

1 Screening and assessment of functional variants by regulatory 2 features from epigenomic data in livestock species

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18 Abstract

19 Single nucleotide polymorphisms (SNPs) and small insertions/deletions (2-50bp) in
20 genomic regulatory regions may impact function, and although widespread, they are
21 largely unexplored in livestock. Here leveraging >500 epigenomic datasets from pigs,
22 cattle, sheep, and chickens, 8-39 million variants were identified with candidate
23 functional confidence. Using our Functional Confidence scoring system, these
24 candidate functional variants were further ranked as High, Moderate, Low, Minimal, or
25 Possible functional confidence by scoring for likelihood of disrupting transcription
26 factor (TF)-chromatin binding based on their presence in eight genomic regulatory
27 features. Predictive reliability analysis of estimated breeding values (EBVs) based on
28 High/Moderate Confidence variants from pig shows a 23~46% increase in reliability
29 compared to EBVs based on general SNPs, illustrating the versatility of Functional
30 Confidence scoring system for identifying potential functional variants in livestock.
31 Therefore, we developed the Integrated Functional Mutation (IFmut) platform and
32 embed the Functional Confidence scoring system for users to effortlessly navigate
33 through epigenomic data or pinpoint specific genomic features/regions, uncover
34 potential function of new variants or previously identified ones. Our work offers the
35 scientific community a powerful and flexible tool, tailor-made for delving deep into
36 variant function, setting a new benchmark in livestock research and breeding strategies.

37 **Introduction**

38 Variants in *cis*-regulatory elements can greatly affect gene expression, consequently
39 influencing organismal phenotype (1-4). While some variants are harmless (i.e., neutral),
40 others can lead to severe diseases or other deleterious effects (5). In human disease
41 research, the identification and characterization of variants are crucial for determining
42 the underlying causes of genetic disorders (6-8). Several large-scale initiatives, such as
43 the 1000 Genomes Project, have facilitated exploration of variants in the human
44 genome (9, 10). By contrast in livestock research, variants are typically studied through
45 resequencing in livestock populations to improve breeding strategies (11-14). The
46 identification of variants is a crucial initial step, but determining whether a variant is
47 functional in livestock species such as pig (*Sus scrofa*), cattle (*Bos taurus*), sheep (*Ovis*
48 *aries*), and chicken (*Gallus gallus*) poses a great challenge.

49 Advances in publicly available bioinformatic analytical toolkits, e.g., the
50 Encyclopedia of DNA Elements (ENCODE) (15), have driven considerable progress in
51 functional variant screening in humans, uncovering previously unrecognized functions
52 of several regions in the human genome. In addition, integrated analysis of expression
53 quantitative trait loci (eQTLs) and variants (16-18), along with establishment of the
54 regulomeDB database (19), has also facilitated identification of *cis*-regulatory elements
55 and trans-acting factors that influence gene expression, revealing a variety of regulatory
56 mechanisms of the human genome. Despite these innovations in human genomic
57 research, the exploration of variants with regulatory function in the genomes of
58 important livestock species remains limited.

59 Genomic selection (GS) (20) has brought about a revolution in livestock and
60 poultry breeding, enabling greater precision in the selection of individuals based on
61 quantitative traits, such as growth rates or disease resistance (21-25). However, GS
62 efficiency relies on SNP markers distributed throughout the genome because all of the
63 QTLs and the SNPs used for these analyses are in linkage disequilibrium (LD) (20,
64 26, 27). Although GS can enhance the reliability of estimated breeding values (EBVs),
65 EBVs based on GS markers ignore the potentially significant source of functional

66 genetic variation available in genomic regulatory elements. Variants in regions
67 containing gene regulatory elements can potentially modulate gene expression (28). By
68 integrating variants and SNPs in regulatory regions into genomic prediction analysis,
69 the reliability of EBVs for target traits may be increased. However, despite their
70 possible value for improving EBV reliability, studies identifying candidate functional
71 variants that may be informative for GS are lacking.

72 In this study, we identified genomic regulatory features in over 500 datasets
73 comprising transposase-accessible chromatin with sequencing (ATAC-seq) data,
74 DNase I hypersensitive site sequencing (Dnase-seq) data, H3 lysine 27 acetylation
75 (H3K27ac) ChIP-seq, and transcription factor ChIP-seq data from pigs, cattle, sheep,
76 and chickens. We then identified the candidate functional variants and employed a
77 scoring system to assess the likelihood of variants affecting regulatory function (i.e.,
78 Functional Confidence Score) based on their presence (or absence) in eight different
79 regulatory features/regions in genomes of the four livestock species. The identified
80 variants were then ranked into five categories (12 sub-categories) based on Functional
81 Confidence scores, in descending order from High, Moderate, Low, and Minimal
82 functional confidence, to possible association with regulatory function. We further
83 tested whether genomic prediction with High and Moderate confidence IFmut variants
84 identified from three tissues of pig could improve the predictive reliability of EBVs
85 over that of EBVs based on 11000 randomly selected SNP markers in pig. We then
86 constructed the Integrated Functional Mutation (IFmut) database and Functional
87 Confidence scoring system to provide a public resource for researchers. This study
88 provides a large database with a versatile and powerful online toolkit, along with a
89 proof-of-concept demonstration of IFmut for exploration of functional variants in
90 fundamental research and molecular breeding of livestock.

91 **Results**

92 **A large scale epigenomic screen of potential functional variants across four
93 livestock species**

94 To screen for functional variants that potentially affect gene expression in livestock, we
95 first obtained SNP and small InDel genomic variants in pig (susScr11), cattle (bosTau9),
96 sheep (oviAri4), and chicken (galGal5) from the Ensembl database, including more
97 than 63 million in pig, over 97 million in cattle, over 63 million in sheep, and over 23
98 million in chickens. Analysis of their distribution and predicted effects using
99 ChIPseeker R package and SnpEff software indicated that more than 90% of the
100 variants were located in non-coding regions in all four species (Fig. 1A-D;
101 Supplemental Fig. 1), aligning well with previous studies that showed variants are
102 highly prevalent in intergenic and intronic regions of the human genome (7, 29, 30).
103 Thus how to identify the potential functional variants from a large pool across four
104 livestock species is still a challenge.

105 Since genomic regulatory features, especially transcription factor binding sites
106 (TF binding sites) identified by epigenomic analyses, can be informative of the potential
107 function of variants (19, 31), we sought to screen for potential function variants in the
108 above libraries using epigenomic data. To this end, we collected 583 total epigenomic
109 datasets (including ATAC-seq, Dnase-seq data, H3K27ac ChIP-seq, TF ChIP-seq and
110 Hi-C) from pigs, cattle, sheep, and chickens generated in previous studies such as the
111 FANNG project (32, 33), and our own previous study (34). After processing raw reads
112 from ATAC-seq, Dnase-seq, or ChIP-seq data using the ENCODE pipeline, we
113 removed 6 samples due to low number (<10000) of significant peaks and 35 samples
114 due to low correlation among biological replicates ($R < 0.8$). Ultimately, 538 datasets
115 from 19 tissues and 12 cell lines met quality control standards (Fig. 1E,F), including
116 256 ATAC-seq, 26 Dnase-seq data, 167 H3K27ac ChIP-seq, 80 TF ChIP-seq (63 for
117 CTCF from pigs, cattle, sheep, and chicken and 17 for RAD21, EGR1, KLF2, KLF4,

118 OSR1, OSR2, SMC2, CAP-H and BRD4 from chicken), and 9 Hi-C datasets from pig
119 (Fig. 1G-J). In total, 125, 193, 24, and 196 total datasets were compiled for pigs, cattle,
120 sheep, and chickens, respectively (Fig. 1E).

121 ENCODE guidelines (<https://www.encodeproject.org/>) were then applied to
122 identify genomic regulatory regions containing basic regulatory features and/or TF
123 binding site-related features using these datasets. In total, more than 350000 non-
124 redundant genomic regions with basic regulatory features were identified across all four
125 species, including open chromatin regions (OCR), H3K27ac significant peaks, and
126 nucleosome-free regions (NFR; Table 1). Furthermore, footprint calling and significant
127 TF binding peak calling followed by genomic mapping with TF motif positional weight
128 matrices (PWMs) in the called features yielded between 41460-171290 non-redundant
129 genome regions with regulatory features related to TF binding sites in each species
130 (Table 1). The total length of non-redundant genomic regulatory regions accounted for
131 approximately 31.56% of the pig reference genome (susScr11), while in cattle
132 (bosTau9), sheep (oviAri4), and chicken (galGal5), these accounted respectively for
133 30.74%, 12.63%, and 37.35% (Table 2).

