Pig genome functional annotation enhances biological interpretations of complex traits and comparative epigenomics

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Abstract

The functional annotation of livestock genomes is crucial for understanding the molecular mechanisms that underpin complex traits of economic importance, adaptive evolution and comparative genomics. Here, we provide the most comprehensive catalogue to date of regulatory elements in the pig (Sus scrofa) by integrating 223 epigenomic and transcriptomic data sets, representing 14 biologically important tissues. We systematically describe the dynamic epigenetic landscape across tissues by functionally annotating 15 different chromatin states and defining their tissue-specific regulatory activities. We demonstrate that genomic variants associated with complex traits and adaptive evolution in pig are significantly enriched in active promoters and enhancers. Furthermore, we reveal distinct tissue-specific regulatory selection between Asian and European pig domestication processes. Compared with human and mouse epigenomes, we show that porcine regulatory elements are more conserved in DNA sequence, under both rapid and slow evolution, than those under neutral evolution across pig, mouse, and human. Finally, we provide novel biological insights on tissue-specific regulatory conservation and demonstrate that, depending on the traits, mouse or pig might be more appropriate biomedical models for different complex traits and diseases in humans through integrating comparative epigenomes with 47 human genome-wide association studies.
Functional elements play essential roles in regulating gene expression in living cells and tissues\(^1\). There have been great efforts on the identification and annotation of functional elements in human and mouse genomes\(^1\)\textsuperscript{-11} as well as other model organisms including *Drosophila*\(^12\) and *C. elegans*\(^13\). Significant enrichment of variants associated with human complex traits within regulatory elements has demonstrated the importance of Encyclopedia of DNA Elements (ENCODE) data\(^14\). Comparative analysis of epigenomes and transcriptomes across species could provide novel insights into the interrogation of underlying molecular mechanisms of human disease\(^8,15\). Genetic variants associated with common illnesses are enriched in human orthologues of mouse regulatory elements identified by ENCODE\(^3\), which validates the mouse could serve as a biomedical model for understanding some human diseases. However, compared with the mouse, pig (*Sus scrofa*) has more anatomical and physiological similarities to humans\(^16\)\textsuperscript{-18}, and has been widely used as a human medical model\(^16,17,19\)\textsuperscript{-21}. The pig is also one of the most important farm animal species for meat production worldwide\(^22\). The genetic improvement of economically important complex traits such as growth, feed efficiency, and health contribute to efficient and sustainable production of animal protein ensuring a secure food supply for a growing world population. Functional annotation of regulatory elements in pig will lay a solid foundation for the identification of causative variants associated with phenotypic variations of complex traits, due to the enrichment of these variants in regulatory regions and the small genetic effect of each variant, making them difficult to discover\(^23\).

Following ENCODE and Roadmap Epigenomics projects\(^8\), the Functional Annotation of Animal Genomes (FAANG) initiative\(^24\), although still in its infancy, has made great progress
towards annotating functional elements in many tissues across multiple domestic species including pigs\textsuperscript{25-30}. Here, we present 95 new genome-wide sequencing datasets from six gut-associated porcine tissues and integrate them with 128 previously published FAANG datasets from eight biologically distinct tissues. The collective interpretation of these datasets yields the most comprehensive annotation of functional elements to date in any domesticated animal species. In addition, we find that tissue-specific regulatory elements were enriched for the potential causative variants of complex phenotypes by integrating a large scale of genome-wide association studies (GWAS) and expression QTL (eQTL) datasets. Furthermore, by combining selection signature of the pig genome, we show that tissue-specific regulatory elements play an important role during the domestication. Finally, we compared porcine functional annotations with complementary datasets from the human and mouse and integrated GWAS datasets concerning 47 human complex traits. These comparisons demonstrate conservation of tissue-specific epigenetic signatures, suggesting that, depends on diseases, pig or mouse may be a better animal model for them than the other one.

Results

Data summary

We integrated 223 genome-wide sequencing datasets from 14 major tissues in pig (Fig. 1a), representing four histone modifications (H3K4me3, H3K4me1, H3K27ac and H3K27me3) measured by Chromatin Immunoprecipitation sequencing (ChIP-seq), chromatin accessibility by the Assay for Transposase-Accessible Chromatin (ATAC-seq), DNA methylation by Reduced Representation Bisulfite sequencing (RRBS), and gene expression by RNA-seq (Supplementary Fig. 1). We produced nearly 9 billion mapped reads with an average remaining rate of 68.81\%
across samples (Supplementary Table 1). Among 14 tissues, we obtained an average of 32,387, 106,849, 72,252, 98,721, and 122,585 peaks for H3K4me3, H3K4me1, H3K27ac, H3K27me3, and ATAC, covering 1.56, 2.78, 2.37, 7.74, and 3.31% of the entire genome, respectively (Fig. 1b, c). Additionally, we utilized 16 CTCF ChIP-seq datasets from eight tissues and four Hi-C datasets from liver to identify topologically associating domains (TADs) and Hi-C loop for associating regulatory elements (enhancers) with potential target genes.

The hierarchical clustering of samples based on the signal intensity of epigenetic marks and gene expression profiles clearly recapitulated sequencing assays, followed by tissue types and biological replicates (Fig. 1d), which was consistent with results of principal component analysis (PCA) (Supplementary Fig. 2). The six assays formed three major clusters: (1) active regulatory regions (H3K4me3, H3K27ac, H3K4me1 and ATAC), (2) Polycomb repression (H3K27me3), and (3) gene expression (RNA-seq). The four active regulatory marks were positively correlated with each other, but were negatively correlated with H3K27me3, especially H3K27ac. The signal intensity of RNA-seq (within gene bodies) showed a weakly positive correlation with active regulatory marks, and a negative correlation with H3K27me3. Overall, all four active regulatory marks exhibited significant enrichments in the upstream of transcription start sites (TSS) of genes across tissues (Fig. 1e).

We present Myosin IA (MYO1A) gene, which is engaged in responses to *Escherichia coli* infection and microvillar membrane morphology in intestinal tissues, as an example to show the complex interplays of regulatory elements and gene expression in Figure 1f. MYO1A is specifically and highly expressed in intestine tissues, and showed specific enrichments of H3K27ac signals around its TSS in intestine tissues but not in other tissues (Fig. 1f). In addition,
the TSS of MYO1A is accessible and was enriched for other active regulatory marks (i.e., H3K27ac, H3K4me3 and H3K4me1) but not for Polycomb repression (H3K27me3) (Fig. 1f).

