Epigenetic clock and DNA methylation analysis of porcine models of aging and obesity

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Abstract DNA-methylation profiles have been used successfully to develop highly accurate biomarkers of age, epigenetic clocks, for many species. Using a custom methylation array, we generated DNA methylation data from \( n = 238 \) porcine tissues including blood, bladder, frontal cortex, kidney, liver, and lung, from domestic pigs (\( \textit{Sus scrofa domesticus} \)) and minipigs (Wisconsin Miniature Swine\textsuperscript{TM}). Samples used in this study originated from Large White \( \times \) Landrace crossbred pigs, Large White \( \times \) Minnesota minipig crossbred pigs, and Wisconsin Miniature Swine\textsuperscript{TM}. We present 4 epigenetic clocks for pigs that are distinguished by their compatibility with tissue type (pan-tissue and blood clock) and species (pig and human). Two dual-species human-pig pan-tissue clocks accurately measure chronological age and relative age, respectively. We also characterized CpGs that differ between minipigs and domestic pigs. Strikingly, several genes implicated by our epigenetic studies of minipig status overlap with genes (ADCY3, TFAP2B, SKOR1, and GPR61) implicated by genetic studies of body mass index in humans. In addition, CpGs with different levels of methylation between the two pig breeds were identified proximal to genes involved in blood LDL levels and cholesterol.

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synthesis, of particular interest given the minipig’s increased susceptibility to cardiovascular disease compared to domestic pigs. Thus, breed-specific differences of domestic and minipigs may potentially help to identify biological mechanisms underlying weight gain and aging-associated diseases. Our porcine clocks are expected to be useful for elucidating the role of epigenetics in aging and obesity, and the testing of anti-aging interventions.

**Keywords** Porcine · Pig · Minipigs · Aging · Development · Epigenetic clock · DNA methylation

**Introduction**

Pigs (*Sus scrofa*) are omnivores that last shared a common ancestor with humans between 79 and 97 million years ago [1, 2]. The domestication of pigs dates back to approximately 10,000 years, where they were domesticated from local wild boars across Eurasia [3, 4]. Since then, a wide variety of domestic and minipig breeds have been selectively bred for agricultural and biomedical purposes. While murine models have been traditionally used in biomedical research, there are added advantages to the use of porcine models in translational research. This includes their comparable size, anatomy, physiology, immunology, metabolism, and genetics with humans [5–7]. At the cellular levels, we have previously demonstrated similar genome-wide DNA methylation patterns between pigs and humans across a range of biomedically relevant tissues [8, 9], further supporting the high relevance of pigs in modeling human disease and development. Indeed, these advantages have been recognized and porcine models are already being used in biomedical research [10–19]. However, the large size and low propensity for atherosclerosis development in particular [20], limited the use domestic pigs as cardiovascular models. To overcome these issues, selectively bred and genetically modified minipigs have emerged over recent years as excellent models of hypercholesterolemia, atherosclerosis, metabolic syndrome, diabetes, and even cancer [13, 21–27]. Due to their smaller size, ease of handling, and genetic manipulability, minipigs are becoming increasingly important animal models for a wide range of human pathologies [12].

The study of any disease would be incomplete without an understanding of how age contributes to the malfunctioning of cells, tissues and organs. Despite this acknowledgement, the contribution of age to pathology has been largely unaddressed, not for the lack of will, but means. In the absence of an accurate way to quantify biological age, time (chronological age) is adopted as a surrogate that is manifestly unsatisfactory, as it remains unresponsive to biological fitness or frailty. The need for a measure of age that is based on biology is clear, and hints that this may be possible emerged when DNA methylation level was observed to change with advancing age. DNA methylation is an epigenetic modification that controls gene expression. The significance of its age-associated change was a subject of speculation until recently, when an array-based technology was developed to accurately measure its level on specific cytosine-phosphate-guanines (CpGs) in the genome. DNA methylation levels allow one to build age estimators (pan tissue clocks) that apply to most cells of the human body [28]. The rate of epigenetic aging was observed to be associated with a wide range of human conditions including mortality risk, Down syndrome, HIV infection and obesity [29–34], indicating that epigenetic age is a measure, at least to some degree, of biological age.