134 Genomic variants positioned within transcription factor binding sites, such as
135 in RegulomeDB, often result in functional consequences (19). Since the above
136 genomic regions containing basic regulatory and TF binding-related features were
137 identified through DNA-TF interaction data, variants detected in these regions were
138 likely to have transcription regulation function in the host livestock species. Using
139 BEDTools, we then determined which variants in our initial calling were located in
140 these regulatory features, which yielded 21005715 (32.90%; Fig. 2A) SNPs and small
141 InDels in pig, while 39157953 (40.32%; Fig. 2B) were detected in the cattle genome,
142 8194045 (12.97%; Fig. 2C) in sheep, and 10896983 (47.05%; Fig. 2D) in chicken,
143 which we collectively designated as potential functional variants.

144 **A scoring system to rank variants by likelihood of functional impacts**

145 In the current study, we identify a multitude of variants with predicted
146 functional/phenotypic consequences in four livestock species, and the number of such
147 variants in each species was positively correlated with proportion of the genome
148 occupied by regulatory features ($R=0.74$; Supplemental Fig. 1E). To further distinguish
149 differences in the likelihood that a regulatory region variant will indeed impact
150 transcription regulation in livestock species, we developed a functional confidence
151 index similar to that used by RegulomeDB for variant classification with human TF
152 ChIP-seq data (19). At present, only 80 TF ChIP-seq datasets are available in livestock,
153 the vast majority of which were generated for CTCF (in total 63), with only chicken
154 having 17 ChIP-seq data for 9 TFs with ChIP-seq data, compared to the 876 TFs
155 covered by 3537 ChIP-seq data in RegulomeDB v.2. Thus, due to the lack of TF ChIP-
156 seq data in livestock, functional confidence scoring instead relied on a combination of
157 ATAC-seq/Dnase-seq (i.e., OCR and footprints) and H3K27ac ChIP-seq (i.e., NFR and
158 significant narrow peaks; Fig3A; Table 3). In addition, quantitative trait loci (QTL) data
159 were also collected, since variants in these regions can also potentially impact
160 agronomic traits (Supplemental Table 1).

161 In the Functional Confidence scoring system, the greater the number of
162 regulatory features used to determine the presence of SNPs/small InDels in TF binding
163 sites, the higher the likelihood that a variant could affect transcriptional regulation
164 (Table 3). Based on the prominent association of NFRs, OCRs and TF footprints
165 (especially those containing fully or partially matching recognition motifs) with
166 transcriptional activation, variants in these regions had the highest likelihood of
167 affecting TF binding and gene expression, and were therefore scored as high functional
168 confidence variants (Category 1). Variants that met these criteria but were never found
169 in NFRs were subsequently scored as moderate functional confidence (Categories 2a-
170 2d), suggesting a moderate likelihood of affecting TF activity. Moreover, within
171 Category 1 and 2, variants were present in QTLs, were assigned higher scores

172 (Categories 1a, 1c, 2a and 2c respectively), whereas variants were not in QTLs, had
173 slightly lower likelihoods (Categories 1b, 1d, 2b and 2d respectively). By contrast,
174 variants found in DNase/ATAC-seq or H3K27ac ChIP-seq data but not in TF footprints
175 or recognition motifs were included in Categories 3 and 4, with low functional
176 confidence and minimal functional confidence, respectively, in their likelihood of
177 affecting TF binding. Finally, Category 5 was reserved for variants detected only by
178 H3K27ac ChIP-seq, and were therefore potentially associated with transcriptional
179 regulation (see Table 3 for a key of criteria).

180 Next, the candidate functional variants identified by our study were ranked based
181 on our Functional Confidence scoring system. Then, Figure 3 shows a summary of
182 variant numbers in each functional confidence category for pig (Fig.3E), cattle (Fig.
183 3B), sheep (Fig. 3C), and chicken (Fig. 3D). Among these variants, a total of 240938,
184 3096314, 280103, and 204100 SNPs/small InDels were included in high and moderate
185 functional confidence categories (Category 1 and 2) Categories 1 and 2, accounting for
186 1.15%, 7.91%, 3.42% and 1.87% of all potential functional variants in pigs, cattle,
187 sheep and chickens, respectively (Fig. 3B-E).

188 To validate the variants in our above analysis were present in population data
189 and that functional confidence scoring could be applied to whole-genome sequencing
190 (WGS) data, we obtained 22926176 minimum allele frequency (MAF>0.047) filtered
191 variants in WGS data from 491 individual pigs across 61 breeds generated in our
192 previous study (34). Among these variants, 7557763 (32.97%) were identified as
193 potentially functional variants, 87002 (1.15%) of which fell into categories 1 or 2 (Fig.
194 3F). Overall, the proportions of variants in each category filtered by MAF from WGS
195 data were similar to that of variants obtained from Ensembl (Fig. 3B,F). These results
196 indicated that taking MAF into account did not affect the proportion of variants in each
197 category, but could reduce the number of candidate functional variants. Thus on animal
198 breeding a lower MAF threshold (e.g. 0.01) have to consider for functional variants to
199 keep their efficiency of animal breeding.

200 **The functional confidence scoring in eQTL classification and EBV reliability**
201 **assessment**

202 Although eQTLs are reportedly associated with gene expression (35, 36), some TF
203 ChIP-seq and DNase-seq studies in humans suggest that more than 50% of eQTLs are
204 not associated with TF binding sites (19, 37), implying that genomic regulatory features
205 could be used to assess the potential regulatory function of eQTLs. To test this possible
206 use of our Functional Confidence scoring system, we obtained cis-eQTL data from
207 adipose, liver, spleen, hypothalamus, kidney, lung, muscle, and rumen of cattle from
208 the farmGTEX database (<https://www.farmgtx.org/>). Among these cis-eQTLs, more
209 than 58% had no classification as potential functional variants (Categories 1-5; Fig. 4A
210 and Supplemental Table 2). Moreover, only a small fraction of cis-eQTLs (~2.60%) in
211 each tissue were scored as high and moderate functional confidence variants (Category
212 1 and 2 variants; Fig. 4B,C). These results indicated that cis-eQTLs could primarily
213 serve as marker loci, but were unlikely to be functional variants that affect transcription.
214 In addition, this analysis provided a proof-of-concept that Functional Confidence
215 scoring system could be used to assess potential regulatory function in cis-eQTL
216 datasets and score for functional confidence.

217 We further validated Functional Confidence scoring system variant
218 identification and functional confidence scoring by genomic prediction with high and
219 moderate functional confidence variants (Category 1 and 2 variants) in pigs. We
220 assessed the predictive reliability of estimated breeding values (EBVs) for two traits,
221 average daily gain (ADG) and backfat thickness (BF) in a large white population
222 (n=874) using a genomic BLUP model with DMU software (38). EBVs were based on
223 four different genomic relationship matrices constructed by four scenarios of SNP
224 markers, including three scenarios using high and moderate functional confidence
225 variants from muscle, liver, or adipose, as well as one scenario that used 11k randomly
226 selected variants from whole genome sequence of pig (Table 4). Overall, the predictive
227 reliability of EBVs for ADG and trait BF was similar among the three scenarios using
228 high and moderate functional confidence variants (~0.31-0.38), whereas the predictive

229 reliability of EBVs was lowest in the scenario using 11K random SNPs (~0.27), despite
230 containing the highest number of SNP markers (11000). Notably, predictive reliability
231 was highest in the scenario based on variants detected in adipose (~0.38), despite using
232 the fewest markers (3861 SNPs). Predictive reliability of EBVs generated with
233 Category 1 and 2 variants from liver was higher than that of functional confidence
234 variants from muscle. Ultimately, the predictive reliability of EBVs increased 23%~46%
235 for the three tissue types by using high and moderate functional confidence variants
236 compared to EBVs based on randomly selected SNP markers. This analysis further
237 validated the use of Functional Confidence scoring system for screening functional
238 variants in genomic data of livestock.