**Prediction and characterization of chromatin states across 14 tissues**

We defined 15 distinct chromatin states by combining all five epigenetic marks across 14 tissues. These states mainly represented promoters (TssA, TssAHet and TssBiv, covering 1.16% of the entire genome), TSS-proximal transcribed regions (TxFlnk, TxFlnkWk and TxFlnkHet, covering 0.92% of the genome), enhancers (EnhA, EnhAMe, EnhAWk, EnhAHet and EnhPois, covering 6.5% of the genome), repressed regions (Repr and ReprWk, covering 13.25% of the genome) and quiescent regions (73.39%) (Fig. 2a–e, Supplementary Table 2). Totally we identified 2,097,958 regulatory elements (exclude Qui) spanning 14 tissues including 39,351 active promoters (TssA), 188,827 active strong enhancers (EnhA), and 142,821 repressors (Repr) (Supplementary Fig. 4a-c). On average, 4.79% of the genome was accessible but did not coincide any other measured epigenetic marks (ATAC islands), indicating that additional epigenetic marks are required to further explore the biological function of such regions. TssA and TssBiv showed the highest enrichment of conserved DNA sequence elements, followed by TSS-proximal transcribed regions and accessible enhancers (EnhA and EnhAMe) (Fig. 2f). In general, TssA and TssBiv showed the highest enrichment at TSS, while other chromatin states showed enrichment at up- and down-stream of TSS (Fig. 2g). For instance, TssAHet and TSS-proximal transcribed states had the highest enrichment around 2kb up-stream of TSS, whereas enhancer states showed the highest enrichment around 20kb up-stream of TSS. Repressed states were enriched around 20kb up- and down-stream of TSS (Fig. 2g).

In general, different chromatin states showed distinct DNA methylation levels (Fig. 2h). Promoter and TSS-proximal transcribed states were hypomethylated compared to nearby sequence
(10kb up- and down-stream of TSS). Among promoter states, TssA had the lowest methylation level, confirming the well-known negative correlation between promoter methylation and gene expression\textsuperscript{33}. The enhancer states showed intermediate methylation levels, among which EnhA and EnhAME had lower methylation levels compared to other enhancers (Fig. 2g), where we also observed that EnhA and EnhAME had more conserved sequence than other enhancers (Fig. 2f). This result suggests accessible enhancers may have more conserved sequences than non-accessible enhancers.

We took chromosome 7 (Chr7) as an example to explore the relationships among chromatin states, individual epigenetic marks, gene density, gene expression, DNA methylation and chromatin conformation (Fig. 2i). For instance, we observed that regions with higher density of genes were characterized by active chromatin states, higher gene expression, more chromatin accessibility, and lower methylation level. The chromatins were more physically interacted (measured by TADs from Hi-C data) within both gene desert and gene rich regions than the rest of genomic regions. To examine the associations of chromatin states with gene expression across tissues, we presented the \textit{VILI} locus (Villin-1), which participates in response to intestinal inflammation\textsuperscript{34}, as an example (Fig. 2j). \textit{VILI} exhibited tissue-specific active promoters and enhancers, as well as high expression in intestinal tissues compared to other tissues. Of particular note, despite the presence of TssA, \textit{VILI} was not expressed in stomach, possible due to the lack of enhancer activity in the upstream of its TSS, indicating that enhancers together with promoters may collectively regulate gene expression. Similar patterns were observed for \textit{MYO1A} and \textit{HNF4G} (Supplementary Fig. 3).

\textbf{Dynamics of chromatin states across genome and tissues}
We clustered the entire genome into 12 modules based on their relative frequency of chromatin states and observed that these modules exhibited distinct enrichments for protein-coding genes, non-coding genes and CpG islands (Fig. 3a). For instance, module 2 (M2) was characterized by active promoters and accessible enhancers, showed the highest enrichment for genes and CpG islands, and the lowest levels of DNA methylation, and the highest gene expression levels (Fig. 3b). Compared to modules 11 and 12, module 10 showed similar enrichment for Polycomb repression but higher enrichment for TssBiv, in which genes exhibited significantly lower expression levels, suggesting the crucial role of TssBiv for regulating gene repression (Fig. 3b). In addition, we noticed that module 1 had high enrichment for TssAHet, high levels of DNA methylation, and high representation of genes located on the X chromosome and these genes are relevant with histone modification Gene Ontology (GO) terms (Supplementary Table 3). This may indicate potential roles of TssAHet in heterochromatin on the Chromosome X.

By examining the distribution of chromatin states among all 14 tissues, we found that enhancer activity was the most variable between tissues, while promoters were least variable (Fig. 3c-d, Supplementary Fig. 4d,e). Among promoters, TssBiv was least constitutive and often switched to TSS-proximal transcribed or quiescent regions between tissues (Fig. 3d). Hierarchical clustering of samples using the signal intensity of H3K4me1 within EnhA clearly separated different tissue types (Fig. 3e), suggesting that the signal intensity of individual epi-mark in enhancers is highly indicative of tissue identity.

To explore the relationship between proximal regulatory elements (within 2kb of TSS of genes) and tissue-specific gene expression, we identified genes with tissue-specific expression (TSE), which were significantly engaged in known biological functions of specific tissues (Supplementary Fig. 5, Supplementary Table 4). We also observed that TSE were enriched for active states
promoters, transcribed regions and enhancers) and depleted for repressed states in the 2kb regions around their TSS in the corresponding tissue compared to other tissues (Fig. 3f). Furthermore, we found that predicted target enhancers of TSE in a tissue were more constitutive among biologically similar tissues compared with other tissues (Fig. 3g), which was consistent with promoters of TSE (Supplementary Fig. 6).

**Functional characterization of tissue-specific chromatin states**

As enhancers were most variable among tissues compared to other chromatin states, we identified an average of 6,895 tissue-specific EnhAs among 14 tissues, ranging from 1,393 in jejunum to 14,811 in skeletal muscle (Fig. 4a). To further investigate the biological functions of such enhancers, we defined three other types of EnhA, including all-common EnhA (shared among all tissues), gut-common EnhA (shared among gut tissues) and brain-common EnhA (shared among brain tissues). Gene Ontology (GO) analysis of putative target genes of these different types of EnhAs revealed distinct biological functions (Fig. 4b, Supplementary Table 5). For instance, all-common EnhAs were involved in fundamental biological processes (e.g., regulation of mRNA catabolic processes and responses to wounding), whereas gut-common EnhAs were significantly involved in intestinal development, digestion and absorption, and immune response. EnhAs that were specifically active in individual gut tissues showed distinct functions, clearly matching the known biological functions of the tissue in question. For example, jejunum-specific EnhAs were involved in biological processes relevant to T cell and lymphocyte function\(^\text{36}\), while colon-specific EnhAs were mainly engaged in stress-activated MAPK cascades\(^\text{37}\) (Fig. 4b). We observed that intestine- and spleen-specific EnhAs shared many immune functions, and brain-specific EnhAs were significantly involved in memory and learning (Fig. 4b). Furthermore, we observed that genes whose topologically associated with tissue-specific EnhAs (Methods) were specifically
highly expressed in the corresponding tissues (Fig. 4c), and that methylation levels of tissue-
specific EnhAs were lower in the corresponding tissues (Supplementary Fig. 7a), indicating that
these tissue-specific enhancers and their methylation level were accurately predicted to regulate
the expression of associated target genes.