It is evident that the extrapolation of this epigenetic clock to other species, especially those such as pigs that are employed as disease models in biomedical research, will greatly facilitate research into the influence of age on pathology. This would also permit the
quantitative testing of interventions that could potentially mitigate age effects on pathology. Towards this end, we aimed to develop epigenetic clocks that apply to both humans and pigs at the same time. To overcome the species barrier, we used a DNA methylation array platform (HorvathMammalMethylChip40) that encompasses CpGs flanked by DNA sequences that are conserved across different species of the mammalian class.

Here, we present highly accurate epigenetic clocks that apply to both humans and different pig breeds: the regular sized domestic pigs, Wisconsin Miniature Swine™, and a cross between domestic and Minnesota minipigs. The human-pig clock provides increases the probability that findings in pigs will translate to humans, and vice versa. We also characterized (1) age-related changes in the porcine methylobe and (2) cytosines that differ between minipigs and regular sized pigs.

Methods

Porcine samples

All animal procedures were approved by the University of Illinois and University of Wisconsin Institutional Animal Care and Use Committee, and all animals received humane care according to the criteria outlined in the Guide for the Care and Use of Laboratory Animals. Porcine whole blood samples \(n=146\) were collected from female Large White X Landrace crossbred domestic pigs \(n=84\), age range 11–2285 days) and Wisconsin Miniature Swine™ \(n=60\), age range 8–1880 days) at the University of Wisconsin-Madison. Whole blood \(n=16\) and tissue samples (bladder, frontal cortex, kidney, liver, lung; \(n=16\)/tissue type) were collected from 16 Large White X Minnesota minipig crossbred pigs \(n=9\) female, \(n=8\) male, age range 29–1447 days) at the University of Illinois at Urbana-Champaign. All blood samples were collected in EDTA tubes, aliquoted, and flash frozen in liquid nitrogen within 10 min of collection. Tissue samples were collected and flash frozen within 10 min of euthanasia. All samples were stored at –80 until processing. Samples were shipped to the University of California, Los Angeles Technology Center for Genomics & Bioinformatics for DNA extraction and generation of DNA methylation data.

Human tissue samples

To build the human-pig clock, we analyzed previously generated methylation data from \(n=1211\) human tissue samples (adipose, blood, bone marrow, dermis, epidermis, heart, keratinocytes, fibroblasts, kidney, liver, lung, lymph node, muscle, pituitary, skin, spleen) from individuals whose ages ranged from 0 to 93. The tissue samples came from three sources. Tissue and organ samples from the National NeuroAIDS Tissue Consortium [35]. Blood samples from the Cape Town Adolescent Antiretroviral Cohort study [36]. Skin and other primary cells provided by Kenneth Raj [37]. Ethics approval (IRB#15–001,454, IRB#16–000,471, IRB#18–000,315, IRB#16–002,028).

DNA methylation data

All methylation data were generated DNA methylation data using the custom mammalian array “HorvathMammalMethylChip40” [38]. Not all CpGs on the array apply to pigs. The particular subset of species for each probe is provided in the chip manifest file at the NCBI Gene Expression Omnibus (GEO) platform (GPL28271). The SeSaMe normalization method was used to define beta values for each probe [39].

Relative age estimation

To introduce biological meaning into age estimates of pigs and humans that have very different lifespan, as well as to overcome the inevitable skewing due to unequal distribution of data points from pigs and humans across age range, relative age estimation was made using the following formula: Relative age = Age / maxLifespan, where the maximum lifespan for the two species was chosen from an updated version of the anAge database [40]. According to the "anAge" data base [40, 41], the maximum lifespan of Sus scrofa is 27 years since one specimen of the riukiuanus subspecies lived for 27 years in captivity [42]. In our models, we chose the more conservative value of 23 years as mentioned in the “Discussion” section.
Clocks and penalized regression

Details on the clocks (CpGs, genome coordinates) and R software code are provided in the Supplement. Penalized regression models were created with glmnet [43]. We investigated models produced by elastic net” regression (alpha = 0.5 in glmnet R function). The optimal penalty parameters in all cases were determined automatically by using a tenfold internal cross-validation (cv.glmnet) on the training set. By definition, the alpha value for the elastic net regression was set to 0.5 (midpoint between Ridge and Lasso type regression) and was not optimized for model performance. We performed a cross-validation scheme for arriving at unbiased (or at least less biased) estimates of the accuracy of the different DNAm based age estimators. One type consisted of leaving out a single sample (LOOCV) from the regression, predicting an age for that sample, and iterating over all samples. We subset the set of CpG probes to those that uniquely mapped to a CpG site in the swine genome. While no transformation was used for the blood clock for pigs, we did use a log linear transformation for the dual species clock of chronological age (Supplement). The accuracy of the resulting clocks was assessed via cross validation estimates of (1) the correlation $R$ between the predicted epigenetic age and the actual (chronological) age of the animal, and (2) the median absolute difference between DNAm age and chronological age (mae).

