxLS and the body weight traits (Table 2).

Sex effect. Four BXD genotypes (B6D2F1, D2B6F1, BXD102, B6) had cases from both males and females. We used these to test for sex effects. All the mouse clocks showed significant age acceleration in male mice, and this effect was particularly strong for the pan-tissue int.EAA (Fig 2c; Data S3). The predicted-maxLS was significantly lower in males (Fig 2d). Entropy on the other hand, was significantly higher in females (Fig 2e).

**Fig 2. Correlates and modifiers of the epigenetic readouts**

(a) For 48 mice, initial body weight (BW0) was measured 1 or 3 days after high fat diet (HFD), and these showed significant correlation between BW0 and epigenetic age acceleration (EAA shown for the universal clock). (b) Weight was first measured at mean age of 4.5 ± 2.7 months, and then at 6.3 ± 2.8 months (BW1). Weight gains during this interval (deltaBW = BW1 – BW0) is a direct correlate of EAA derived from the interventional clock in both normal chow (control diet or CD; n = 210) and HFD mice (n = 128). (c) For the BXD genotypes with samples from both males and females, males have higher age acceleration and this sex effect is highly significant for the pan-tissue interventional clock (int.EAA). Bars represent mean ± standard error; 40 females (26 CD, 14 HFD) and 18 males (10 CD, 8 HFD). Males had (d) lower predicted maximum-lifespan, and (e) lower average methylome entropy. (f–i) The residual plots display the direction of association between metabolic traits and DNA methylation (n = 276 cases with metabolic data). After adjusting for chronological age, diet and body weight, serum cholesterol has inverse associations with (f) predicted maximum lifespan (p = 0.002), and (g) methylome entropy (p = 9.1E-06). Serum glucose level has inverse associations with (h) epigenetic age acceleration derived from the universal clock (p = 0.005), and (i) methylome entropy (p = 0.003).
Association with metabolic traits

276 cases with DNAm data also had fasted serum glucose and total cholesterol, and we examined whether these metabolic traits are associated with three of the DNAm readouts: the univ.EAA, predicted-maxLS, and DNAm entropy. We applied linear regression with age, diet and final body weight as covariates, and this showed significant effects of cholesterol on predicted-maxLS (p = 0.002), and entropy (p = 9E-06) (Table S1). To visualize how cholesterol levels associate with these, we plotted the residual values after the respective predictor and outcome variables were adjusted for age, diet, and BWF. The residual plot shows an inverse association between cholesterol and predicted-maxLS (Fig 2f). For entropy, similar to how it related with weight, higher cholesterol predicted lower entropy. Cholesterol had no significant association with univ.EAA (Table S1).

Glucose had an unexpected inverse association with the univ.EAA that indicates lower age acceleration with higher glucose (p = 0.005) (Fig 2h; Table S1). Higher glucose was also associated with lower entropy (p = 0.003) (Fig 2i). Glucose was not associated with predicted-maxLS (Table S1).

Association with strain longevity

We next obtained longevity data from a parallel cohort of female BXD mice that were allowed to age on CD or HFD. We evaluated whether the DNAm readouts were informative of strain-level lifespan. Since the strain lifespan was determined from female BXDs, we restricted this to only the female cases. For strains with natural death data from n ≥ 5, we computed the minimum (minLS), 25th quartile (25Q-LS), mean, median lifespan, 75th quartile (75Q-LS), and maximum lifespan (maxLS) (Data S1). Specifically, we postulated (a) an accelerated clock for strains with shorter lifespan (i.e., inverse correlation), (b) a direct correlation between predicted-maxLS and observed lifespan, and (c) higher entropy with shorter lifespan.

Overall, the EAA measures showed the expected inverse correlation trend with the lifespan summaries, and this was highly significant for the universal clock (Table S2; Fig 3a,b). For the mouse clocks, this effect was significant for the liver int.EAA (Table S2). When separated by diet, these correlations became weaker, but the negative trend remained consistent.

The DNAm entropy had an inverse correlation trend with strain lifespan (Table S2). This was nominally significant only for the strain maxLS when CD and HFD groups were combined (r = −0.13, p = 0.02) but became non-significant when separated by diet.

The predicted-maxLS showed a positive correlation trend with the lifespan summaries, and this was significant for the observed strain maxLS (Fig 3d). When separated by diet, the predicted-maxLS remained a significant correlate of strain maxLS only in the CD group.
Genetic analysis of epigenetic age acceleration and predicted-maxLS

The EAA traits had modest to high heritability, and averaged at 0.50 for mouse clocks (Table 2). The predicted-maxLS had heritability of 0.66 on CD, and 0.70 on HFD. Another way to gauge level of genetic correlation is to compare between members of strains maintained on different diets. All the EAA traits, and the predicted-maxLS had high strain-level correlations between diets, indicating an effect of background genotype that is robust to dietary differences (Table 2).