239 **Development of the Integrated Functional Mutation database for screening
240 candidate functional variants in livestock species**

241 In order to facilitate screening for candidate functional variants in livestock species, we
242 integrated genomic variants with epigenomic datasets in a single database, the
243 Integrated Functional Mutation (IFmut) database. This database contains 65124531
244 potential functional variants from the genomes of pig, cattle, sheep, and chicken (Fig.
245 5A), as well as the 538 aforementioned epigenomic datasets from 19 tissues and 12 cell
246 lines across the four species (Fig. 5B). In addition, the IFmut database
247 (<http://www.ifmutants.com:8210/#/home>) has a user-friendly web interface that
248 enables users to query variants of interest, different genomic regions, or browse
249 epigenomic signal viewers.

250 In the first module, users can use a "Quick Search" function on the homepage to
251 search for a specific dbSNP ID or search specific genomic regions for a variant of
252 interest. Details, such as genomic location, conversion type, and functional confidence
253 score (defined in the following section) about the queried variant, if stored in the IFmut
254 database, are then listed at the bottom of the homepage (Fig. 5C). Clicking on an
255 SNVID in the search hits will direct the user to a new page containing information about
256 the regulatory feature(s) associated with queried variant of interest (Fig. 5D). In the

257 third module, users can search for “Affected motif” to facilitate hypothesis generation
258 about the potential effects of a variant on TF binding. Searches in this module return
259 logos plots of conservation of the potentially affected TF motif(s) and a table containing
260 the predicted effect on TF binding, and the affected gene symbol of the TF motif etc.
261 (Fig. 5E,F). In the fourth module, the “JBrowse” features allows users to view ATAC-
262 seq, Dnase-seq and ChIP-seq (H3K37ac and TFs) signals or Hi-C interaction heatmaps
263 (for Pig) around the variant of interest, as well as nearby genes (Fig. 5G). For this
264 purpose, each epigenomic dataset in the IFmut database is accompanied by BigWig and
265 genome annotation files that can be loaded in the right sidebar of JBrowse (Fig. 5G;
266 Supplemental Fig. 2), allowing users to examine epigenomic signals or annotation data
267 around queried variants in greater detail.

268 To facilitate further exploration of potential functional variants, IFmut also
269 provides hyperlinks to other databases: (i) For variants in pigs and cattle, users can click
270 on hyperlinked SNVIDs to access the IAnimal database (<https://ianimal.pro/>), which
271 contains additional information, such as genotype and major allele frequency (Fig. 5H).
272 (ii) Clicking on the “TAD/TAD Boundary” feature of IFmut entries that contain
273 topologically associating domain (TAD) information related to genomic variants in pig
274 will also direct users to the IAnimal database, allowing a subsequent search for genes
275 within that TAD or TAD boundary (Fig. 5I). (iii) Since ChromHMM Chromatin States
276 uses epigenomic information (such as ChIP-Seq data for various histone modifications)
277 across one or more human cell types to facilitate annotation of non-coding genome
278 regions, this function can be used for comparative genomics analysis to identify
279 regulatory feature-containing regions. The “ChromHMM Chromatin States” section in
280 the IFmut database can thus be used to map variant-containing genomic regulatory
281 regions in the four livestock species to corresponding chromatin regions in the human
282 hg38 genome (<https://genome-asia.ucsc.edu/>; Fig. 5J) by LiftOver (39).

283 It should be noted that IFmut also incorporates the details of functional
284 confidence scoring for each variant and provides the tool for scoring novel variants. For
285 such variants that are not yet included in IFmut, and are the subject of a user query, a
286 dialog box will prompt the user to categorize their variant using an embedded “Variant

287 scoring tool" (Fig. 6A). Upon clicking the "OK" button, a window is displayed
288 containing the classification results for the variant of interest (Fig. 6A). Queried SNPs
289 can also be loaded into JBrowse to visualize the relevant epigenetic data (Fig. 6B).

290 **Discussion**

291 Previous studies have shown that the majority of SNPs and small InDels are located in
292 non-protein-coding genomic regions (7, 40-44), and thus interpreting whether and how
293 a variant may affect function remains considerably challenging (45-47). Evaluating
294 perturbation effects of variants on TF binding sites in TF ChIP-seq data is a
295 demonstrably effective way for identifying potential functional variants in the human
296 genome (19). However, available TF ChIP-seq data is still comparatively lacking in
297 livestock, posing an obstacle for this approach of screening functional variants in
298 regulatory genomic features. To overcome this limitation, we compiled the IFmut
299 database of candidate SNP and small InDel functional variants in or near TF binding
300 sites in ATAC-seq/Dnase-seq and H3K27ac ChIP-seq datasets.

301 ATAC-seq/Dnase-seq analyses can largely capture TF binding footprints in full
302 range of open chromatin regions across the genome, and have been widely used for this
303 purpose in human and livestock research (48-52). At present, the TF binding sites
304 capturing in livestock were primarily relay on the ATAC-seq/Dnase-seq rather than the
305 TF ChIP-seq. Then our Functional Confidence scoring system used ATAC-seq/Dnase-
306 seq data to identify TF binding sites, which is different with the approach in
307 RegulomeDB based on TF ChIP-seq (19). Overall, our scoring approach was more
308 suitable the current study of functional mutations in livestock for abundant ATAC-
309 seq/Dnase-seq datasets in these species, as well as the design idea of using the ATAC-
310 seq/Dnase-seq data to identify TF binding sites to rank functional mutations can also
311 be transplanted to related research works on other species.

312 Further, our scoring approach identified five main categories of candidate
313 functional variants in pig, cattle, sheep, and chicken. We primarily focused on SNP and
314 InDel variants in the High and Moderate confidence groups (Categories 1 and 2), since

315 these variants were ranked based on their elevated likelihood of producing an effect in
316 livestock breeding and production. Genomic predictions with these High and Moderate
317 variants showed that Functional Confidence scoring could increase the predictive
318 reliability of EBVs in pigs compared to a larger set of randomly selected SNPs. These
319 findings suggested that our scoring system can guide the identification of important
320 variants, and could therefore drive advances in genetic improvement of livestock.

321 It is well-known that increasing the number of SNP markers can also increase
322 the predictive reliability of GEBVs (genomic EBVs (20)). Nevertheless, in this study,
323 we found that genomic prediction with 11000 random SNPs from across the pig genome
324 resulted in markedly lower GEBV reliability than that in some scenarios where even
325 only one third the number of High/Moderate confidence SNPs from IFmut were used.
326 Furthermore, this genomic prediction analysis also indicated that adipose tissue was
327 more strongly associated with average daily gain and backfat thickness than muscle or
328 liver. This finding might be at least partially explained by adipose function as the major
329 site of energy storage and insulation in pigs (53), and provides direct evidence that the
330 selection of candidate functional SNPs can guide genomic breeding efforts in pigs.

331 As variants play important roles in genomic breeding in livestock, a number of
332 sequencing-related databases have been developed for animal research, such as
333 AnimalQTLdb (54), Animal-ImputeDB (55), Animal-eRNADB (56), and IAnimal (57),
334 and range from one omics data type to comprehensive multi-omics data collections.
335 However, tools for identifying candidate functional variants, visualizing relevant
336 evidence in epigenetics data, and scoring for confidence in their function are still
337 unavailable for mining these databases. We therefore designed the IFmut platform to
338 allow users to retrieve and explore genomic, and epigenetic data related to the possible
339 function of a variant, as well as a Functional Confidence scoring tool for assessing new
340 variants of interest identified by users alongside those in IFmut and across multiple
341 livestock species. Overall, the Functional Confidence classification data for SNPs and
342 small InDels in the four species in IFmut, along with the tools for further exploration,
343 can facilitate investigations of functional impacts of variants

344 **Methods**

345 **Data collection**

346 Genome variants VCF data of pig (susScr11), cattle (bosTau9), and chicken (galGal5)
347 were downloaded from ensemble database (<http://ftp.ensembl.org/pub/>), genome
348 variants VCF data of sheep (oviAri4) was from NCBI Single Nucleotide Polymorphism
349 Database (https://ftp.ncbi.nih.gov/snp/organisms/archive/sheep_9940/VCF/00-All.VCF.gz). We also used whole-genome sequencing (WGS) data from 491 individual
350 pigs across 61 breeds generated in our previous study(29). QTL data of four livestock
351 were downloaded from Animal QTL database (<https://www.animalgenome.org/cgi-bin/QTldb/>). In cattle, we also downloaded the best variants cis-eQTL data from the
352 farmGTEEx database (https://cgtex.roslin.ed.ac.uk/wp-content/plugins/cgtex/static/rawdata/Full_summary_statisites_cis_eQTLs_FarmGTEEx_cattle_V0.tar.gz). The TF ChIP-seq, H3K27ac ChIP-seq, and ATAC-seq data in pig,
353 cattle, sheep, and chicken, Hi-C data in pig, and Dnase-seq in chicken were downloaded
354 from NCBI Sequence Read Archive (<http://www.ncbi.nlm.nih.gov/sra/>). A total of 579
355 raw epigenomic datasets were collected from multiple projects in NCBI, of which 75
356 datasets from pig were from our previous study (34).
357

361 **Sequencing data analysis**

362 To adhere to the ENCODE standard, we primarily refer to the analysis methods used in
363 our previous study for processing ChIP-seq and ATAC-seq data (34).