EWAS and functional enrichment

Epigenome-wide association studies (EWAS) was performed in each tissue separately (bladder, blood, cerebral cortex, kidney, liver, lung) using the R function “standardScreeningNumericTrait” from the “WGCNA” R package [44]. Next, the results were combined across tissues using Stouffer’s meta-analysis method. The functional enrichment analysis was done using the genomic region of enrichment annotation tool [45]. CpGs implicated by our EWAS were filtered for CpG position information, lifted over to the human genome using UCSC’s Liftover tool and fed into the online functional analysis tool GREAT using the default mode, to obtain a list of significantly enriched functions for both positive and negative EWAS hits in the different tissues.

Transcription factor enrichment and chromatin states

The FIMO (Find Individual Motif Occurrences) program scans a set of sequences for matches of known motifs, treating each motif independently [46]. We ran TF motif (FIMO) scans of all probes on the HorvathMammalMethyl40 chip using motif models from TRANSFAC, UniPROBE, Taipale, Taipaledimer, and JASPAR databases. A FIMO scan $p$ value of $1E-4$ was chosen as the cutoff (lower FIMO $p$ values reflect a higher probability for the local DNA sequence matching a given TF motif model). This cutoff implies that we would find almost all TF motif matches that could possibly be associated with each site, resulting in an abundance of TF motif matches. We caution the reader that our hypergeometric test enrichment analysis did not adjust for CG content.

Results

In total, we analyzed 238 tissue samples mainly from blood (Table 1). Blood samples were obtained from three pig lines: a cross between the Large White and Landrace domestic pig breeds, the Wisconsin Miniature Swine minipig, and a cross between the Large White domestic breed (maternal line) and the Minnesota minipig (sire). From the latter pig line (i.e., the domestic minipig cross), methylation profiles were obtained from DNA isolated from bladder, brain (frontal cortex), kidney, liver, and lung tissue. Unsupervised hierarchical clustering revealed that the samples clustered by tissue type (Supplementary Fig. 1). Random forest predictors were fitted to three different outcomes: (1) pig breed (Sus scrofa domesticus versus Sus scrofa minusculus), (2) tissue type, and (3) sex. The three classifiers exhibited perfect accuracy, with respective (out-of-bag) error rates of zero.

Predictive accuracy of the epigenetic clock

To arrive at unbiased estimates of the epigenetic clocks, we applied cross-validation analysis with the training data. For the development of the basic pig clock, this consisted of pig blood, bladder, frontal cortex, kidney, liver, and lung DNA methylation profiles. For the generation of human-pig clocks however, the training data was constituted by human and pig DNA methylation profiles. Cross-validation
analysis reports unbiased estimates of the age correlation $R$ (defined as Pearson correlation between the age estimate (DNAm age) and chronological age) as well as the median absolute error.

From these analyses, we developed four epigenetic clocks for pigs that vary with regard to two features: species and measure of age. The resulting two human-pig clocks mutually differ by way of age measurement. One estimates chronological ages of pigs and humans (in units of years) based on methylation profiles of 638 CpGs, while the other employs the methylation profiles of 542 CpGs to estimate relative age, which is the ratio of chronological age of an animal to the maximum lifespan of its species; with resulting values between 0 and 1. This relative age ratio is highly advantageous because it allows alignment and biologically meaningful comparison between species with very different lifespans such as pig and human, which cannot otherwise be afforded by direct comparison of their chronological ages.

As indicated by its name, the pure pig clock, constituted by 120 CpGs, is highly accurate in age estimation in all porcine tissues ($R=0.97$ and median error 0.22 years, Fig. 1A). The pan-tissue clocks exhibit high age correlations in individual porcine tissues ($R>0.90$, Supplementary Fig. 1). The human-pig clock for chronological age is highly accurate when DNA methylation profiles of both species are analyzed together ($R=0.98$, Fig. 1B), and remains so when restricted to pig tissue samples ($R=0.97$, Fig. 1C). Similarly, the human-pig clock for relative age exhibits high correlation regardless of whether the analysis is applied to samples from both species ($R=0.98$, Fig. 1D) or only to pig samples ($R=0.96$, Fig. 1E). The use of relative age circumvents the clustering of data points of pigs and humans to opposite parts of the curve, which is evident in Fig. 1C. These highly accurate array of porcine clocks are readily usable with immediate effect in porcine models of diseases and conditions, and the human-pig clock of relative age is particularly exciting as it allows comparison between human and pigs based on their relative positions within the lifespans of both species.