To explore potential tissue-specific transcription factors (TF), first we identified motifs that
were significantly enriched in tissue-specific EnhAs (Fig. 4d, Supplementary Fig. 8a), such as
MEF2A, HNF1B, and HNF4A1 in muscle, liver and intestinal tissues, respectively, which was in
line with previous findings in humans\(^8\). In addition, we found the binding motif of HNF4G, which
participates in the renewal of intestinal stem cells in mice\(^38\) and is specifically active in intestine
(Supplementary Fig. 3b), and is enriched in most intestine-specific EnhAs, whereas CDX2, a major
regulator of intestine-specific genes involved in cell growth and differentiation and is highly
expressed in jejunum over duodenum and ileum\(^39,40\), and its motif is specifically enriched in
jejunum-specific EnhAs. The expression levels of the inferred TFs were higher in the
corresponding tissue than in other tissues (Supplementary Fig. 8b,c), indicating that these tissue-
specific enhancers are hotspots for TF activity and play important roles in the tissue-specific
regulation of gene expression. We further observed that genes linked to tissue-specific EnhAs were
significantly associated with biologically relevant complex diseases in humans and mice (Fig. 4e,
Supplementary Fig. 7b, Supplementary Table 6). For example, colon-specific EnhAs were
associated with recurrent bacterial infection diseases, and cecum-specific EnhAs were
significantly associated with bruising susceptibility diseases.

We also explored potential function for tissue-specific promoters (TssA), and found that
promoters also showed tissue-specific regulatory (TSR) function, but to a lesser degree than
enhancers (Supplementary Fig. 9, Supplementary Table 7).
Chromatin states predictions enhanced the biological interpretations of adaptive evolution and complex traits in pigs

To determine whether genomic regions associated with adaptive evolution are significantly enriched in regulatory elements (REs), we first identified selection signatures (the top 5% of regions measured by Fst) by comparing wild with domesticated pigs in Asian and European populations separately (Supplementary Table 8). We found that genomic regions under selective pressure were most enriched for TssA and TSS-proximal transcribed regions, followed by enhancers, with similar patterns in both Asian and European populations (Fig. 5a, Supplementary Fig. 10a). In examining tissue-specific regulation, our analysis revealed that the all-common TssA were significantly enriched within regions under selective pressure in both populations (Fig. 5b). Interestingly, spleen-specific REs were most enriched in Asian pig domestication, whereas cortex-specific REs were most enriched in European pig domestication (Fig. 5b). Consequently, tissue-specific gene regulation may have played an essential role in the adaptive selection processes that resulted in Asian and European pig domestication. This result was also in agreement with the observation that Asian domesticated pigs being more disease resistant, whereas European domesticated pigs are more active and aggressive.

To ask whether SNPs associated with complex traits in pigs are enriched in regulatory regions, we integrated GWAS signal enrichment analysis for 44 complex traits (Supplementary Table 9) with all 15 chromatin states, and demonstrated that GWAS signals were most enriched in TssA (Fig. 5c), which was consistent with previous findings in humans. We also found that enrichment for variants associated with complex traits was significantly positively correlated with signatures of selection (Supplementary Fig. 10b,c). We then asked if tissue-specific REs were involved in genetic control of specific complex traits. To answer this question, we conducted GWAS signal
enrichment analysis for average daily gain (ADG) in three separate breeds (i.e., Duroc, Landrace and Yorkshire), with emphasis on tissue-specific TssA and EnhA. As we expected, muscle, adipose, liver, and gut-common regulatory elements were the most relevant for ADG (Fig. 5d). In further examining the top ADG QTLs in Landrace (Fig. 5e), we found that the top hit SNPs that are within a muscle-specific EnhA (Fig. 5f) that appears to target two genes (ZNF532 and ALPK2) based on TAD and Hi-C loop derived from CTCF and Hi-C data. Among all seven genes within this QTL, ALPK2 plays important roles in cardiogenesis and was upregulated in the longissimus dorsi muscle in Wannanhua compared with Yorkshire pig45,46, and was the only gene specifically expressed in muscle (Fig. 5f-h). Additional evidence from the eQTLs in muscle showed the highest enrichment in accessible enhancers (EnhA and EnhAMe) compared to other chromatin states (Supplementary Fig. 10d), suggesting that genetic regulatory variants are more likely to influence gene expression through perturbing enhancers. In summary, these results together demonstrated the important role of functional genome annotation for interpreting the molecular mechanisms underpinning complex traits, adaptive evolution and gene regulation.

Comparative analysis of pig, mouse and human epigenomes

The distribution of individual histone epigenetic marks and chromatin accessibility with respect to genomic features (e.g., 5’UTRs and exons) was consistent between pig, mouse, and human (Supplementary Fig. 11). To determine if the chromatin states are similarly conserved between these species, we predicted 15 chromatin states in mouse and human based on the same epigenetic marks in pig. The resulting chromatin state predications demonstrated general similarity among the three species in terms of genome coverage, genomic distribution and sequence conservation (Fig. 6a, Supplementary Fig. 12).
To explore the relationship between the epigenome and DNA sequence conservation among three species, we divided each genome into regions corresponding to 50 different levels of sequence conservation (0<sup>th</sup>-49<sup>th</sup>) (Methods). Our results revealed that the majority of chromatin states showed higher conservation levels in sequences under both rapid and slow evolution than those under neutral evolution, following a U-shaped distribution<sup>7</sup> (Fig. 6b, Supplementary Fig. 13a). We also found that the densities of chromatin states and gene elements followed the similar U-shaped distribution (Supplementary Fig. 13b,c), supporting the hypothesis that conserved epimodifications may buffer negative selective pressures by providing the genome more elastic room to adapt<sup>47</sup>. Furthermore, we categorized orthologous genes into 50 groups based on the degree of conservation of gene expression between species and observed that genes with more conserved expression levels also demonstrated more conserved TssA and TssBiv signatures (Fig. 6c). In further examining sequence extremely conserved (49<sup>th</sup>) or extremely variable regions (0<sup>th</sup>), genes linked to TssA shared by human and pig are involved in basic biological processes, such as ncRNA metabolic process and mRNA catabolic process (Supplementary Fig.14). For the sequence extremely conserved region(49<sup>th</sup>), we found that genes proximal (± 2 kb) by human-specific (comparing with pig) TssA in brain (e.g., Foxg1<sup>48</sup>) are engaged in neuron fate commitment, cerebral cortex development, learning and memory (Fig. 6d, Supplementary Fig. 15, Supplementary Table 10).