To uncover genetic loci, we applied QTL mapping using mixed linear modeling that corrects for the BXD kinship structure. First, we performed the QTL mapping for the EAA derived from the different mouse and universal clocks, with adjustment for diet and body weight. EAA from the two interventional clocks had the strongest QTLs (Data S4). The pan-tissue int.EAA had a significant QTL on Chr11 (90–99 Mb) with the highest linkage at ~93 Mb (p = 3.5E-06; equivalent to a LOD score of 4.7) (Fig 4a). Taking a genotype marker at the peak interval (BXD variant ID DA0014408.4 at Chr11, 92.750 Mb), we segregated the BXDs homozygous for either the D2 (DD) or the B6 (BB) alleles. Strains with DD genotype at this locus had significantly higher int.EAA (Fig 4a inset).

The liver int.EAA had the peak QTL on Chr19 (35–45 Mb) with the most significant linkage at markers between 38–42 Mb (p = 9E-07; LOD score of 5.2) (Fig 4b). We selected a marker at the peak interval (rs48062674 at Chr19, 38.650 Mb), and the BB genotype had significantly higher int.EAA compared to DD (Fig 4b inset). The QTL map for the univ.EAA did not reach genome-wide significance (Fig 4c). However, there were nominally significant peaks at the Chr19 (p = 0.0004), and Chr11 (p = 0.004) intervals.

Fig 3. Predictors of strain longevity

BXD genotypes with shorter life expectancy tended to have a more accelerated universal clock (univ.EAA). This inverse relation is depicted for the (a) 25th and (b) 75th quartile age at natural death for female BXDs kept on either normal chow (CD) or high fat diet (HFD). (c) Age acceleration from the liver interventional mouse clock (int.EAA) also showed a similar inverse correlation with strain longevity, but effect was significant only when both diets were included (here illustrated with median lifespan). (e) The predicted maximum lifespan had a significant direct correlation with the observed strain maximum lifespan. Analysis in n = 302 female BXDs; 191 CD and 111 HFD.
We next performed QTL mapping for DNA methylation entropy with adjustment for major covariates (diet,

![Manhattan plots](image.png)

**Fig 4. Genetic linkage analysis**
The Manhattan plots represent the location of genotyped markers (x-axis), and linkage – \( \log_{10}p \) (y-axis). (a) The peak quantitative trait locus (QTL) for age acceleration from the pan-tissue interventional clock (int.EAA) is on chromosome (Chr) 11 at ~93 Mb. The inset shows the mean (± standard error) trait values for BXDs the are homozygous for the C57BL/6J allele (BB; grey) versus BXDs homozygous for the DBA/2J allele (DD; black). (b) The liver-specific int.EAA has a peak QTL on Chr19 (~38 Mb). Trait means by genotype at this locus are shown in inset. (c) The linkage statistics are weaker for the EAA derived from the universal clock (univ.EAA). However, there are consistent nominally significant peaks on the Chr11 and Chr19 loci. (d) The DNA methylation based predicted maximum lifespan also maps to Chr19, but the peak markers are at ~47.5 Mb.
chronological age, and body weight). No locus reached genome-wide significance (DataS4). There were modest QTLs on Chrs 11 and 19. However, the Chr11 region is slightly distal to the markers linked to the EAA traits (minimum p = 0.009 at Chr11, ~103.7 Mb). The Chr19 locus somewhat overlapped the QTL for EAA, but the peak marker (minimum p = 0.0009) is slightly distal at ~48 Mb (Data S4).

The predicted-maxLS had a significant QTL on Chr19 (Fig 4d; Data S4) with the peak markers between 44–48 Mb (p = 2E-07; LOD score of 5.9). This overlaps the EAA QTL, but the peak markers are distal (rs30567369 at 47.510 Mb). At this locus, mice with the BB genotype had significantly higher predicted-maxLS (Fig 4d inset).

To identify regulatory loci that are consistent across the different EAA measures, we applied a multi-trait analysis and derived the linkage meta-p-value for the mouse and universal EAA traits. The peaks on Chrs 11 and 19 attained the highest consensus p-values (Fig S1a; Data S3). Additional consensus peaks (at –log10meta-p > 6) were observed on Chrs 1 (~152 Mb), and 3 (~54 Mb).

We focus on the Chrs 11 and 19 QTLs and refer to these as EAA QTL on Chr 11 (Eaaq11), and EAA QTL on Chr 19 (Eaaq19). Eaaq11 extends from 90–99 Mb. For Eaaq19, we delineated a broader interval from 35–48 Mb that also encompasses the peak markers for the predicted-maxLS, albeit these may be separate loci related to EAA (~39 Mb of Eaaq19), and predicted-maxLS (~47 Mb of Eaaq19).

We performed marker-specific linkage analyses for each of the mouse and universal clocks using a regression model that adjusted for diet. With the exception of the liver int.EAA, all the EAA traits had nominal to highly significant associations with the representative Eaaq11 marker (DA0014408.4), and the DD genotype had higher age acceleration (Table 3). Mean plots by genotype and diet shows that this effect was primarily in the CD mice (Fig S1b). The effect of this locus appeared to be higher for the pan-tissue clocks compared to the corresponding liver-specific clocks. This marker in Eaaq11 was not associated with either entropy or predicted-maxLS.