EWAS of chronological age

Although several hundred CpGs were used to construct the epigenetic clocks described above, these were merely a subset of all CpGs, which changed with advancing age. There are many more age-associated CpGs that are not used for the purpose of estimating porcine age, but are nevertheless very important when we seek to identify CpGs with methylation levels that are associated with age through EWAS. In total, 34,540 probes from the HorvathMammalMethylChip40 are aligned to loci that are proximal to 5,209 genes in the \textit{Sus\_scrofa.Sscrofa11.1.100} genome assembly. Due to the high inter-species conservation of the probes on the array, findings from the pig methylation data can probably be extrapolated to humans and other mammalian species. EWAS of chronological age revealed clear tissue-specific DNAm change in pigs (Fig. 2A). Age-associated CpGs in one tissue tend to be poorly conserved in another tissue (Supplementary Fig. 3). However, the poor conservation and differences in $p$ value ranges in our analyzed tissue types may reflect the limited sample size in non-blood tissues.
A nominal $p$ value $< 10^{-4}$ was set as the cutoff for significance. The top age-associated CpGs and their proximal genes for the individual tissues are as follows: bladder, $ANO4$ exon ($z=5.8$); blood, $EN1$...
promoter (z = 26); frontal cortex, FGF9 exon (z = 5.6); kidney, TNRC6A exon (z = −7.5); liver, NRSAl exon (z = 8.6); and lung, UNC79 5′UTR (z = 8.1). Despite poor conservation across tissues, there are nonetheless age-related CpGs that are common to all tissues, and these were identified through meta-analysis of these six tissue samples to be hypermethylation in EN1 promoter (z = 18), HCN1 exon (z = 17), UNC79 5′UTR (z = 17), LHFPL4 exon (z = 17), and NR2F2 exon (z = 17). The upset plot analysis identified several CpGs with conserved DNA methylation patterns in at least four pig tissues (Fig. 2B). The most conserved DNA methylation pattern was hypermethylation of the SP8 promoter in all tissues, with the exception of the brain. Genes whose expressions are regulated by SP family transcriptional factors are essential for proper limb development. Aging-associated CpGs in different tissues were distributed in genic and intergenic regions that are defined relative to transcriptional start sites (Fig. 2C). There is little enrichment of age-related CpGs in any genetic features across all tissues, with the exception of consistent hypermethylations in promoters and 5′UTRs. This result is consistent with a higher positive association of CpG island methylation with age than non-island CpGs in all tissues (Fig. 2D). These features suggest that a substantial amount of age-associated CpGs are likely to impact gene expression.

To specifically explore the potential impact of age-related porcine CpG methylation on gene expression, we analyzed putative transcription factors binding sites for such methylation. From these we identified 20 transcription factor binding motifs that exhibit age-related methylation changes (Fig. 3A). The corresponding transcription factors control the expression of genes which are involved in many different cellular activities. For example, hypomethylation in the SP1 motif in blood and cortex indicates greater access of the SP1 protein to some of its binding sites with increasing age. However, the outcome of this is difficult to predict as SP1 activates the transcription of many genes that are involved in diverse cellular processes ranging from cell growth, apoptosis, the immune response and chromatin remodeling. The challenge in predicting downstream events from single transcription motifs can be partly addressed by collective analysis of multiple transcription factor motifs. Such an analysis identified age-associated methylation changes for SMAD3, SP1, SP3, and E2F1 transcription factor binding motifs, which are implicated in telomerase regulation (p = 6E−7). We briefly mention that the effects of telomere length and telomerase activity in porcine tissues appears to be comparable with that of humans [47].