Next, we evaluated the evolutionary basis of complex traits in humans. Heritability enrichment analysis of 47 complex traits across 15 chromatin states that were mapped from pigs to orthologous regions in humans found that promoters and TSS-proximal transcribed regions were most enriched for variants (Fig. 6e). We further revealed that the more conserved (species-shared) chromatin states showed significantly higher enrichment of complex traits heritability than the more
divergent (species-specific) chromatin states (Fig. 6f). Then we further examined the role of tissue-specific gene regulation on human complex traits. Our heritability enrichment analysis of complex traits, based on human orthologous regions of tissue-specific EnhAs identified in pigs, demonstrated that tissue-specific enhancers were significantly enriched for the corresponding human complex traits relevant to biological functions of specific tissues (Fig. 6g). For instance, the lung-specific EnhAs were significantly enriched for the heritability of lung forced expiratory volume 1 (FEV1), liver for fasting glucose and cholesterol, colon for Crohn's disease, and cortex for intelligence (Fig. 6g).

Finally, we sought to determine if this annotation of regulatory elements substantiated the use of pig as an appropriate animal model for different human diseases by comparing human, mouse and pig epigenomes in specific tissues. In brain cortex, the mouse-human shared EnhAs exhibited significantly higher heritability enrichment than the pig-human shared EnhAs for most brain-relevant traits, such as attention deficit hyperactivity disorder (ADHD), intelligence, depression and reaction time, with the exception of Alzheimer’s disease, for which heritability was significantly enriched in pig-human shared EnhAs rather than the mouse-human shared EnhAs (Fig. 6h). This was in line with previous findings that pigs have been used as a biomedical model for studying Alzheimer’s disease\(^{21,49}\). Similar observations were found in intestine (Crohn’s disease and inflammatory bowel disease (IBD), but not colorectal cancer, which demonstrated more heritability in the pig-human shared EnhAs) (Fig. 6i) and in adipose (body mass index (BMI), body fat percentage, waist-hip ratio and weight with significantly higher heritability enrichments in the pig-human shared EnhAs) (Fig. 6j). Similar results showed in promoter (TssA) (Supplementary Fig.16). Our findings suggest the pig could be a better biomedical model for certain human traits and diseases than mouse, and vice versa.
Discussion

In this study, we provided the most comprehensive catalog of porcine regulatory elements to date, spanning 14 tissues including six gut-associated tissues, and characterized the dynamic chromatin state landscape across these tissues and uncovered extensive tissue-specific regulation of gene expression.

The annotation of functional elements in human and mouse has proven highly effective for the identification of causative variants of complex traits\textsuperscript{23,27}. Our results also demonstrated that variants of complex traits and eQTLs of growth-related traits were significantly enriched in the active promoters and enhancers annotated by this study. Specifically, we speculate that a potential causative SNP, which was associated with average daily gain and which was found within a muscle-specific enhancer, may regulate the expression of \textit{ALPK2}\textsuperscript{45,46}, a gene demonstrating muscle-specific expression (0.5Mb away). In addition, our annotation of functional elements in pigs allows us to evaluate the potential role of regulatory elements on pig domestication. Our analysis illustrated that signatures of domestication were significantly enriched in porcine regulatory elements. Specifically, genetic variants in the spleen-specific promoters were enriched during Asian pig domestication, whereas variants within cortex-specific promoters were enriched during European pig domestication. This novel insight may reflect the observed distinct phenotypic difference between Asian (more disease resistance\textsuperscript{41}) and European domesticated pigs (more active and aggressive\textsuperscript{42,43}). Further investigation is warranted to deepen our understanding of genetic selection and domestication in the pig. This regulatory element atlas will serve as a valuable source for the livestock community to inform GWAS and eQTL findings, genomic
selection program, and genome editing strategies, as well as to enhance our understanding of genome evolution and adaptation. With continued efforts by the FAANG Consortium, more epigenomic data will be available from diverse samples such as reproduction related tissues, additional developmental stages, and different physiological states. The systemic integration of “omics” data, for instance, the on-going pig GTEx effort will contribute additional insight into the biological mechanisms that underpin agronomic traits, and thereby enhancing genetic improvement of economically important phenotypes.

Finally, this atlas of functional elements provided a unique opportunity for comparative epigenomic analysis between human, mouse and pig, the results of which can inform which species constitute the most appropriate biomedical model(s) for specific human diseases. We observed that regions under positive or negative selective pressure demonstrated higher conservation of epigenetic signatures (such as TssA, TssBiv and TxFlnk) than those under neutral selective pressure, further confirming the hypothesis that elasticity of regulatory conservation may play an important role in the evolution of the less conserved regions (impact of negative selection pressure). Recently evolved liver enhancers (i.e., species-specific) are often associated with genes that show evidence for being under positive selection. Such enhancers have been further demonstrated to actively affect gene expression, although they have a lower effect than those enhancers shared across species when the comparison is controlled for number of enhancer elements acting on that gene. However, the human-specific promoters in brain tissues were enriched in intelligence related genes, which suggests a critical role for epigenomic regulation of novel biological function in humans in the sequence most evolutionarily conserved regions. It is widely accepted that neither mouse nor pig is universally appropriate to serve as an animal model for every human disease. Gene regulatory networks play significant roles in controlling
phenotypic variance of complex traits, including most human diseases. In examining heritability enrichment of 47 complex traits in humans, our epi-conservation analysis among three species by comparing pig-human vs. mouse-human shared enhancers in different tissues revealed novel insights and potential underlying molecular mechanisms as to why pig might be a more appropriate animal model for certain human diseases than mouse and vice versa. This new line of evidence aligns well with many studies of human diseases using either mouse or pig as an animal model. Our study provides a new basis for understanding genetic regulation of complex traits, such as human diseases by focusing on regulatory network conservation across different mammalian species. While the findings from our study are exciting, they are not yet conclusive. More epigenomic data from additional tissues and cell types, as well as additional species such as non-human primates, along with more experimental studies will be needed to extend and functionally validate these mechanisms that underpin complex traits and diseases.