For proximal Eaaq19, the representative marker (rs48062674) was associated with all the EAA traits and the BB mice had higher age acceleration on both diets (Fig S1c). This marker was not associated with entropy, and had only a weak effect on predicted-maxLS (Table 3). When we performed the same analysis with the marker on distal Eaaq19 (rs30567369), the association with EAA became weaker, and the association with predicted-maxLS became much stronger (Table 3). This suggests that the proximal part of Eaaq19 is related to EAA while the distal part is related to predicted-maxLS.

We also tested if these peak markers were associated with the recorded lifespan phenotype and we found no significant association with the observed lifespan of the BXDs.
Association of EAA QTLs with body weight trajectory

Since gain in body weight with age was an accelerator of the clocks, we examined whether the selected markers in Eaaq11 and Eaaq19 were also related to body weight change. We retrieved

Table 3: Marker specific linkage analyses for epigenetic age acceleration, predicted maximum lifespan, and body weight trajectory

| Predictor | Outcome          | Estimate | Std Error | t Ratio | p      |
|-----------|------------------|----------|-----------|---------|--------|
| Eaaq11    | EAA, pan         | 0.096    | 0.023     | 4.184   | 3.8E-05|
|           | EAA, liver       | 0.067    | 0.017     | 3.880   | 0.0001 |
|           | dev.EAA, pan     | 0.077    | 0.025     | 3.041   | 0.003  |
|           | dev.EAA, liver   | 0.037    | 0.020     | 1.878   | 0.06   |
|           | int.EAA, pan     | 0.153    | 0.029     | 5.278   | 2.5E-07|
|           | int.EAA, liver   | -0.033   | 0.025     | -1.284  | 0.20   |
|           | univ.EAA         | 0.101    | 0.025     | 4.057   | 6.3E-05|
| Eaaq19    | EAA, pan         | -0.083   | 0.028     | -2.954  | 0.003  |
| rs48062674 | EAA, liver       | -0.137   | 0.020     | -6.972  | 2.0E-11|
| Chr19, 38.650 Mb     | dev.EAA, pan     | -0.206   | 0.029     | -7.218  | 4.3E-12|
|           | dev.EAA, liver   | -0.124   | 0.023     | -5.461  | 9.9E-08|
|           | int.EAA, pan     | -0.143   | 0.035     | -4.028  | 7.1E-05|
|           | int.EAA, liver   | -0.250   | 0.027     | -9.238  | 4.6E-18|
|           | univ.EAA         | -0.145   | 0.029     | -4.932  | 1.3E-06|
|           | Pred-maxLS       | -0.100   | 0.048     | -2.086  | 0.04   |
| Distal Eaaq19 | int.EAA, liver | -0.079   | 0.026     | -2.995  | 0.003  |
| rs30567369   | univ.EAA         | -0.053   | 0.026     | -2.012  | 0.05   |
| Chr19, 47.510 Mb   | Pred-maxLS       | -0.383   | 0.036     | -10.781 | 3.9E-23|

Mixed model for longitudinal change in body weight

| Predictor | Outcome   | Estimate | Std Error | t Ratio | p      |
|-----------|-----------|----------|-----------|---------|--------|
| Eaaq11    | Body weight | 0.619    | 0.345     | 1.794   | 0.07   |
| DA0014408.4[DD] | Number of observations = 6885; number of individuals = 2112 | |
| Eaaq19    | Body weight | -1.847   | 0.374     | -4.945  | 7.6E-07|
| rs48062674[DD] | Number of observations = 6132; number of individuals = 1852 | |
| Distal Eaaq19 | Body weight | -1.619   | 0.363     | -4.458  | 8.3-06 |
longitudinal weight data from a larger cohort of the aging BXD mice that were weighed at regular intervals. After excluding heterozygotes, we tested the effect of genotype. Concordant with the higher EAA for the DD genotype at Eaaq11 in the CD group, the DD genotype in the CD group also had slightly higher mean weight at older adulthood (12 and 18 months; Fig 5a). However, this marker had no significant association with body weight when tested using a mixed effects model (p = 0.07; Table 3). In proximal Eaaq19, it was the BB genotype that exhibited consistently accelerated clock on both diets, and the BB genotype also had higher average body weight by 6 months of age (Fig 5b), and this locus had a significant influence on the body weight trajectory (p = 7.6E-07; Table 3). The nearby marker on distal Eaaq19 also showed a similar pattern of association with body weight (Table 3).

Fig 5. Body weight trajectory by diet and genotype

Body weight was measured at regular age intervals (x-axis) from (a) 2112 BXD mice that were homozygous at the Leaaq11 marker (DA0014408.4; 842 BB, 1279 DD), and (b) 1852 BXD mice that were homozygous at the proximal Leaaq19 marker (rs48062674; 1252 BB, 600 DD). Mice were maintained on either control diet (CD) or high fat diet (HFD). The graphs show the segregation of body weight over time by diet and genotype. Mean ± standard error; heterozygotes were excluded.