While identification of genes proximal to age-associated CpGs, as performed above, are useful, the likelihood of their potential effects on cells is difficult to gauge. This can be partly addressed by carrying out analysis to identify enrichment of implicated genes in specific pathways, pathologies and biological processes (Fig. 3). This analysis highlighted the following features impacted by age-related CpG methylation changes in porcine cells: organism development, the nervous system and metabolism; all of which have also been found to be implicated in epigenetic aging of humans and other species (Fig. 3B). Furthermore, genes proximal to hypermethylated CpGs are associated with H3K27Me3 marks and are often polycomb protein EED targets in porcine tissues. EED is a member of the multimeric Polycomb family protein complex that maintains the transcriptional repressive states of genes. These proteins also regulate H3K27Me3 marks, DNA damage, and senescence states of the cells during aging [48].

Studying differences across pig lines

With the DNA methylation datasets we have accrued from the different porcine breeds, we are in a position to identify CpGs that differ between domestic and minipig breeds. We compared DNA methylation profiles from blood of domestic pigs with those from Wisconsin Miniature Swine™. We found the mean methylation across CpGs located in CpG islands is higher in minipigs than in the other two pig lines (Supplementary Fig. 4). Similarly, the average rate of change in methylation across island CpGs is increased in minipigs (Supplementary Fig. 4).

We analyzed individual CpGs using two different multivariate models. In the first, the DNA methylation levels of a given CpG were regressed on age and breed (minipig versus reset) to identify CpGs that are associated with aging in both breeds (age main effect). This model was also used to identify CpGs with significantly different basal methylation levels between breeds (Minipig main effect). The second model identified age-related CpGs with different
EWAS of chronological age

(A) Bladder
- ANO4
- EN1
- FGFBP4
- PAX6
- FOXD3
- NQO1
- OCA2

(Blood)
- EN1
- FGFBP4
- PAX6
- FOXD3
- NQO1
- OCA2

(C) Frontal Cortex
- OSR2
- FGF9
- HOXC5
- ETV2
- HOXA3
- ETH1
- EN1

(Kidney)
- TNRC6A
- SYNGAP1
- RAB11FIP4
- PABPC4
- MAPKAP1
- RASBN1
- PAX8
- VRK2
- ATXN7
- SRED2
- RNF144A

Direction of association
- Red: Hypermethylated
- Blue: Hypomethylated

CpG count

Upset plot of the significant CpGs

D) Associations with chronological age

Bladder
- ANO4
- EN1
- FGFBP4

Blood
- FGFFBP4
- PAX6
- FOXD3
- NQO1
- OCA2

Frontal Cortex
- OSR2
- FGF9
- HOXC5
- ETV2
- HOXA3
- ETH1

Kidney
- TNRC6A
- SYNGAP1
- RAB11FIP4
- PABPC4
- MAPKAP1
- RASBN1
- PAX8
- VRK2
- ATXN7
- SRED2
- RNF144A

Intergenic upstream
- Promoter
- Gene body
- LTR
- threcUTR
- threcUTR

Intergenic downstream
- Intergenic upstream
- Intergenic downstream

CpG location relative to closest TSS

Tissue
- Bladder
- Blood
- Frontal Cortex
- Kidney
- Liver
- Lung

Meta analysis

CpG count

T-test, p = 2.2e-1
rates of methylation changes between these two pig breeds independent of the direction of change. At a genome-wide significance set at $p < 1 \times 10^{-8}$, 10,167 age-associated CpGs were shared between both pig breeds, while 825 CpGs had different baseline methylation levels, and the rate of methylation change of 32 age-associated CpGs were significantly different between the two (Fig. 4A). Thus, while age has the largest impact on methylation of these CpGs, there is an inherent species-specific difference in basal methylation levels of a substantial number of CpGs, which is expected given the overt differences between the breeds. The top CpGs with divergent rate of methylation change between the breeds are proximal to the $MGST1$ exon, $SON$ 3'UTR, and $TFAP2B$ exon (Fig. 4A–C). CpGs with different basal methylation levels between the two breeds were located within the $OSBP$ exon, $ENCI$ intron, and $CTNNBL1$ upstream regions (Fig. 4A). In total, eight categories of CpGs can be defined based on the direction of their methylation change with age in the two pig breeds. While methylation of most age-associated CpGs was altered in similar directions in both breeds (hypo or hyper in both axes), some were clearly in opposite directions (hyper in one axis and hypo in the other). The $LMNA$ intron is an example region that displayed extreme divergence. While its intron was hypomethylated in age in the domestic pig, it was hypermethylated in minipigs (Fig. 4C). An enrichment analysis of age-related CpGs that are exclusive to either breed implicated pathways involved in development, survival, cancer, and growth (Fig. 4D).