364 **ChIP-seq**

365 **Mapping and Quality control**

366 The ENCODE ChIP-seq pipeline (https://github.com/kundajelab/chipseq_pipeline)
367 was utilized to process the ChIP-seq datasets of the four species in a strict manner. The
368 raw reads from each dataset were aligned to the respective reference genome assemblies
369 (susScr11, bosTau9, oviAri4, galGal5) using BWA v0.7.17 (58). Subsequently, the
370 removal of low MAPQ reads (<25), unmapped reads, mate unmapped reads, not

371 primary alignment reads, and duplicate reads using Picard v1.126
372 (<https://broadinstitute.github.io/picard>) and SAMTools v1.9 (59).

373 The read coverage of genomic regions between replicate filtered BAM files was
374 computed using the multiBamSummary bins function of deepTools v2.0 (60). A bin
375 size of 2 kb was used to assess genome-wide similarities. The resulting read coverage
376 matrix obtained from the multiBamSummary step was used to calculate the Pearson
377 correlation coefficients between two replicate filtered BAM files. The non-duplicated
378 BAM file of replicates with a Pearson correlation coefficient ≥ 0.8 were merged, and
379 the remaining replicates with a correlation coefficient < 0.8 were excluded from further
380 analysis.

381 **Identification of nucleosome free region**

382 The HOMER (61) were utilized to detect nucleosome-free regions (NFR). The
383 makeTagDirectory command was used to generate tag directories for the H3K27ac IP
384 and input data using the merged non-duplicated BAM file obtained from the “Mapping
385 and Quality control” steps. Subsequently, the findPeaks command with the -nfr option
386 was applied to identify NFR peaks, requiring at least 10,000 peaks per data, and finally
387 excluding the scaffold regions.

388 **Identification of TF binding sites and H3K27ac narrow peaks**

389 The identification of TF binding sites and H3K27ac narrow peaks was carried out using
390 MACS2 v2.1.0 (62) and deepTools v2.0 (60), as described in greater detail in the
391 methods section of our previous study (34).

392 **ATAC-seq**

393 **Mapping, quality control and peak calling**

394 The ATAC-seq datasets of four species were processed following the ENCODE
395 ATAC-seq pipeline (https://github.com/kundajelab/atac_dnase_pipelines). The
396 preprocessing steps included checking and trimming adapters using Cutadapt v1.14
397 (<https://cutadapt.readthedocs.io/en/stable/>). The ATAC-seq reads were then aligned to
398 the susScr11, bosTau9, oviAri4, and galGal5 reference genome assemblies using

399 Bowtie2 v2.3.4.1. After alignment, low MAPQ reads (<25), unmapped reads, mate
400 unmapped reads, not primary alignments, reads failing platform, and duplicates were
401 removed using SAMTools v1.9 (59) and Picard v1.126
402 (<https://broadinstitute.github.io/picard>) software. The mitochondrial reads were further
403 removed from the mapped BMA file using BEDTools v2.26.0 (63) to generate effective
404 reads, which were subsequently used for peak calling. MACS2 v2.1.0 (62) was
405 employed to call peaks for each replicate individually, using parameters: genome size
406 (-g), p-value threshold (0.01), peak model (--nomodel), shift size (--shift), extension
407 size (--extsize), and other options (--B, --SPMR, --keep-dup all, --call-summits). And
408 generate a data set of at least 10,000 peaks for further analysis.

409 **Dnase-seq**

410 **Mapping, quality control and peak calling**

411 For the Dnase-seq datasets of chicken, the ENCODE Dnase-seq pipeline
412 (https://github.com/kundajelab/atac_dnase_pipelines) was followed. With the
413 'dnase_seq' parameter specified to indicate Dnase-seq data, and the others were
414 consistent with the above ATAC-seq analysis.

415 **Identification of open chromatin region**

416 In the peak calling step, peaks with $P < 10^{-5}$ were considered significant and selected
417 for further analysis. These significant narrow peaks were filtered based on replicates
418 with high Pearson correlation coefficients ($R > 0.8$). The peaks from these replicates
419 were merged using BEDTools v2.26.0 (63), requiring at least 50% overlap between
420 peaks in each replicate. The merged peaks represent open chromatin regions.
421 Furthermore, the BAM files from highly correlated replicates ($R > 0.8$) were merged to
422 generate signal tracks using MACS2 v2.1.0 (62). This step helps to visualize the signal
423 intensity and distribution of chromatin accessibility across the genome.

424 **Identification of footprints in ATAC-seq and Dnase-seq**

425 The footprint analysis was primarily performed as the following steps: (i) the board
426 peaks were called from the merged ATAC-seq or Dnase-seq data using the MACS2

427 v2.1.0 broad module (62, 64); (ii) the broad peaks meeting the criteria of $P < 10^{-10}$ and
428 $10^{-10} < P < 10^{-5}$ overlapping OCR were merged with BEDTools v2.26.0 (63) as
429 significant broad peaks; (iii) the Hmm-based IdeNtification of Transcription factor
430 footprints (HINT) framework of Regulatory Genomics Toolbox (RGT) v0.13.2 (65)
431 was employed to analysis footprints using the significant broad peaks. The HINT
432 framework was utilized with specific parameters depending on whether ATAC-seq or
433 Dnase-seq data was used (--atac-seq or --dnase-seq) and considering paired-end
434 sequencing data (--paired-end). The organism information (--organism=) was also
435 specified; and (iv) the cutoff value for footprint score was determined as more than the
436 20% quantile of all footprint score generated by the HINT framework of GRT v0.13.2
437 (65).

438 **Transcription factor motif mapping in genome function region**

439 The transcription factor motif mapping was primarily performed as the following steps:
440 (i) OCR, NFR, TF binding sites and footprint in OCR regions were merged into a BED
441 file; (ii) The fasta-get-markov command from the MEME Suite
442 (<https://github.com/cinquin/MEME>) software was used to generate a .fa.bg file and
443 “bedtools getfasta” command generate .fa file corresponding .bed file of step (i); (iii)
444 The fimo command in MEME Suite (--max-stored-scores 5000000) used to map motif
445 in the genome; and (iv) the fimo mapped results of pig, cattle, sheep filtered by $P <$
446 $5*10^{-6}$, and chicken filtered by Pvalue $< 5*10^{-7}$.

447 **Prediction of transcription factor motif effects**

448 In addition, potential functional variants (Categories 1-5) located within footprint
449 regions were analyzed using the motifbreakR (66) package in R v4.0. The motifDB
450 database, specifically JASPAR 2018 (67), was selected as the data source for predicting
451 the transcription factors to which the SNPs may bind.

452 **Hi-C**

453 The Hi-C data of two-week-old LW pigs were from our previous study (34), and the
454 other Hi-C data were downloaded from GEO under accession number GSE153452 at

455 <http://ncbi.nlm.nih.gov/geo>, including the cells of pig from zygotes, 4 cell stage and
456 morula of in vitro fertilization (IVF), and pig embryonic fibroblasts (PEFs). These
457 downloaded data were processed using the HiC-Pro (version 2.11.1) pipeline to
458 produce the ICE normalization contact matrices (68). The insulation score of the ICE
459 matrix was calculated by using the following options: -is 480000 -ids 320000 -im
460 iqrMean -ss 160000. Furthermore, the insulation method was utilized to define the
461 topologically associating domain (TAD) structure (insulation/boundaries).