Candidate genes for epigenetic age acceleration

There are several positional candidate genes in Eaaq11 and Eaaq19. To narrow the list, we applied two selection criteria: genes that (1) contain missense and/or stop variants, and/or (2) contain non-coding variants and regulated by a cis-acting expression QTL (eQTL). For the eQTL analysis, we utilized an existing liver transcriptome data from the same aging cohort.18 We identified 24 positional candidates in Eaaq11 that includes Stxbp4, Erbb2 (Her-2 oncogenic gene), and Grb7 (growth factor receptor binding) (Data S5). Eaaq19 has 81 such candidates that includes a cluster of cytochrome P450 genes, and Chuk (inhibitor of NF-kB) in the proximal region, and Pcgf6 (epigenetic regulator) and Elov3 (lipid metabolic gene) in the distal region (Data S5).

For further prioritization, we converted the mouse QTL regions to the corresponding syntenic regions in the human genome, and retrieved GWAS annotations for these intervals.33 We specifically searched for the traits: epigenetic aging, longevity, age of menarche/menopause/puberty, Alzheimer’s disease, and age-related cognitive decline and dementia. This highlighted 5 genes in Eaaq11, and 3 genes in Eaaq19 (Table S4). We also identified a GWAS study that found associations between variants near Myof-Cyp26a1 and human longevity,30 and a meta-GWAS that found gene-level associations between Nkx2–3 and Cutc, and epigenetic aging (Table S4).28

Gene expression correlates of EAA and predicted max-LS
Liver RNA-seq data was available for 153 of the BXD cases that had DNAm data (94 CD, and 59 HFD). We used this set to perform transcriptome-wide correlation analysis for the univ.EAA. To gain insights into gene functions, we selected the top 2000 transcriptome correlates (|r| ≥ 0.37, p ≤ 2.8E-06; Data S6) for functional enrichment analysis. These top correlates represented transcript variants from 1052 unique genes and included a few positional candidates (e.g., Ikzf3, Kif11, Cep55, Cyp2c29, Cyp2c37). Only 62 transcripts from 36 unique genes were negatively correlated with univ.EAA, and this set was significantly enriched (Bonferroni correct p < 0.05) in oxidation-reduction, and metabolic pathways (Data S7; Fig 6a). These functional categories included the cytochrome genes, Cyp2c29 and Cyp2c37, located in Eaaq19. This set was also highly liver specific. The positive correlates were enriched in a variety of gene functions, and was not a liver-specific gene set (Data S7). Taking the top 10 GO categories, we can broadly discern two functional domains: immune and inflammatory response, and mitosis and cell cycle (Fig 6a). To verify that these associations are robust to the effect of diet, we repeated the correlation and enrichment analysis in the CD group only (n = 94). Again, taking the top 2000 correlates (|r| ≤ 0.30; p ≤ 0.003), we found the same enrichment profiles for the positive and negative correlates (Data S7).

Next, we performed the correlational analysis using liver proteomic data that was available for 164 of the BXDs with DNAm data. The proteome data quantifies over 32000 protein variants from 3940 unique genes. We took the top 2000 protein correlates of univ.EAA (|r| ≥ 0.27, p ≤ 6.0E-04) (Data S8). This represented protein levels from 563 unique genes. 1139 protein variants (215 genes) had negative correlations, and similar to the mRNA correlates, there was enrichment in oxidation-reduction and metabolic processes. This set was also enriched in liver

![Fig 6. Gene expression correlates of the universal clock](image)
genes, and included pathways related to lipid and steroid metabolism, epoxygenase p450 pathway, and xenobiotics (Fig 6b; Data S9). These categories were populated by the cytochrome genes including candidates in Eaaq19 (e.g., Cyp2c29, Cyp2c37). The positive proteome correlates showed a different functional profile than the transcriptomic set. These were enriched in genes related to transport (includes apolipoprotein such as APOE), cell adhesion, protein translation, protein folding, and metabolic pathways related to glycolysis and gluconeogenesis (Data S9).

We performed a similar transcriptome and proteome analysis for the predicted-maxLS. For mRNA, both the negative and positive correlates were enriched in metabolic pathways including glucose and lipid metabolism (Data S10, S11). Similarly, the positive and negative protein correlates of predicted-maxLS converged on oxidation-reduction processes (included cytochrome genes located in proximal Eaaq19) and metabolic pathways (Data S12, S13).

**Discussion**

The goals of this study were to examine the aging methylome at conserved CpGs, test the performance of the DNAm based biomarkers, and identify potential modifiers and genetic drivers. HFD had a strong age-accelerating effect that concurs with the association between EAA and obesity in humans.25,34,35 Age-acceleration due to diet manifested within the first 1 to 3 days of transitioning from normal lab chow to HFD. Even among the CD mice, higher weight gain at a younger age was associated with an accelerated clock.