Our analysis of differentially methylated CpGs between domestic and minipigs implicated genes that regulate weight. This particularly interesting finding prompted us to query whether any of these identified genes are associated with weight differences in humans. To this end, we overlapped the EWAS genes with a large GWAS meta-analysis of body-mass index (BMI) in humans, which included 681,275 participants in UK Biobank and GIANT_BMI consortium [49]. Strikingly, several of the EWAS-identified genes had genetic variants that are associated with increased BMI in humans (Fig. 5). The top genes include $ADCY3$, $TFAP2B$, $SKOR1$, and $GPR61$, which had numerous SNPs in different gene regions associated with human BMI. Thus, our EWAS analysis of pig breeds of overtly different sizes implicates genes that were implicated by a published GWAS of human BMI.

**Discussion**

We have previously developed several human epigenetic clocks from DNA methylation profiles that were derived from various versions of human Illumina DNA methylation arrays. As these arrays are specific to the human genome, a critical step toward crossing the species barrier was the development and use of a mammalian DNA methylation array that profiles 36,000 CpGs with flanking DNA sequence that are highly conserved across numerous mammalian species. The employment of this array to profile 238 porcine tissues from 3 pig lines represents the most comprehensive epigenetic dataset of domestic pigs thus far. These data allowed us to construct four highly accurate DNA methylation-based age estimators for three pig lines. Two of these clocks apply only to pigs: the pan-tissue clock which was trained on methylation profiles of six tissues is expected to apply to most pig tissues. The blood clock on the other hand, was trained using only blood DNA methylation data. These two highly accurate porcine clocks are readily and easily included in porcine-based models of diseases and health conditions. This will encourage investigations into the relationship between age and
A

Enrichment of TFs in CpGs located in the promoter

| Gene Symbol | Blood | Frontal Cortex | Kidney | Liver | Liver | meta |
|-------------|-------|----------------|--------|-------|-------|------|
| V_SP1SP3_Q4 | hyper | hypo           | hypo   | hypo  | hypo  | hyper |
| V_SP1_Q6   |       |                |        | hypo  | hypo  |      |
| V_SP1_Q4_01|       |                |        | hypo  | hypo  |      |
| V_RNF96_01 |       |                |        | hypo  | hypo  |      |
| V_AP2_Q6_01|       |                |        | hypo  | hypo  |      |
| Smad3_secondary |      |                |        |       |       |      |
| MA0073.1-RREB1 |   |                |        |       |       |      |
| MA0068.1-Pax4  |   |                |        |       |       |      |
| MA0048.1-NHLH1 |   |                |        |       |       |      |
| GCM1_FIGLA_GCM_bHLH_386 |   |                |        |       |       |      |
| E2F1_EOMES_E2F_Tbox_330 |   |                |        |       |       |      |

B

Enrichment for GO phenotypes, upstream regulators

| Gene Symbol | Blood | Liver | Lung | meta |
|-------------|-------|-------|------|------|
| BENPORATH_ES_WITH_H3K27ME3                          |       |       |      |      |
| BENPORATH_EED_TARGETS                                |       |       |      |      |
| DNA binding                                          |       |       |      |      |
| single-multicellular organism process                |       |       |      |      |
| multicellular organismal process                     |       |       |      |      |
| multicellular organismal development                 |       |       |      |      |
| anatomical structure development                     |       |       |      |      |
| sequence-specific DNA binding                        |       |       |      |      |
| transcription factor activity                        |       |       |      |      |
| nucleic acid binding                                 |       |       |      |      |
| transcription factor activity                        |       |       |      |      |
| nervous system phenotype                             |       |       |      |      |
| sequence-specific DNA binding                        |       |       |      |      |
| abnormal nervous system morphology                  |       |       |      |      |
| decreased brain size                                 |       |       |      |      |
| BENPORATH_SUZ12_TARGETS                              |       |       |      |      |
| lethality during fetal growth through weaning        |       |       |      |      |
| BENPORATH_PRC2_TARGETS                               |       |       |      |      |
| abnormal somatic nervous system morphology          |       |       |      |      |
| organ development                                   |       |       |      |      |
| V$CEBPA_01                                           |       |       |      |      |
| abnormal neuron morphology                           |       |       |      |      |
| regionalization                                      |       |       |      |      |
| abnormal olfactory lobe morphology                   |       |       |      |      |
| MIKKELEN_IPS_WITH_HCP_H3K27ME3                       |       |       |      |      |
| perinatal lethality                                  |       |       |      |      |
| small olfactory bulb                                 |       |       |      |      |
| complete perinatal lethality                         |       |       |      |      |
| CTTTAAR_UNKNOWN                                      |       |       |      |      |
| abnormal olfactory bulb development                 |       |       |      |      |
| abnormal neuron proliferation                        |       |       |      |      |
| organ morphogenesis                                  |       |       |      |      |
| dorsal/ventral pattern formation                     |       |       |      |      |
| V$GFI1_01                                            |       |       |      |      |
| V$NIKX22_01                                          |       |       |      |      |
| truncation of digits                                 |       |       |      |      |
| ectopic dopaminergic neuron                          |       |       |      |      |
| absent cerebellar lobules                            |       |       |      |      |
| abnormal digit pigmentation                          |       |       |      |      |
of life. In this study the highest scoring porcine blood CpG is one that resides in the promoter of the engrailed gene (EN1), which is a member of the Hox gene family. EN1 also scored the highest in meta-analyses of all the tissues. It is as yet unclear how methylation of these loci are involved with aging, but their increased methylation with age hints at the possible role of cellular identity and differentiation. This is no doubt an exciting avenue of exploration in which these epigenetic clocks will be essential.