462 **Variants distribution statistics**

463 SNPs and small InDels were annotated using ChIPseeker package in R v3.6.0, the
464 parameter of annotatePeak was setted that including level="transcript",
465 assignGenomicAnnotation=TRUE, genomicAnnotationPriority=c("Promoter",
466 "5UTR", "3UTR", "Exon", "Intron", "Downstream", "Intergenic"), annoDb=NULL,
467 addFlankGeneInfo=FALSE, sameStrand=FALSE, ignoreOverlap=FALSE,
468 ignoreUpstream=FALSE, ignoreDownstream=FALSE). Next, the reference genome
469 file (fasta) and annotation files (gtf) were used with the snpEff v4.5 software to predict
470 the effects of SNPs on known genes (java -Xmx8g -jar snpEff.jar genome -i .bed).

471 **The identification and filtering of pig variants.**

472 **Identification of SNP and small InDel in pig**

473 A total of 491 whole-genome sequences from 61 pig breeds were obtained from our
474 previous study (29). The method of data processing was consistent with our previous
475 article (34).

476 **Chromatin state discovery and characterization**

477 The chromatin states of human genome (hg38) were downloaded from the NIH
478 Roadmap Epigenomics program
479 (<https://egg2.wustl.edu/roadmap/data/byFileType/chromhmmSegmentations/ChmmModels/imputed12marks/>). The genome coordinates of human genome chromatin states
480 were converted into those of pig genome (SusScr11), chicken genome (GalGal5), cattle
481

482 genome (BosTau9), and sheep genome (OviAri4) by LiftOver, respectively. The
483 positions of SNPs from pig, chicken, cattle, and sheep were used to overlap with the
484 converted genome coordinates of chromatin states by BEDTools v2.26.0. In addition,
485 the converted genome coordinates of chromatin states with the same SNP were merged,
486 and the merged genome coordinates were transformed into those of human genome.

487 **Performances of genomic predictions**

488 **Dataset**

489 The phenotypic dataset used for genomic prediction were obtained from a national pig
490 nucleus herd in North China. In this study, we used phenotypic recordings for two
491 productive traits: 30-100 kg average daily gain (ADG) and 100 kg backfat thickness
492 (BF). All the phenotypic records for the traits were obtained at the same time point,
493 allowing a 10-kg deviation from the final bodyweight (100 ± 10 kg). All of the
494 phenotypes were recorded between the year early 2018 and October 2022. Based on the
495 traced pedigree, there were 11 lines existing in such pig population. For each pig line,
496 DNA samples were collected from about 80 distantly related pigs and were sequenced
497 by DNBSEQ-T7 platform with an averaged $5\times$ coverage. In total, 874 pigs were
498 sequenced. After quality controls, which includes a genotype missing rate below 10%,
499 a call rate of SNPs above 90%, and a minimum allele frequency (MAF) above 1%,
500 18460807 (18000K) SNPs were kept and analyzed in the following study. Missing
501 genotypes were imputed using software Beagle version 5.3. Among the 874 sequenced
502 pigs, 872 pigs had phenotypes of ADG data, meanwhile 867 pigs had BF recordings.
503 Environmental factors including such as genders, herds, and physical units were
504 completely recorded.

505 **Genomic Best Linear Unbiased Prediction (GLUP) models**

506 The breeding values (EBV) for different traits were estimated using the following
507 GBLUP models:

508
$$\mathbf{y} = \mathbf{X}\mathbf{b} + \mathbf{Z}\mathbf{u} + \mathbf{e},$$

509 where \mathbf{y} represents a column vector of phenotypic values for each trait; \mathbf{b} represents a

510 vector of fixed effects, including sex effect, herd effect and physical units effects; \mathbf{u}
511 represents a vector of random additive genetic effects; \mathbf{e} represents a vector of residual
512 effects. Matrices \mathbf{X} and \mathbf{Z} are corresponding design matrices associated with these
513 effects. The GBLUP model assumes a normal distribution for the random additive
514 effects and residual effects, as $\mathbf{u} \sim N(0, \mathbf{G}\sigma_u^2)$, where \mathbf{G} is genomic relationship matrix
515 constructed as Vanraden method 1; $\mathbf{e} \sim N(0, \mathbf{I}\sigma_e^2)$, where \mathbf{I} is an identity matrix. The
516 additive genetic variance and residual variance are denoted by σ_u^2 and σ_e^2 , respectively.

517 **Scenarios of constructing genomic relationship matrices**

518 In this study, GBLUP models with four different genomic relationship matrices (\mathbf{G})
519 were used to estimate the GEBVs for both ADG and BF traits. Four different sets of
520 SNP markers were used for constructing the corresponding \mathbf{G} matrix. In scenario 1,
521 sequenced SNP markers that were with top 1 and top 2 muscle scores (1+2 muscle,
522 10544 SNPs) were calculated the \mathbf{G} matrix. Similarly, sequenced SNP markers that
523 were with top 1 and top 2 liver scores (1+2 liver, 6049 SNPs) and with top 1 and top 2
524 adipose scores (1+2 adipose, 3801 SNPs) were used for constructing \mathbf{G} matrices in
525 scenarios 2 and 3, respectively. In scenario 4, 11000 (11K) randomly selected SNP
526 markers were used for constructing \mathbf{G} matrix. Scenario 4 were repeated for three times
527 in the study.

528 **Predictive Reliabilities**

529 The mean predictive reliabilities of GEBVs were determined by employing the
530 subsequent formula (Mrode, 2005):

$$531 \quad r^2 = \sum \left(1 - \frac{SEP_i^2}{\sigma_g^2} \right) / N,$$

532 where r^2 is reliability of GEBVs and i denotes an individual animal i ; SEP represents
533 the standard error that is associated with the predicted GEBVs; σ_g^2 represents the
534 additive genetic variance and N is the number of used animals.

535 **Data access**

536 All track of ATAC-seq, ChIP-seq (H3K27ac and TFs) and Hi-C, as well as the candidate
537 functional variants and their Functional Confidence score are available at

538 <http://www.ifmutants.com:8210/#/home>.