Somewhat surprising was how entropy related to the metabolic traits. Epigenetic entropy increases with age, and is likely an indicator of the increase in stochastic noise over time.7,36 In biological systems, entropy is kept at bay by the uptake of energy, and investment in maintenance and repair.37 As HFD increased entropy (possibly due to higher cellular heterogeneity and adiposity of liver tissue), we expected entropy to be higher with higher body weight. But instead, entropy had an inverse correlation with weight, an effect that was primarily in the CD mice. Higher levels of serum glucose and total cholesterol were also associated with lower entropy. The reason for this is unclear, and we can only speculate that the enhanced energy consumption in mice that had higher metabolic substrates may have kept the methylome in a more ordered state. Despite this, mice with higher entropy also tended to have higher EAA. Entropy had a modest negative correlations with both the the DNAm based predicted-maxLS, and the known strain-level maxLS. The predicted-maxLS on the other hand, showed no direct association with diet or body weight, but higher total cholesterol and higher EAA were associated with shorter predicted-maxLS.

For the BXDs, life expectancy is highly dependent on the background genotype, and mean lifespan varies from under 16 months for strains such as BXD8 and BXD13, to over 28 months in strains such as BXD91 and BXD175.19,22,24 While the EAA from the universal was inversely correlated with strain lifespan data, the predicted-maxLS had a direct correlation with observed strain maxLS. We note that the predicted-maxLS overestimated the strain max-LS by 0.7 to 3 years (median error of +1.6 years). Nonetheless, the correlation between individual-level predicted-, and strain-level observed maxLS is remarkable considering that both the universal clock and max-LS predictor are pan-mammalian, and species- and tissue-agnostic.16,17
results suggest that these universal epigenetic predictors of biological aging and lifespan are informative of the subtle and normative lifespan variation in a family of inbred mice. The analysis between the epigenetic readouts and lifespan was also an indirect comparison. Unlike the comparison with body weight and metabolic traits, which were traits measured from the same individual, the lifespan data are strain characteristics computed from a parallel cohort of mice that were allowed to survive till natural mortality. Nonetheless, this indirect comparison demonstrates that these multi-species epigenetic predictors capture genotype-dependent effects on aging and lifespan within a species.

We tested different versions of the mouse DNAmAge clocks, and these appeared to capture slightly different aspects of epigenetic aging. For instance, the interventional clocks were sensitive to diet and early weight change, but not related to BW0 in the CD mice. Instead, BW0 had a significant accelerating effect on the liver developmental clock (dev.EAA).

Our goal was to take these different clocks and identify regulatory loci that were the most stable and robust to the slight algorithmic differences in building the clocks. A notable candidate in Eaaq11 is Syntaxin binding protein 4 (Stxbp4, aka, Synip), located at 90.5 Mb. Stxbp4 is a high-priority candidate due to the concordant evidence from human genetic studies. The conserved gene in humans is a replicated GWAS hit for the intrinsic rate of epigenetic aging.26-28 In the BXDs, Stxbp4 contains several non-coding variants, and a missense mutation (rs3668623), and the expression of Stxbp4 in liver is modulated by a cis-eQTL. Stxbp4 plays a key role in insulin signaling,38 and has oncogenic activity and implicated in different cancers.39,40 Furthermore, GWAS have also associated STXB4 with age of menarche.41,42 Eaaq11 corresponds to the 17q12-21 region in humans, and the location of additional oncogenic genes, e.g., ERBB2/HER2, GRB7, and BRCA1.43 The mouse Brca1 gene is a little distal to the peak QTL region and is not considered a candidate here, although it does segregate for two missense variants in the BXDs. Erbb2 and Grb7 are in the QTL region, and Erbb2 contains a missense variant (rs29390172), and Grb7 is modulated by a cis-eQTL. Nr1d1 is another candidate in Eaaq11, and the co-activation of Erbb1, Grb7, and Nr1d1 has been linked to breast and other cancers.44,45

Eaaq19 was consistently associated with EAA from all the clocks we evaluated, and also with body weight gains, irrespective of diet. The predicted-maxLS also maps to this region, and DNAm entropy may also have a weak association with markers at this interval. The EAA traits have peak markers in the proximal part of Eaaq19 (around the cytochrome cluster), and the predicted-maxLS peaks in the distal portion (over candidates like Elovl3, Pcgf3). Two candidates in Eaaq19 have been implicated in epigenetic aging in humans based on gene-level meta-GWAS: NK homeobox 3 (Nkx2-3, a developmental gene), and CutC copper transporter (Cutc).28 Eaaq19 is also the location of the Cyp26a1-Myof genes, and the human syntenic region is associated with longevity, metabolic traits, and lipid profiles.30,46,47 Another noteworthy candidate in Eaaq19 is Chuk, a regulator of mTORC2, that has been associated with age at menopause.41,48 Clearly, Eaaq19 presents a complex and intriguing QTL related to the different DNAm readouts, and potentially metabolic traits. Both Eaaq19 and Eaaq11 exemplify the major challenge that follows when a genetic mapping approach leads to gene- and variant-dense regions.49,50 Both loci have several biologically relevant genes, and identifying the causal gene (or genes) will require a more fineScaled functional genomic dissection.
The gene expression analyses highlighted metabolic pathways related to lipids, glucose, and proteins for both the univ.EAA and predicted-maxLS. Other enriched pathways were mitosis and cell division, and immune processes, but this was specific to the positive transcriptomic correlates. The more compelling evidence is for the cytochrome P450 genes, which are both positional candidates, as well as expression correlates at the transcriptomic and proteomic levels. These genes have high expression in liver, and have major downstream impact on metabolism.\(^{51-53}\) One caveat is that these CYP genes are part of a gene cluster in Eaaq19 that includes transcripts with cis-eQTLs (e.g., Cyp2c66, Cyp2c39, Cyp2c68), and the tight clustering of the genes, and proximity of trait QTL and eQTLs may result in tight co-expression due to linkage disequilibrium.\(^{54}\) Nonetheless, the cytochrome genes in Eaaq19 are strong candidate modulators of EAA that calls for further investigation.