Methods

Animals and tissues

Procedures for tissue collection followed the Animal Care and Use protocol (#18464) from the Institutional Animal Care and Use Committee (IACUC), University of California, Davis. We collected five gut-associated tissues (stomach, jejunum, duodenum, ileum, and colon) of two Yorkshire littermate male pigs at six months of age from Michigan State University. Cecum from two female hybrid pigs (Yorkshires X Hampshires, five months) were obtained at University of California, Davis meat lab. Tissues were first flash frozen in liquid nitrogen, and then stored at –80 °C until further processing.
Library construction and sequencing

We performed ChIP-seq (H3K4me3, H3K4me1, H3K27ac and H3K27me3) experiments on flash-frozen tissue samples using the iDeal ChIP-seq kit for histones (Diagenode Cat.#C01010059, Denville, NJ), as previously described\(^29\). Libraries were sequenced on the Illumina’s HiSeq 4000 with 50 bp single-end reads. ATAC-seq libraries were generated from frozen tissue samples by a modified protocol (https://figshare.com/articles/dataset/Final_ATAC_protocol_docx/13891268) according to the protocol of Omni-ATAC\(^54\) and cryopreserved nuclei\(^55\). The sequencing was performed on Illumina’s NextSeq with paired-end 40 bp reads (PE40). For the RRBS-seq experiments, DNeasy Blood & Tissue Kit (Qiagen, Hilden, Germany) was used for extraction of DNA from frozen tissues. The samples were sent to Novogene (Sacramento, CA, USA) for library construction and sequencing by Illumina HiSeq 4000 PE150. Total RNA isolated from flash-frozen tissue by Zymo Quick-RNA™ Miniprep Kit (Irvine, CA, USA). RNA-seq libraries were constructed by NEBNext® Poly(A) mRNA Magnetic Isolation Module kit (NEB #E7490) and NEBNext® Ultra™ Directional RNA Library Prep Kit for Illumina (NEB #E7720, New England Biolabs (NEB), Ipswich, MA) and sequenced on Illumina HiSeq 4000 with PE100.

Data processing and data summary

In total, 95 new datasets including ChIP-seq (H3K4me3, H3K4me1, H3K27ac, H3K27me3, input control), ATAC-seq, RRBS, RNA-seq in two biological replicates of six gut-associated tissues were generated. We also integrated additional 144 existing pig epigenomic datasets including ChIP-seq (H3K4me3, H3K4me1, H3K27ac, H3K27me3, CTCF, input control), ATAC-seq, RRBS, RNA-seq in the same two biological replicates of eight core tissues (Adipose, Cerebellum, Cortex, Hypothalamus, Liver, Lung, Muscle, Spleen) from our FAANG pilot project (PRJEB14330)\(^29\), and four Hi-C pig liver datasets from publicly available dataset (PRJEB27364)\(^30\).
UC Davis FAANG Functional Annotation Pipeline (https://github.com/kernco/functional-annotation) was applied to process the ChIP-seq, ATAC-seq, and RNA-seq as previously described\(^{29}\). Briefly, the susScr11 genome assembly and Ensembl genome annotation (v100) were used as references for pig. Sequencing reads were trimmed with Trim Galore!\(^{56}\)(v.0.6.5), and aligned with STAR\(^{57}\)(v.2.5.4a) or BWA\(^{58}\) (v0.7.17) to the respective genome assemblies. Alignments with MAPQ scores less than 30 were filtered using Samtools\(^{59}\)(v.1.9). For RNA-seq, gene counts were determined using htseq-count\(^{60}\)(v.0.13.5), and then trimmed mean of M-values (TMM) and transcript per million (TPM) normalization were performed using EdgeR (v3.32.0) and StringTie2 (v.1.3.3), respectively\(^{61}\). For ChIP-seq, after the filtering, duplicates were marked and removed using Picard (v.2.18.7). Regions of signal enrichment (“peaks”) were called by MACS2\(^{62}\) (v.2.1.1). Various quality metrics (e.g., JSD, Supplementary Table 1) were calculated following the method described in our previous study\(^{29}\). RRBS data were processed using Bismark\(^{63}\) (v.0.22.1) with parameter set in RRBS pipeline (https://github.com/zhypan/Functional-Annotation-of-Pig). We called the Hi-C contacts using the Juicer pipeline\(^{64}\) with default parameters.

The global correlations among assays, tissues, and biological replicates were performed by deepTools\(^{65}\)(v.3.5.0). Briefly, the Z-score normalized read signals of all samples with step of 1k bp window were calculated by multiBigwigSummary and were presented by plotCorrelation. The signal of marks along with protein coding genes were generated by deepTools\(^{65}\) (computeMatrix scale-regions function) with parameter -a 2500 -b 2500. The Z-score was used to normalize bigWig of five marks as input files.

**Annotation of chromatin states**

ChromHMM\(^{66}\) (v.1.20) was used to train the chromatin state prediction model by integrating ChIP-seq (H3K4me3, H3K4me1, H3K27ac, H3K27me3, and input control) and ATAC-seq data.
from two biological replicates of 14 tissues. The same tissue of two biological replicates were collectively considered as one tissue epigenome. The 15-state model was chosen, as it presented maximum number of states with distinct epigenetic mark combinations. We labelled these 15 chromatin states based on their combinations of histone modifications and enrichment around TSS. Then the fold enrichment of each chromatin state for each external gene element (e.g., exon, CpG islands) was calculated by \((C/A)/(B/D)\), where \(A\), \(B\), \(C\), \(D\) are the number of bases in a chromatin state, a gene element, overlapped between a chromatin state and a gene element, in the genome, respectively. In addition, we also computed chromatin state fold enrichment in mammalian conserved elements which identified from Multiple Sequence Alignments (MSA) using the Genomic Evolutionary Rate Profiling (GERP) software based on 103 mammals (ftp://ftp.ensembl.org/pub/release-100/bed/ensembl-compara/103_mammals.gerp_constrained_element/). The methylation level of each state and its up- and down- stream 10kb was calculated by the computeMatrix scale-regions function of deepTools with parameters --binSize 500, --regionBodyLength 2000 and --skipZeros.

**Clustering of large-scale chromatin structure**

To examine genome-wide chromatin structure, we first divided the genome (excluding chrUn) into 1,224 fragments of 2Mb in length. Then we calculated the state frequencies (state bin/total bin) in each 2Mb fragment for each tissue and then the average frequency across tissues. To identify modules, column clustering was performed by \(k\)-means=12, and rows were clustered using \(k=3\). In addition, we calculated number of protein-coding, lncRNA, and CpG islands for each 2Mb fragment by BEDTools (v.2.29.2). We also calculated the average TPM of protein coding gene and average methylation level across 14 tissues in each 2Mb fragment. Then the average gene
expression and methylation level in each of the 12 modules were calculated and a Student’s T-test was performed with parameter setting ref.group = "M3".

**Chromatin state variability**

For each state, we first obtained regulatory regions across 14 tissues (RRATs) (Supplementary Fig. 4a-c) using BEDtools merge function (any regulatory region between two tissues overlapped by 1 bp was merged), then we calculated the total genomic length for each tissue (GL) and the total combined genomic length (TGL) for RRATs. The relative state coverage per tissue was derived by GL/ TGL (Supplementary Fig. 4d). Finally, followed by the order from high to low based on the GL/TGL value in each tissue, we calculated the total genomic length of accumulated tissues (aGL) by adding one tissue each time until all 14 tissues were added, and the cumulative state coverage was calculated as aGL/ TGL. States whose cumulative coverage changed faster than others were considered to be less constitutive (more variable) states.