539 **Competing interest statement**

540 The authors declare no competing interests.

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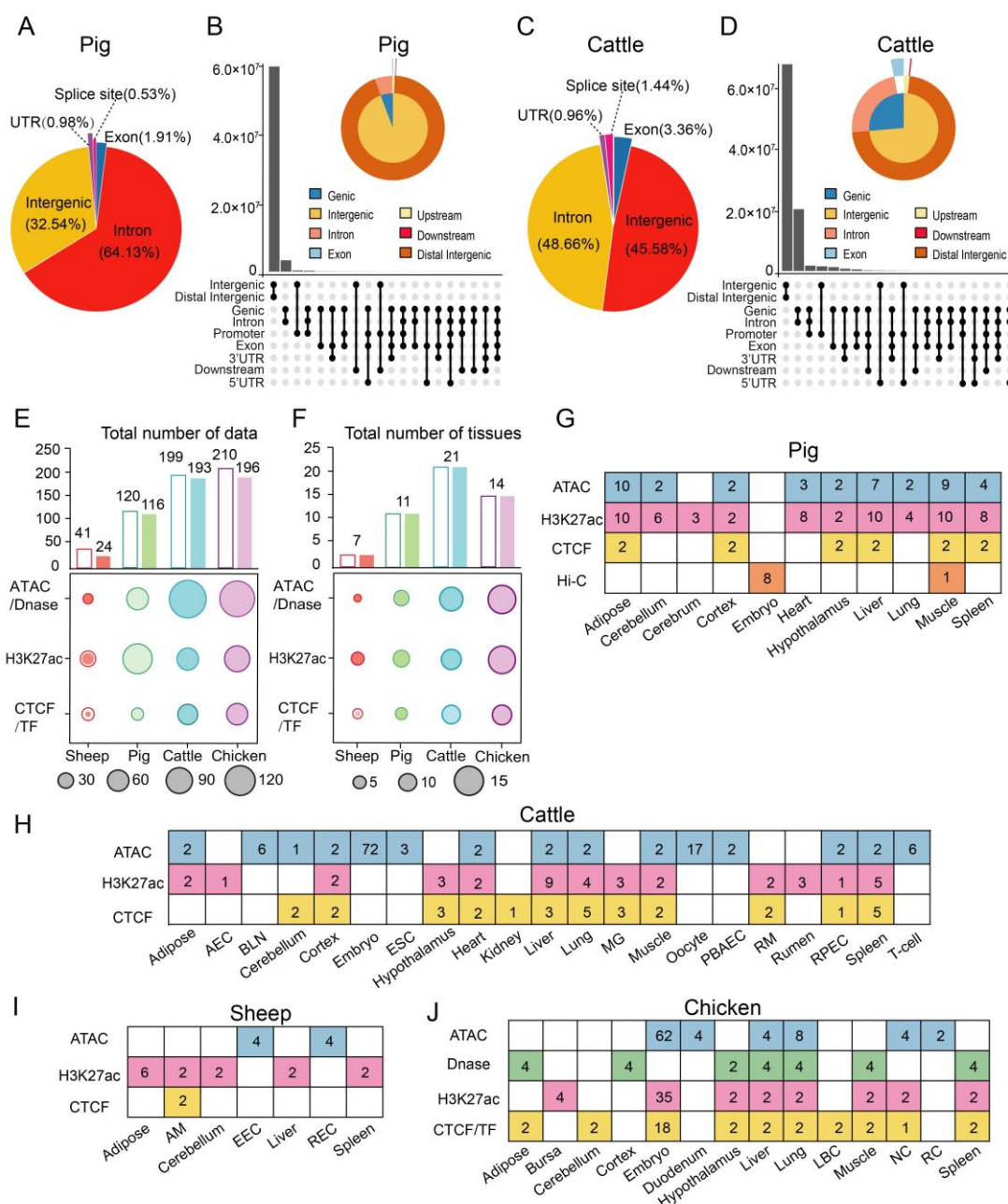
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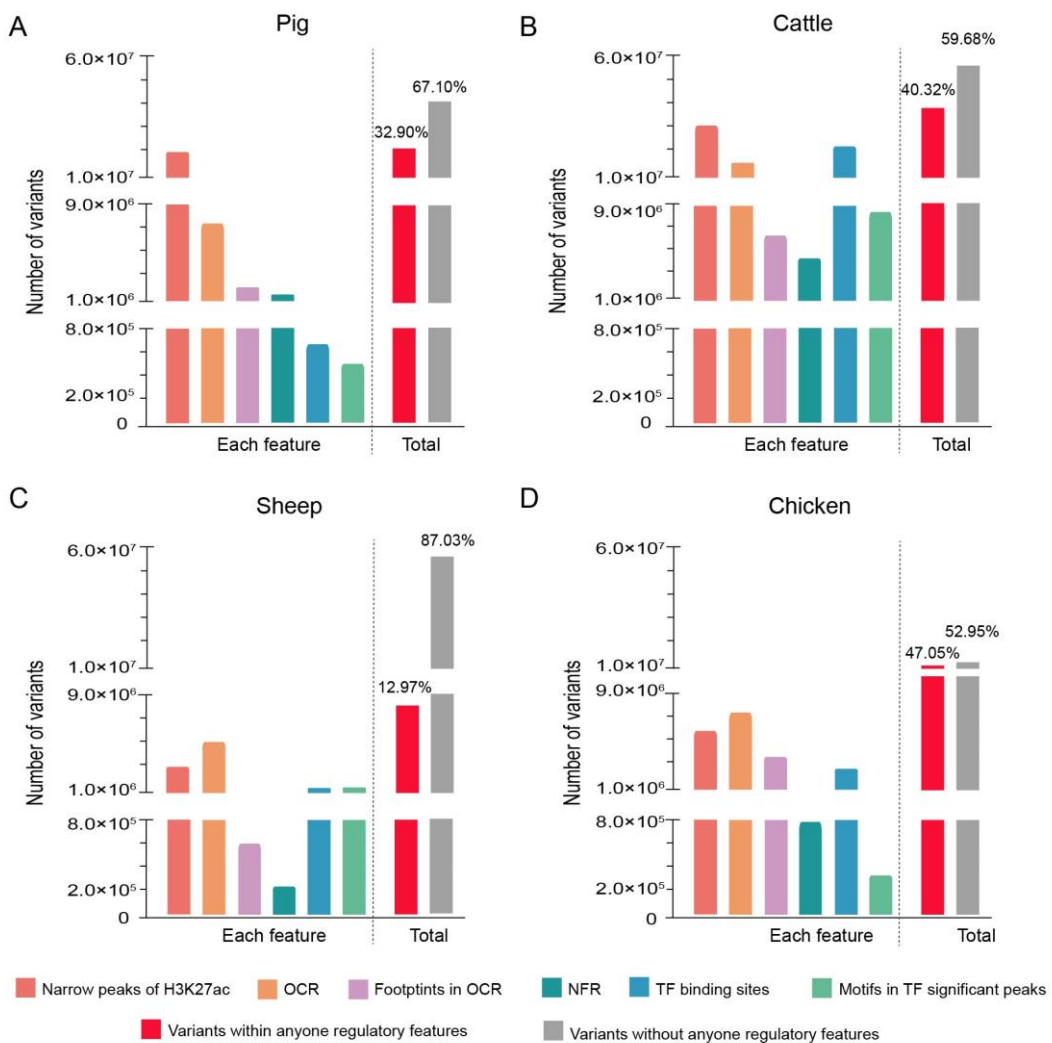
717

718 **Figures and Tables**



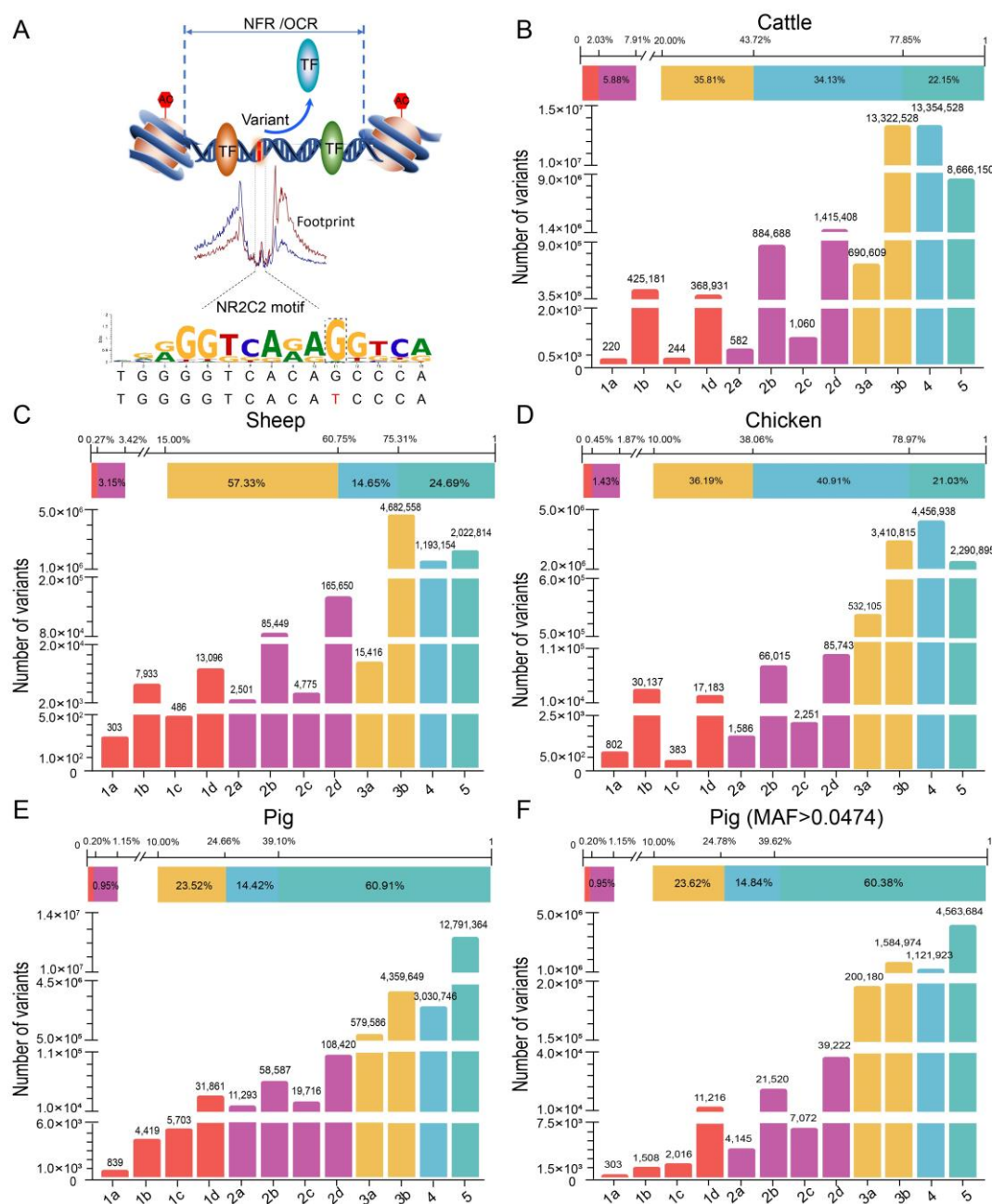
719 **Figure 1.** Genomic distribution of variants and their predicted effects on annotated genes and
720 epigenomic datasets collection in livestock. **(A)** Percent distribution of variants in different
721 regions in the susScr11 reference genome. **(B)** Regions within genome predicted to be affected
722 by variants based on annotated genes in the susScr11; black dots indicate regions within a gene
723 predicted to be simultaneously affected a variant. **(C)** Distribution of variants in the bosTau9
724 reference genome assembly. **(D)** Predicted effects of cattle variants based on annotated genes in the bosTau9
725 genome assembly. **(E)** Statistical summary of epigenomic datasets for the four species. Histogram
726 of total numbers of datasets obtained for each species; empty columns (left) are raw data and
727 filled columns (right) show number of datasets after filtering and quality control. Bubble size
728 represents number of different epigenetics datasets; outer circles, raw data; inner circles, cleaned