Aside from Eaaq11 and Eaaq19, loci with evidence of consensus QTLs were also detected on Chr 1 and 3. We do not delve into these in the present work, but the Chr3 interval is near genes associated with human epigenetic aging (ift80, trim59, kpna4).\(^{26,28}\) However, this QTL is dispersed across a large interval, and the peak markers do not exactly overlap these human EAA GWAS hits. While we have focused on Eaaq11 and Eaaq19, these other loci also present potentially important regions for EAA.

In summary, we have identified two main QTLs—Eaaq11 and Eaaq19—that contribute to variation in two DNA readouts: EAA, and predicted-maxLS. Eaaq11 contains several genes with oncogenic properties (e.g., Stxbp4, Erbb2), while Eaaq19 contains a dense cluster of metabolic genes (e.g., Elov13, Chuk, the cytochrome genes). We demonstrate that metabolic profile and body weight are closely related to epigenetic aging. The convergence of evidence from genetic and gene expression analyses indicates that genes involved in metabolism and energy balance contribute to the age-dependent restructuring of the methylome, which in turn forms the basis of the epigenetic predictors of aging and lifespan.

### Materials and Methods

#### Biospecimen collection and processing

Samples for this study were selected from a larger colony of BXD mice that were housed in a specific pathogen-free (SPF) facility at the University of Tennessee Health Science Center (UTHSC). All animal procedures were in accordance with a protocol approved by the Institutional Animal Care and Use Committee (IACUC) at the UTHSC. Detailed description of housing conditions and diet can be found in.\(^{18,19}\) Mice were given *ad libitum* access to water, and either standard laboratory chow (Harlan Teklad; 2018, 18.6% protein, 6.2% fat, 75.2% carbohydrates), or high-fat chow (Harlan Teklad 06414; 18.4% protein, 60.3% fat, 21.3% carbohydrate). Animals were first weighed within the first few days of assignment to either diets, and this was mostly but not always prior to introduction to HFD. Following this, animals were weighed periodically, and a final time (BWf) when animals were humanely euthanized (anesthetized with avertin at 0.02 ml per g of weight, followed by perfusion with phosphate-buffered saline) at specific ages for tissue collection. The present work utilizes the biobanked liver specimens that were pulverized and stored in -80 °C, and overlaps samples described in Williams et al.\(^{18}\) DNA was extracted using the DNeasy Blood & Tissue Kit from Qiagen. Nucleic
acid purity was inspected with a NanoDrop spectrophotometer, and quantified using a Qubit fluorometer dsDNA BR Assay.

**Methylation array, quality check, and entropy calculation**

DNA samples from ~350 BXD mice were profiled on the Illumina Horvath Human MethylChip40 array. Details of this array are described here Arneson et al.\textsuperscript{15} The array contains probes that target ~36K highly conserved CpGs in mammals. Over 33K probes map to homologous regions in the mouse genome, and data from these were normalized using the SeSame method.\textsuperscript{55} Unsupervised hierarchical clustering was performed to identify outliers and failed arrays, and those were excluded. We also performed strain verification as an additional quality check. While majority of the probes were free of DNA sequence variants, we found 45 probes that overlapped variants in the BXD family. We leveraged these as proxies for genotypes, and performed a principal component analysis. The top principal component (PC1 and PC2) segregated the samples by strain identity, and samples that did not cluster with the reported strains were removed. After excluding outliers, failed arrays, and samples that failed strain verification, the final liver DNAm data consisted of 339 samples.

For entropy calculation, we used 27966 probes that have been validated for the mouse genome using calibration data generated from synthetic mouse DNA.\textsuperscript{56} Shannon entropy was calculated for each sample using the R package, “entropy” (v1.2.1) with method = “ML”: maximum likelihood.\textsuperscript{57}

**Clock estimation and maximum lifespan predictor**

The development of the universal pan-tissue epigenetic clocks of age, and the universal maximum lifespan predictor are described in Lu et al.\textsuperscript{16} and Li et al.\textsuperscript{17} respectively. For the present work, we utilized the universal clock that predicts relative age, defined as individual age relative to the maximum lifespan of its species, followed by inverse transformation to estimate DNAmAge.\textsuperscript{16}

The mouse specific clocks reported in this study were built with a similar strategy as described in Lu et al.\textsuperscript{16} using elastic net regression. First, a log-linear transformation was applied to the chronological age using the function:

$$f(Age) = \begin{cases} 
\frac{Age}{1.2 + 0.06} + \log(1.2 + 0.06) - \frac{1.2}{1.2 + 0.06}, & Age > 1.2 \\
\log(Age + 0.06), & Age \leq 1.2 
\end{cases}$$