**Chromatin state switching between tissues**

Chromatin state switching between tissues was calculated by pairing two tissues. Given a pairing of A and B tissues, we first counted total bins of chromatin state “e” in A (TbAe), then obtained the overlap bins of chromatin state “e” (Obe) in A and B, then computed the state switching probabilities using Obe/TbAe for the tissue A to B transition and Obe/TbBe for the tissue B to A transition. By averaging these calculations for a pair of tissues, we obtained the pair switching probabilities. We calculated the state switching probabilities in between intestinal tissues, between brain tissues (Supplementary Fig. 6a,b) and between 8 distinguishable tissues (jejunum, cortex, adipose, liver, lung, muscle, spleen).
Hierarchical epigenome clustering

We first calculated the an epigenetic mark’s signal confidence scores ($-\log_{10}(\text{Poisson P value})$) within 200 bp of the genomic regions for each mark of each sample as described in http://jvanheld.github.io/stats_avec_RStudio_EBA/practicals/02_peak-calling/peak-calling_report.html#data_sets. Then, we extracted a specific mark’s signal confidence score of each sample for specific state of RRATs regions. For example, we extract H3K4me1 signal confidence scores for EnhA. After combining all samples’ mark confidence scores for each tissue and each state, we constructed a distance matrix using the ward.D2-linkage hierarchical clustering following by Euclidean distance method in R.

Promoter enrichment analysis of tissue-specific expressed genes among 14 tissues

To evaluate how chromatin state changes at promoter region of TSE genes across tissues, we first performed a Student’s $t$-test among 14 tissues to identify tissue-specific expressed genes based on TPM. We further grouped some tissues into different sub-groups such as small intestine (Jejunum, Ileum, Duodenum), large intestine (Cecum, Colon), and brain (Cortex, Cerebellum, Hypothalamus), and identified tissue-specific expressed genes by excluding the tissues in the same sub-group. Then we selected genes with the top 5% $t$-value as TSE genes. The biological process of GO enrichment for these TSE genes were identified by WebGestalt2019 using the default significance level (FDR< 0.05). Then we calculated the chromatin state fold enrichment of TSE (up and down stream 2000bp around TSS) in each tissue and the change in enrichment by TSE enrichment in specific tissue minus other tissues.

Chromatin state switching of target enhancer (EnhA) of TSE gene
To evaluate how enhancers of TSE genes switch among tissues, we first identified the target enhancers of TSE genes following the method described in our recent study. Briefly, we generated the predicted TADs from CTCF ChIP-seq data by FIMO following the method described in Oti, et al. Then we predicted the enhancer-gene pairs according to the Spearman’s rank correlation of every possible combination of regulatory element H3K27ac signal and gene expression value within each TAD. Benjamini-Hochberg adjustment (FDR < 0.05) was used to define putative interacting pairs. The enhancers in the enhancer-gene pairs that corresponded to TSE genes were considered as TSE genes’ target enhancers. Finally, we computed enhancer state switching probabilities of TSE genes among tissues using the method described above.

**TSR of enhancer, promoter and their putative functional regulation**

For strong enhancer (EnhA) identified in each tissue, we counted the bins of overlapping RRATs by comparing to other tissues. If the number of bins ≥ 1, the tissue of this RRATs region would be assigned 1, otherwise it was assigned 0. We generated a total of 17 modules of tissue-specific regulatory elements (TSR) enhancers. The 17 modules included all-common (presented in all tissues), gut-common (presented in all 5 intestinal tissues), brain-common (presented in all 3 brain tissues) and 14 tissue-specific modules. The same method was used to obtain TSR for promoters (1_TssA). In addition, we performed enrichment analyses (GO, Human Phenotype Ontology (HPO), Mouse Phenotype) based on genes proximal to TSR using the GREAT tool with default parameters except for TSR promoters (proximal 2kb upstream, 1kb downstream, plus distal up to 3kb). We used a cut-off of FDR<0.05 for both the binomial and the hypergeometric distribution-based tests.
The motifs of tissue-specific EnhAs were identified by HOMER\textsuperscript{73} (v.4.11) with cutoff FDR<0.05. We selected the top three enriched or tissue function relevant motifs for each tissue as the candidate tissue-specific EnhAs motifs and generated a total of 51 motifs enriched in tissue-specific EnhAs. In addition, we used these 51 motifs as known TF motifs to conduct the enrichment for all tissues by HOMER. The mRNA expression of corresponding TFs in pigs were used to calculate the correlation with motif enrichment.

**Selection signature enrichment analysis of chromatin state**

A total of 406 whole genome sequence datasets (Supplementary Table 8) in pigs (Asian wild (58) and domestic pigs (129), European wild (35) and domestic pigs (184)) were trimmed by Trimmomatic\textsuperscript{74} (v.0.39), mapped by BWA (0.7.17), and marked duplicates by GATK\textsuperscript{75} (v4.1.4.1) MarkDuplicates with default parameters. The SNPs of Gvcf for each sample were called by GATK HaplotypeCaller. All Gvcf were then combined and the variants for each sample were called by GenotypeGVCFs. After SNP calling, the variants were filtered using VariantFiltration (QD < 2.0, MQ < 40.0, FS > 60.0, SOR > 3.0, MQRankSum < -12.5, ReadPosRankSum < -8.0) to remove low-quality SNPs. We then performed Fst analysis between Asian wild and domestic pigs, and between European wild and domestic pigs, and calculated the fold enrichment of selection signature for chromatin states using the same method for gene elements enrichment described above.

**GWAS and eQTL signal enrichment of chromatin state**

The pig GWAS data of 44 traits was described previously\textsuperscript{76,77}(Supplementary Table 9). First, we filtered out all SNPs with minor allele frequency below 0.5%, with a large deviation from Hardy--
Weinberg proportions ($P<1.0^{-6}$), or with a R2 value of the imputation accuracy estimated by Minimac4 of less than 0.4. We performed GWAS signal enrichment of 44 pig complex traits (3 daily gain related, 20 lipid related, and 21 feed efficiency related) for each chromatin state across 14 tissues using a 10,000 times genotype cyclical permutation tests. The eQTLs data in pig muscle with FDR<0.05 were used to calculate the fold enrichment for the chromatin states using the same method above.