729 data. (F) Statistical summary of epigenetics datasets for different tissue types in the four species.
730 Empty histograms (left) are number of tissue types with raw datasets; filled histograms (right) are
731 number of tissue types with cleaned datasets. Bubble size indicates number of different tissues
732 represented in each data type; outer circles are raw datasets; inner circles are cleaned datasets.
733 (G-J) Summary of different quality-controlled epigenomic datasets and represented tissue types
734 in (G) pig, (H) cattle, (I) sheep, and (J) chicken. AEC, aortic endothelial cells; BLN, bronchial
735 lymph node; ESC, embryonic stem cells; MG, mammary gland; PBAEC, primary bovine aortic
736 endothelial cells; RM, renal medulla; and RPEC, rumen primary epithelial cell. AM, alveolar
737 macrophage; EEC, esophagus epithelium cells; and REC, rumen epithelium cells. LBC,
738 lymphoma B-cell; NC, neural crest; RC, retinal cell.

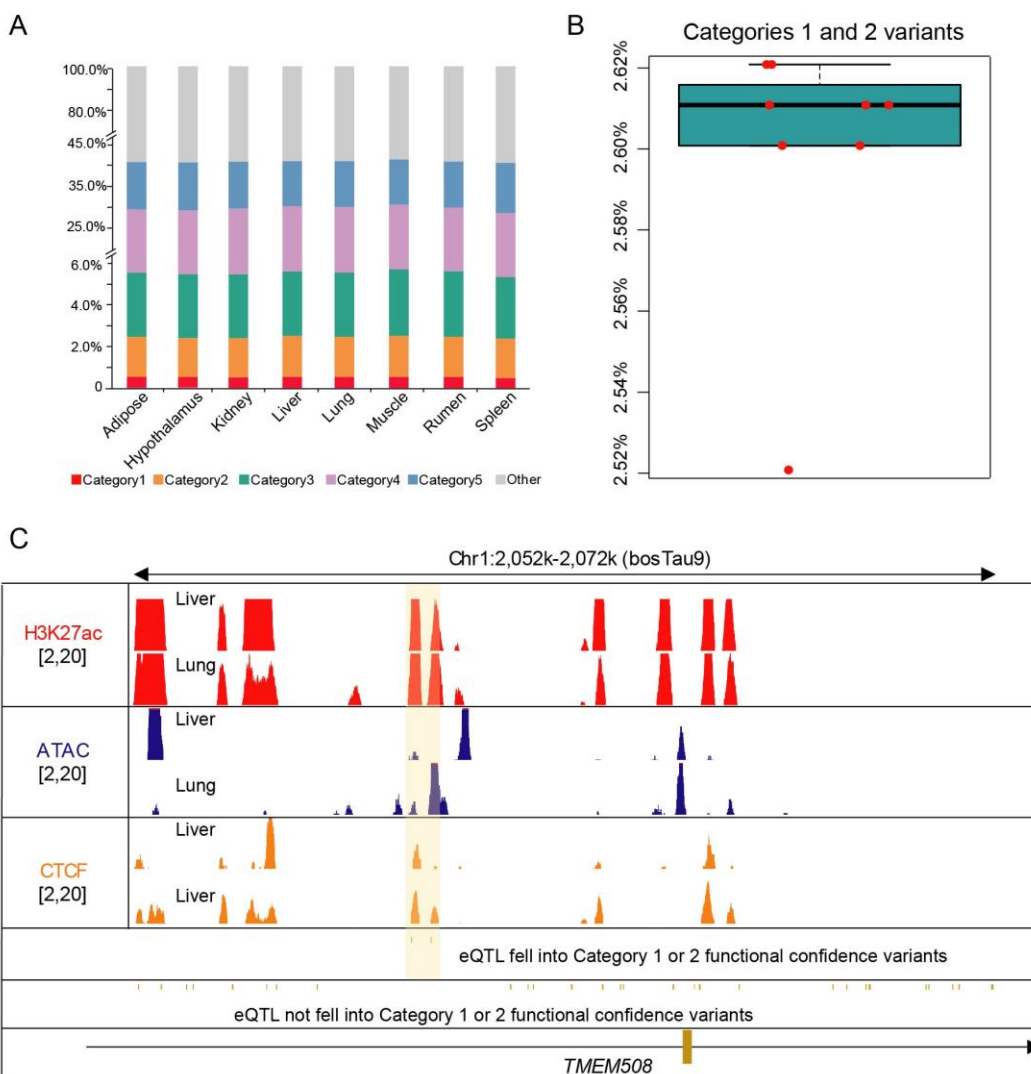


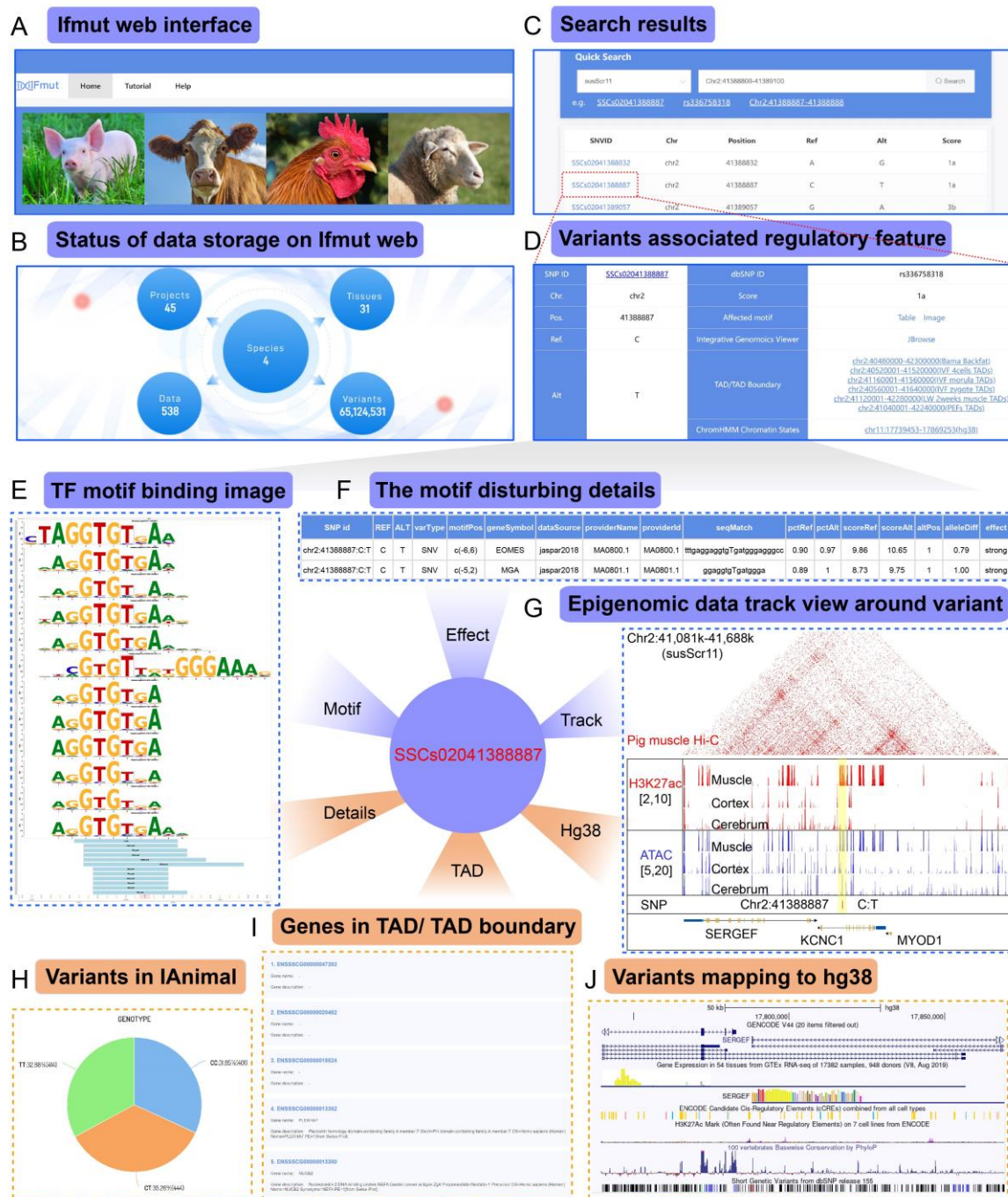
739 **Figure 2.** Statistical summary of candidate functional variants and their distribution in genomic
740 regulatory features in four livestock species. (A) Total number of candidate functional variants
741 distributed in each of 7 regulatory features, including narrow peaks in H3K27ac, open chromatin
742 region (OCR), footprints in OCR, nucleosome free regions (NFR), recognition motifs in
743 significant transcription factor (TF) peaks in ChIPseq, and TF binding sites, or associated with
744 no regulatory features in epigenomic data from (A) pig, (B) cattle, (C) sheep, and (D) chicken.

745 Figure 3 revised



746 **Figure 3.** Confidence scoring of candidate functional variants. (A) Illustration of design principle
747 of Functional Confidence scoring system. (B-E) Statistical summary of candidate functional variant
748 distribution among confidence subcategories in (B) cattle, (C) sheep, (D) chickens, and (E) pigs.
749 Bar at the top shows the proportional distribution of main confidence categories among total
750 candidate functional variants for each livestock species. (F) Number of candidate functional variants
751 in each subcategory filtered by minor allele frequency (MAF>0.047) from 491 whole genome
752 sequencing datasets in pigs.





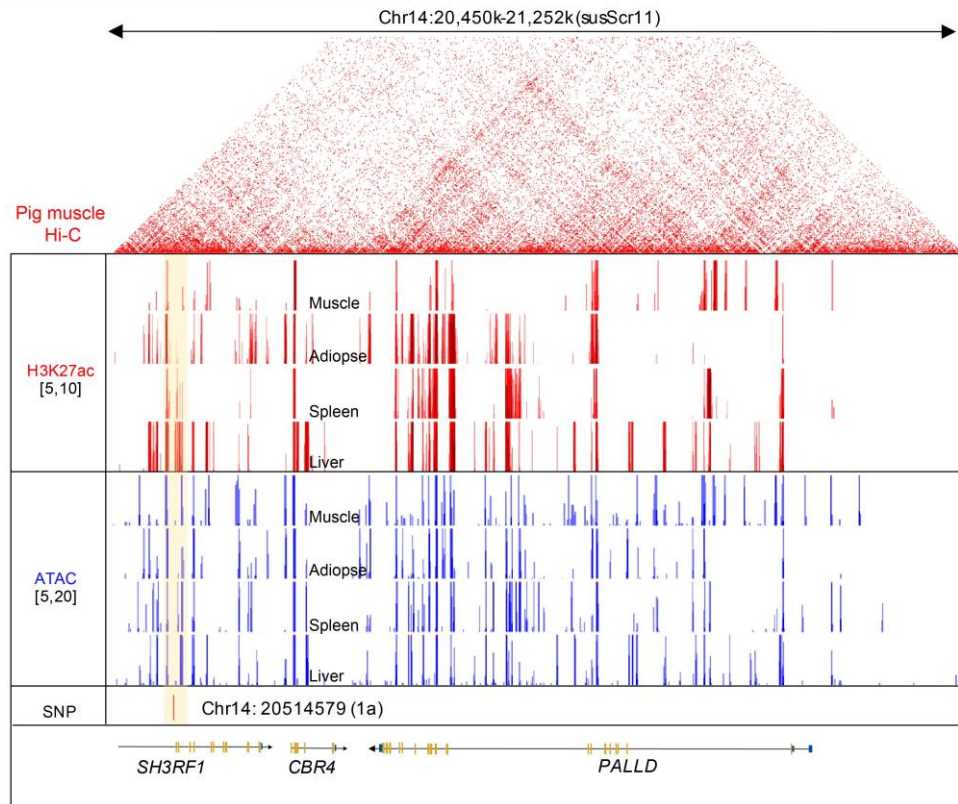
759 **Figure 5.** Overview of the Integrated Functional Mutation (IFmut) database. (A) IFmut home
760 page, containing information about the four livestock species. (B) Graphic summary of datasets
761 available in IFmut organized by search type. (C) Example of search results generated with the
762 "Quick Search" function, including variant chromosomal location, conversion type, and