The considerable similarities between humans and pigs allows for the testing of age-mitigation interventions as well as factors that impact longevity in pigs, which have a shorter lifespan and are much more amenable subjects for controlled trials. To accurately translate age-related findings from pigs to humans however requires a correct and accurate measure of age-equivalence. We fulfilled this need through a two-step process. The first, which we described above is the generation of dual-species clocks (human-pig), one of which is as accurate in estimating pig age as it is for human age; in chronological units of years. The second is the expression of pig and human ages in respect to the maximum recorded ages of their respective species (species lifespan), i.e. 23 years for pigs and 122 years for humans. It is difficult to find reliable lifespan data for domestic pigs.

According to the “anAge” database [40, 41], the maximum lifespan of Sus scrofa is 27 years since one specimen of the riuikuianus subspecies lived for 27 years in captivity [42]. In our models, we chose the more conservative value of 23 years after consulting with experts. We are not sure whether minipigs live longer lives than regular sized domestic pigs. If in fact pigs behave the same as dogs (and several other species), one would expect domestic pigs to live considerably less long than minipigs. Minipigs are expected to live between 12 and 18 years but precise estimates of maximum lifespan appear to be unknown [50, 51]. In our own database, we assigned the Wisconsin Miniature Swine™ the Latin name “Sus scrofa minusculus” with an estimated maximum lifespan of 23 years. Domestic pigs experience a remarkably fast growth period which might influence their lifespan, their aging process, and mean methylation levels. This might explain why the mean methylation across CpGs located in CpG islands is higher in minipigs than in the other two pig lines (Supplementary Fig. 4).
Our study is limited in that all animals were younger than 6.3 years old while the maximum observed lifespan of domestic pigs (Sus scrofa domesticus) appears to be 23. We hypothesize that the lack of training data on older animals resulted in porcine clocks that will under-estimate the ages of older animals. However, we expect that the correlation between age and its DNA methylation based estimate will continue to be high in older animals. Further, we hypothesize that CpGs that correlate with age in
younger animals also correlate with age in older animals. These hypotheses derive from our studies of humans and many other species where these hypotheses have been validated [31, 52].

The mathematical operation of generating a ratio eliminates chronological units of time and produces a value that indicates the age of the organism in respect to the maximum age of its own species. This allows a meaningful and realistic cross-species comparison of biological age. For example, the biological fitness of a 20-year-old pig, which is very old, is not equivalent to that of a 20-year-old human, who is young. However, a pig with a relative epigenetic age of 0.5 is more comparable to a human of similar relative epigenetic age. Collectively, the ability to use a single mathematical formula to measure epigenetic age of different species and the replacement of chronological units of time with proportion of lifespan, are two significant innovations that will propel cross-species research as well as cross-species benefits.