**Interspecies conservation of chromatin state**

We collected data from ENCODE, Roadmap Epigenomics and published articles (9 tissues in human and 7 tissues in mouse, Supplementary Table 11,12), including ChIP-seq (H3K4me3, H3K4ac, H3K4me1, H3K27me3, Input), ATAC-seq, DNase-seq, and RNA-seq. In total, we obtained six matched tissues (small intestine, liver, spleen, lung, adipose, cortex) among pig, human, and mouse. All the data were processed following the same pipeline used in pig. The GRCh38 (human) and GRCm38 (mouse) assemblies with Ensembl annotations (v100) were used for data analysis. Chromatin states of human and mouse were also trained by ChromHMM and 15 chromatin states were identified. To explore the relationship between sequence conservation and epi-conservation among the three mammals, we first divided the genome into 50 equal sized sets (0th-49th) with increasing average PhyloP scores using the method detailed by Xiao et al. Briefly, the human genome was divided into 15 million 200 bp segments. Then average PhyloP score (100 vertebrate genomes) was computed for each 200 bp segment. These genomic segments were divided into 50 equal sized sets from the fastest changing sequence (smallest PhyloP scores) to the most conserved (greatest PhyloP scores). (Supplementary Fig. 13d). To quantify epigenomic conservation, we downloaded the whole genome alignments UCSC chain files among human
(hg38), pig (SusScr11), mouse (mm10) and processed as described in the UCSC Genome Wiki website (http://genomewiki.ucsc.edu/index.php/HowTo:_Syntenic_Net_or_Reciprocal_Best) to derive reciprocal best chains. Then we converted genomic coordinates between assemblies using the UCSC Liftover tool (https://genome.sph.umich.edu/wiki/LiftOver) based on 0.65 sequence identity. All the chromatin states in pig and mouse were lifted over to human. The conservation rate (0~1) of each region of each state from pig to human was calculated based on state region coverage of pig over human. If there was no overlap it was assigned 0, if completely occupied it was assigned 1. The same analysis was conducted for pig to mouse and mouse to human. Furthermore, we performed genomic and epigenomic conservations for every pair of mammalian species in each tissue. Finally, we conducted the same analysis on mammalian conserved score based Genomic Evolutionary Rate Profiling (GERP) using 103 mammalian genomes (ftp://ftp.ensembl.org/pub/release-
100/compara/conservation_scores/103_mammals.gerp_conservation_score/)

To examine the biological relevance of sequence extremely variable (0th-2th sets) and conserved regions (47th-49th sets), we extracted the human-pig shared and human-specific chromatin state TssA from these regions. Then the GREAT tool with parameter of proximal 2kb upstream, 1kb downstream, plus distal up to 3kb was used to conduct GO function enrichment analysis.

**Expression conservation versus epi-conservation**

The TPM of 14302 orthologous genes from pig, human, and mouse were used to identify differentially expressed genes in each tissue using the Student’s t-test. We sorted the genes by p-value within each species and divided them into 50 equally sized sets. Then we calculated the average epi-conservation score of states in the 20kb region around TSS of gene in each set.
Heritability enrichment of human complex traits in chromatin state

To explore how conserved or species-specific chromatin states affect complex traits in humans, we extracted six types of species-share or species-specific regulatory elements (all_shared, human_mouse_shared, human_pig_shared, human_specific, mouse_specific, pig_specific). We applied stratified linkage disequilibrium score regression (LDSC) to partition heritability of 47 human complex traits into distinct functional categories, which revealed which functional regions explained more genetic variation of complex traits from an evolutionary point of view. These functional categories included six types of species-shared/specific regulatory elements, chromatin state of each tissue, and TSR of EnhA/TssA. We calculated the stratified LD scores using 1000G Phase 3 European human samples, where only HapMap3 SNPs with INFO≥0.9 and MAF>0.05 in 1000G European samples were used (the 100G samples and default SNP weights were obtained from https://github.com/bulik/ldsc).

The GWAS summary statistics for 47 human complex traits were obtained from public databases (Supplementary Table 13), with an average sample size of 321,978 (all European ancestry) and a high quality overlap with HapMap3 panel. In addition, these GWAS results have a mean χ^2 statistics of > 1.02 and a heritability Z-score of > 4. We also performed default quality control for each GWAS summary statistics by LDSC to remove GWAS SNPs that are with MAF≤0.01, genotype call rate ≤ 0.75, INFO ≤ 0.9, out-of-bounds P-value, duplicated Rsid, strand ambiguous variants and extreme large χ^2 statistics. The results of LDSC regression for base model, which has not been partitioned heritability, are available in Supplementary Table 14.
Data availability

All high-throughput sequencing data in this study were deposited in European Nucleotide Archive (ENA) with accession number PRJEB37735 and PRJEB14330. All raw data are also available through the FAANG portal (https://data.faang.org/dataset). All processed data are publicly available at https://doi.org/10.6084/m9.figshare.13480425. Chromatin states of pig, mouse, and human are available in Genome Browser: http://genome.ucsc.edu/s/zhyp/susScr11_15_state_14_tissues_new; http://genome.ucsc.edu/s/zhyp/mm10_7tissues_chr_state; http://genome.ucsc.edu/s/zhyp/hg38_9tissue_chr_state.

Code availability

The pipeline for RNA-seq, ATAC-seq, DNase-seq and ChIP-seq processing is available at https://github.com/kernco/functional-annotation. RRBS pipeline and other processing codes are publicly available at https://github.com/zhyp/Functional-Annotation-of-Pig.

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**Author contributions**

H.Z., L.F. Z.P. P.R., C.W. E. and C.K.T. conceived and designed the study. C.E., Y.W., K.C. and Z.P. were responsible for sample collection. Z.P., Y.W. and M.H. performed ChIP-seq, ATAC-seq, RNA-seq. Z.P., N.T. and K.W. contribute for RRBS data collection. Z.P., Y.Y. L.F. and C.K. conducted bioinformatic analysis. Z.X., G.S., GS. S., MS. L., M.F., and P. KM. were responsible for pig GWAS data analysis. H.Y. and L.B. responsible for pig selection signature collection. Z.P., L.F., Y.Y. and H.Z. wrote the initial draft of the manuscript. M.H. C.W. E., P. R., and C.K. T. revised manuscript. All co-authors contributed to the final manuscript.