This is similar to the age transformation described in the original Horvath pan-tissue human clock, but with offset at 0.06, and adult mouse age at 1.2.\textsuperscript{8} Following this, elastic net regression was implemented to regress the transformed chronological age on the CpG beta-values in the training set using the glmnet() function in R.\textsuperscript{58} The training set included the mouse samples reported in\textsuperscript{16,17} but excluded the BXDs. The training was therefore BXD unbiased. For pan-tissue clocks, all the samples were used, and for the liver specific clocks, the training was limited to liver samples. The alpha was set at 0.5, and the optimal lambda parameter was determined by 10-fold cross-validation (function cv.glmnet). This selected subsets of ‘clock
CpGs’ and coefficients (see Table S2 for clock CpG IDs, intercept, and coefficients). DNAmAge was then calculated as:

\[
\text{DNAmAge} = f^{-1}\left(\frac{b_0 + b_1 CPG_1 + b_2 CPG_2 + \ldots + b_i CPG_i}{b_0 + b_1 + b_2 + \ldots + b_i}\right)
\]

where \(b_0\) is the intercept, and \(b_1\) to \(b_i\) are the coefficients, and \(CPG_1\) to \(CPG_i\) denote the beta-values for the respective clock CpGs, and \(f^{-1}\) denotes the inverse function of \(f\).

### Statistics

Statistical analyses between the epigenetic predictors and continuous variables (body weight, strain lifespan) were based on Pearson correlations, and t-test was used to evaluate the effect of categorical predictors (sex, diet).

Two metabolic traits were downloaded from the bioinformatics platform GeneNetwork 2 (GN2) (1) fasted serum glucose, and (2) fasted serum total cholesterol (more information on how to retrieve these data directly from GN2 are provided in Data S14). Association with metabolic traits was examined using multivariable linear regression (the R equations are provided in Table S1). For visualization, residuals for both the predictor and outcome variables were extracted after regressing on age, diet, and BWF using the R code: residuals(lm( ~ age + diet + BWF)).

Longevity data (defined as age at natural death) was also downloaded from GN2 (Data S13). Males were excluded and strain-by-diet lifespan summary statistics were derived. Only strain-by-diet groups with 5 or more observations were included in the correlational analyses with the epigenetic predictors.

### Genetic analyses

The broad sense heritability within diet was estimated as the fraction of variability that was explained by background genotype\(^{20,60,61}\). For this, we applied a simple anova: aov(EAA ~ strain), and heritability was computed as: \(H^2 = SS_{strain}/(SS_{strain} + SS_{residual})\), where \(SS_{strain}\) is the strain sum of squares, and \(SS_{residual}\) is the residual sum of squares.

All QTL mapping was done on the GN2 platform, and these traits can be accessed from this website\(^{59}\) (trait accession IDs provided in Data S14). In the GN2 home page, the present set of BXD mice belongs to the Group: BXD NIA Longevity Study, and GN2 provides a direct interface to the genotype data. All QTL mapping was done for genotypes with minor allele frequency \(\geq 0.05\) using the genome-wide efficient mixed model association (GEMMA) algorithm\(^{31}\), which corrects for the BXD kinship matrix. For the EAA traits, diet, weight at 6 months, and final weight were fitted as cofactor. Chronological age had not correlation with EAA and this was not included as a cofactor (including age does not change the results). Genome-wide linkage statistics were downloaded for the full set of markers that were available from GN2 (3720 markers as of early 2021). For the combined p-values, QTL mapping was done separately using GEMMA for each EAA derived from all the unbiased mouse and universal clocks. Fisher’s p-value combination was then applied to get the meta-p-value\(^{32}\). We used this method to simply highlight loci that had consistent linkage across the different EAA measures. QTL mapping for entropy, major covariates—age, diet, BW1, and BWF—were included as co-factors. QTL
mapping for predicted-maxLS was done without co-factors (including variables such as age, weight, and diet did not change the results).

For marker specific linkage, we selected SNPs located at the peak QTL regions (DA0014408, rs48062674, rs30567369), and grouped the BXDs by their genotypes (F1 hybrids and other heterozygotes were excluded from this), and marker specific linkage was tested using ANOVA. rs48062674 and rs30567369 are reference variants that is already catalogued in dbSNP,\(^6^2\) and is used as a marker in the QTL mapping. DA0014408.4 is an updated variant at a recombinant region in the Chr11 interval and within the peak QTL interval.\(^2^0\) Genotypes at these markers for individual BXD samples are in Data S1.

For marker specific QTL analysis for EAA, we performed linear regression using the data in Data S1. Heterozygotes at the respective markers were excluded, and we applied the following regression model for each of the mouse and universal EAA traits separately: lm(EAA ~ genotype + diet). To test the effect on body weight change, body weight data measured at approximately 4 (baseline), 6, 12, 18, and 24 months were downloaded from GN2 (Data S14). Detailed description of these weight data are in Roy et al.\(^1^9\) We then applied a mixed effects regression model using the lme4 R package\(^6^3\): lmer(weight ~ age + diet + genotype + (1|ID)), where ID is the identifier for individual mouse.