In addition to age-related epigenetic changes, we also compared DNA methylation profiles between domestic and minipigs. CpGs with substantially

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**Fig. 4** EWAS of multivariate regression models for pig breed and age. The figure visualizes the results of 2 different linear regression models which used individual CpGs as dependent variable. The 2 linear models differ in terms of the underlying covariates: two covariates (age and pig breed) or three covariates (age, pig breed, and interaction effect). A Manhattan plots of DNAm aging loci that are shared between minipigs and domestic pigs (aging main effect), basal breed differences in DNAm levels (minipig main effect), and the interaction of breed and aging, which represent the loci with a divergent DNAm aging pattern between minipigs and domestic pigs. The analysis is done by multivariate regression models with or without (to estimate the main effect) interaction term for age and breeds. For breeds, the domestic pig is the reference variable to estimate the direction of change. Sample sizes: domestic pigs, 98; minipigs, 60. The red line in the Manhattan plot indicates \( p < 1 \times 10^{-35} \). B Scatter plots of DNAm aging between minipigs and domestic pigs. The highlighted CpGs are the loci with significant DNAm aging interaction between breeds at a 5% FDR rate. In total, eight categories of interaction were defined based on the aging Z-statistic of each breed. C Age versus methylation levels for select CpGs with significant interaction terms between breed and age. D Enrichment analysis of the genes proximate to CpGs related to age (shared between breeds), minipig, and age/minipig interaction. The gene-level enrichment was done using GREAT analysis [45] and human Hg19 background. The background probes were limited to 23,087 probes that were mapped to the same gene in the pig genome. The top CpGs were selected at a \( p < 1 \times 10^{-5} \) and based on beta values of association for up to 500 in a positive or negative sign.

**Fig. 5** Overlapping EWAS results in pigs with GWAS results in humans. EWAS of aging, minipig, and aging/pig breed interaction identifies genes with genetic variants associated with human body mass index. The GWAS is based on summary statistics of BMI meta-analysis of 681,275 participants in the UK Biobank and GIANTBMI consortium [49]. The coloring is based on the genes identified by one or multiple EWAS of age, minipig, and age/minipig interaction (inter). The labels indicate the top SNP from each of the top 30 gene regions.

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different basal methylation levels between the two breeds were identified proximal to the OSBP gene. This is of great significance, as OSBP is a transport protein that translocates sterols from lysosomes into the nucleus where the sterol represses the expression of the LDL receptor gene, HMG-CoA reductase gene and HMG synthase gene. The reduction of LDL receptor increases the risk of atherosclerosis and cardiovascular disease. Indeed, mice with ablated LDL receptor genes develop plaques in their aortas, while wild-type mice are free of such plaques. Compounding the effects on LDL receptor levels is the effect on expression of HMG-CoA and HMG synthase, which are members of the intracellular pathway that synthesize cholesterol. These features are of particular interest given the fact that while domestic pigs are refractive to atherosclerotic plaque development, minipigs are susceptible and hence used as models for cardiovascular disease. Another feature of interest is the identification of differentially methylated CpGs associated with genes involved in the development of obesity. The matching of EWAS targets with a large GWAS meta-analysis of body mass index (BMI) in humans led to the identification of overlapping genes included TFAP2B, which influences the effect of dietary fat on weight [53], GPR61, which is involved in the regulation of food intake and body weight [54, 55], and ADCY3 and SKOR1, both of which are associated with obesity and BMI [56–59]. Collectively, these associations points to the contribution of epigenetic control and influence on weight gain and obesity, in addition to highlighting the translational relevance of porcine models for cardiovascular disease and obesity-related research. It is interesting to note that the Dnmt3a locus is also differentially methylated between these two breeds. It is tempting to speculate that this may be an upstream event that impacts on the downstream methylation differences described above.

As porcine biomedical models for a wide range of age-related disorders are currently in use or being developed, including Alzheimer’s disease [60], cardiovascular disease [61], diabetes [22], and cancer models [21, 23, 24], the availability of these epigenetic clocks will extend the use of porcine models for aging, and possibly obesity studies.

To rigorously evaluate anti-aging treatment effects, we recommend to randomize treatment assignment and to collect several tissue types, e.g., skin samples (ear punches) and blood samples since treatment effects can be tissue/cell type specific as is the case in menopausal hormone therapy [62]. To average out technical variability in DNAm age estimates, we recommend to collect at least two tissue samples before treatment and at least two samples after treatment per animal.

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Author contribution SH conceived of the study and wrote the article. The remaining authors helped with the statistical analysis or the data generation. All authors reviewed and edited the article.

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Declarations

Conflict of interest 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.

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