**Competing interests**

The authors declare no competing interests.
Additional information

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Figures and legends:

**Fig. 1** Data summary of epigenomic information across tissues and marks.

- **a**, Tissues assayed by this study.
- **b,c**, Average peak number and genome coverage for each epigenetic mark in each tissue.
- **d**, The Pearson correlations among assays, tissues, and biological replicates (P348 and P350) based on the normalized signal in 1kb windows stepped across the whole genome.
- **e**, Average epigenetic mark signals proximal to protein coding genes. TSS, transcription start site. TES, transcription end site.
- **f**, Epigenetic signal at the *MYO1A* locus according to different assays and in different tissues. Vertical scale UCSC reads signal 0-200 for RNA-seq, 0-100 for H3K27ac and H3K4me3, 0-50 for other marks and ATAC-seq.
Fig. 2 Chromatin landscape across 14 tissues.

a, b, Definitions and abbreviations of 15 chromatin states. c, Emission probabilities of individual epigenetic marks for each chromatin state. The color from white to deep blue indicate emission probability (0-1). d, Genomic coverages of each chromatin state. M±SD, means ± standard deviation e, Average enrichments of chromatin states for genomic annotations, including CpG islands, genes, TSS/TES_1K (± 1 kb around transcription start/end sites), expressed genes (TPM >=0.1), repressed genes (TPM < 0.1) in each tissue. f, Fold enrichments of chromatin states
for non-coding mammalian conserved elements from Genomic Evolutionary Rate Profiling (GERP). g, Density of each chromatin state in positions relative to gene transcription start sites (TSS). h, Average methylation level of chromatin states in jejunum. i, Hi-C (250kb resolution), predicted chromatin states, and epigenetic signal, and normalized methylation level landscape in jejunum across the Chr7. j, Chromatin state landscapes and mRNA expression at VIL1 locus (chr15:120,459,825-120,493,312, susScr11) across 14 tissues in pig. Vertical scale UCSC reads signal 0-200 for RNA-seq.
Fig. 3 Genome-wide chromatin state dynamics across tissues.

a, Clustering of 2-Mb intervals (1,224 columns) into modules (M1-M12) based on average chromatin state frequency across tissues in each interval. b, Average mRNA expression (TPM) of genes and average methylation level of 2-Mb intervals belonging to each module. The statistical significances for comparisons were calculated using t test, where “*” means $P < 0.05$, “**” means $P < 0.01$, “***” means $P < 0.001$. c, Chromatin state variability based on cumulative genome coverage fraction. Dash line = 0.75. d, Chromatin state switching between all tissues. e, Hierarchical epigenome clustering using H3K4me1 signal in EnhA states. f, Chromatin state
enrichment relative to promoters of genes with jejunum-specific expression genes, relative to muscle. Chromatin state switching of target enhancers (EnhA) of jejunum-specifically expressed genes in other tissues.
Fig. 4 Tissue-specific strong enhancers (EnhA) and their potential functions in 14 tissues.
a, The number and enrichment distribution of 17 modules of TSR (strong enhancers (EnhA)) in tissues. TSR: tissue-specific regulatory elements. The top colors represent 17 modules of strong enhancers (column) referred to the legend on the right side. The side colors represent 14 tissues (row) referred to the legend on the right side. b, Functional enrichment of proximal genes for each module based on gene ontology (GO) biological processes. The columns represent 17 modules of strong enhancers. The rows represent GO terms in each module. All GO terms are presented in Supplementary table 5 (The notes in heatmap are summary function of nearby GO terms enrichment (up-noted from jejunum to spleen, down-noted for lung, muscle and adipose)). c, The mRNA expression (TPM) of EnhAs’ putative target genes in each module. The columns represent the genes in each module, the rows represent each tissue. d, The enrichment of transcription factor motifs in each module. The columns represent 17 modules of EnhAs. The rows represent motif in each module. All enriched motifs are presented in Supplementary Fig. 8a. e, Enrichment for human phenotypes in each module, based on proximal genes. The columns represent 17 modules of EnhAs. The rows represent phenotypes in each module. All phenotypes’ enrichment are presented in Supplementary table 7. The notes in heatmap are summary function of nearby phenotypes enrichment (each color stands for each tissue).
Fig.5 Chromatin state plays an important role in pig domestication and complex traits.

a, Domestication selection signature enrichment of chromatin states in Asian and European pigs. ASD: Asian pig domestication. EUD: European pig domestication. Dash-line = 1, above dash line means significant enrichment. b, Domestication selection signature enrichment in tissue-specific promoters (TssA) between Asian and European pigs. Dash-line = 1, above 1 dash line means significant enrichment. c, Genome-wide association studies (GWAS) signal enrichment within chromatin states across 14 tissues and 44 complex traits in pigs. The statistical significances for comparisons were calculated using a t test, where “***” means $P < 0.001$. d, GWAS signal enrichment of promoter (TssA) and strong enhancer (EnhA) tissue-specific regulatory elements (TSR) in average daily gain (adg) of three pig populations (dd: Duroc, ll: Landrace, yy: Yorkshire). Dash line=$-\log_{10}(P=0.05)$, over dash line means significantly high enrichment. e, Manhattan plot of average daily gain in the Landrace population. f, Chromatin states in the genomic region where GWAS hits for each tissue (dashed rectangle box includes a muscle-specific enhance where SNPs of GWAS hits locate; two arrows in red were predicted from CTCF TAD and H3K27ac signal that suggest the muscle-specific enhancer may target $ZNF532$ and $ALPK2$). g, Hi-C loop (25kb resolution) depiction between a muscle-specific enhancer and putative target genes. Purple shading for the Hi-C data means loop intensity (auto-scale). Two highlighted Hi-C loop with the red circles are potential contacts between a muscle-specific enhancer and $ZNF532$ and $ALPK2$. h, Expression (normalized and centered TPM) of genes proximal to the muscle-specific enhancer.
**Fig. 6** Interspecies conservation of chromatin states.

(a) 15 chromatin states predicated in three species. The colors from white to deep blue indicate emission probability from 0 to 1. (b) Relation between sequence conservation and epigenomic conservation across six tissues. 50 genomic regions were ordered from the fastest changing (0th), to neutral (20th), and to slowest changing (49th) in terms of sequence conservation (Supplementary Fig. 13d). Epigenome conservation (see method) of chromatin states (b) within...
these regions was calculated for pig to human. Value in each region for each chromatin state was then plotted. c, Relation between expression conservation and epigenomic conservation across six tissues. Expression conservation was based on expression of 14,302 orthologous genes among 3 species. Regions were ordered from the biggest difference in expression (0th), to the smallest difference (49th). d, GO enrichment was based on genes proximal to (± 2 kb) human-specific TssA in sequences of extreme conservation sets (49th). Count refers to the number of genes. e, Human GWAS signal enrichment in different chromatin states. dash line=1, over dash line means significant high enrichment. Error bars represent standard error around the estimates of enrichment. Same meaning for dash line and error bars in following sub-figures. f, Human GWAS enrichment in 6 groups of species-specific or shared EnhA. hpm_share stands for human-pig-mouse shared. g, GWAS enrichment of pig tissue-specific enhancer (EnhA) in humans. *means significant enrichment (FDR<0.05). h,i,j, Different GWAS enrichments between human-pig and human-mouse shared strong enhancers (EnhA) in brain cortex, small intestine, and adipose, respectively.

Supplementary information