**Bioinformatic tools for candidate gene selection**

Sequence variation between B6 and D2 in the QTL intervals (Chr11:90–99 Mb, and Chr19:35–48 Mb) were retrieved from the Wellcome Sanger Institute Mouse Genomes Project database (release 1505 for GRCh38/mm10).\(^6^4\)-\(^6^6\) Positional candidates were required to contain at least one coding variant (missense and/or nonsense variants), or have non-coding variants with evidence of cis-regulation in liver tissue of the BXDs. Cis-eQTLs for the candidate genes were obtained from the liver RNA-seq data described in \(^1^8\). An interface to search and analyze this transcriptome data is available from GN2, and is catalogued under Group: BXD NIA Longevity Study; Type: Liver mRNA; and Dataset: UTHSC BXD Liver RNA-seq (Oct 19) TMP Log2. This data was also used for the transcriptome-wide correlations analysis for univ.EAA in the 153 cases that had both DNAm and RNA-seq data. We considered the top 2000 highest correlated transcripts, and the list of transcripts were collapsed to a non-redundant list of gene symbols, and this was uploaded to the DAVID Bioinformatics Database (version 6.8) for GO enrichment analysis.\(^6^7\),\(^6^8\) Similarly, proteome correlational analysis was carried out using the data: Group: BXD NIA Longevity Study; Type: Liver Proteome; and Dataset: EPFL/ETHZ BXD Liver Proteome CD-HFD (Nov19).\(^1^8\).

For human GWAS annotations, we navigated to the corresponding syntenic regions on the human genome by using the coordinate conversion tool in the UCSC Genome Browser. The Chr11 90–95 Mb interval on the mouse reference genome (GRCh38/mm10) corresponds to human Chr17:50.14–55.75 Mb (GRCh38/hg38) (40.7% of bases; 100% span). The Chr11 95–99 Mb interval in the mouse corresponds to human Chr17:47.49–50.14 Mb (29.3% of bases, 57.9% span), and Chr17:38.19–40.39 Mb (20.7% of bases, 44.1% span). Likewise, for the Chr19 QTL, the mm10 35–40 Mb corresponds to hg38 Chr10:89.80–95.06 Mb (32.2% of bases, 89.2% span), 40–45 Mb corresponds to hg38 Chr10:95.23–100.98 Mb (46.6% of bases, 95.6% span), and 45–48 Mb corresponds to hg38 Chr10:100.98–104.41 Mb (46.5% of bases, 100% span). We then
downloaded the GWAS data for these regions from the NHGRI-EBI GWAS catalogue,\(^{33}\) and retained the GWAS hits that were related to aging.

**Data availability**

The full microarray data will be released via NCBI’s Gene Expression Omnibus upon official publication. Genome annotations of the CpGs can be found on Github https://github.com/shorvath/MammalianMethylationConsortium. Individual level BXD data are available on www.genenetwork.org on FAIR+ compliant format; data identifiers, and way to retrieve data are described in Data S14.

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**Author contributions.** KM contributed to the data, conceived portion of the study, and performed statistical analysis and drafted the article. ATL, CZL, AH contributed to the data analysis and in computing the epigenetic clocks and predictor. JVS contributed to the lab work. RWW conceived of the BXD Aging Colony, and provided access to the BXD biospecimen and data. SH developed the array platform, and built the epigenetic clocks and predictor. All authors contributed to, and approved the manuscript.

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**Competing interests.** SH is a founder of the non-profit Epigenetic Clock Development Foundation, which plans to license several of his patents from his employer UC Regents. The other authors declare no conflicts of interest.

**Ethics approval.** All animal procedures were in accordance to protocol approved by the Institutional Animal Care and Use Committee (IACUC) at the University of Tennessee Health Science Center.

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a CD: $r = -0.02$, ns  
HFD: $r = -0.16$, p = 0.09  

b CD: $r = -0.15$, p = 0.04  
HFD: $r = -0.23$, p = 0.01  

b CD: $r = -0.06$, ns  
HFD: $r = -0.16$, p = 0.09  

d CD: $r = 0.23$, p = 0.002  
HFD: $r = 0.1$, ns
a) Negative and positive mRNA correlates
- GO:0055114~oxidation-reduction process
- GO:0007049~cell cycle
- GO:0006955~immune response
- GO:0051301~cell division
- GO:0045087~innate immune response
- GO:0006954~inflammatory response
- GO:0002376~immune system process
- GO:007067~mitotic nuclear division
- GO:0056935~chemotaxis
- GO:0051607~defense response to virus
- GO:0050729~reg of inflammatory response

Fold enrichment

b) Negative protein correlates
- GO:0019373~epoxyxygenase P450
- GO:0006695~cholesterol biosynth
- GO:0016126~sterol biosynthetic
- GO:0006730~1-carbon metabolic
- GO:0008202~steroid metabolic
- GO:0006944~steroid biosynthetic
- GO:0006749~glutathione metabolic
- GO:0055114~oxidation-reduction
- GO:0008152~metabolic
- GO:0006629~lipid metabolic

Fold enrichment