763 confidence score. (D) Variant-associated regulatory features in the SNVID query results of Quick
764 Search are linked to pages containing information such as motif affecting, TAD/boundary, and
765 ChromHMM of human genome conservation region, which are further linked to source data,
766 external databases, etc. (E) Clicking the "Image" link in the "Affected motif" column in (D) takes
767 the user to logos plots of nucleotide conservation in TF recognition motif(s) potentially affected
768 by a queried variant. (F) Clicking "Table" in the "Affected motif" column in (D) takes the user
769 to a page containing the predicted effect on TF binding, and the affected gene symbol of the TF

770 motif etc. details about the potentially affected TF motif(s). *(G)* The track view of variant through
771 "JBrowse" function in *(D)*. This function will bring the user to track views and feature
772 visualization for regions containing the queried variants in ATAC-seq and ChIP-seq (H3K37ac)
773 data, Hi-C interaction heatmaps (for pig), and nearby genes. *(H)* Clicking on SNVID hyperlinks
774 in *(D)* brings the user to the IAnimal database (<https://ianimal.pro/>) to obtain additional
775 information, such as genotype and major allele frequency in pig or cattle. *(I)* A subsequent search
776 for genes within topologically associating domains (TADs) or TAD boundaries that contain user
777 queried variant provides links to information about those genes in the IAnimal database
778 (<https://ianimal.pro/>). *(J)* Users can also perform comparative genomics between predicted
779 variant-affected regions in livestock and corresponding chromatin regions in the human hg38
780 reference genome mapped using LiftOver.

A **Epigenomic data track view around variant**



B **Functional confidence scoring variant**



781 **Figure 6.** Functional confidence scoring and epigenomic data visualization functions in IFmut for
782 user analysis of novel candidate variants. (A) Functional Confidence Scoring tool in IFmut. For
783 variants of interest not stored in IFmut, users are prompted with the option to conduct Functional
784 Confidence scoring using the tool in IFmut. (B) Epigenomic data visualization to assess novel
785 variants. To examine the evidence underlying the IFmut functional confidence score for a variant of
786 interest, users can follow a link to the JBrowse tool showing epigenomic tracks, TAD regions, and
787 nearby genes.

Table 1. Number of regulatory features detected in livestock species.

Regulatory feature	pig	cattle	sheep	chicken
Basic regulatory feature				
NFR	200,450	328,909	58,655	103,504
OCR	441,818	780,224	418,560	600,364
Narrow H3K27ac peaks	163,307	162,841	57,167	69,527
Footprints in OCR	2,318,229	4,002,427	978,982	3,157,063
Total non-redundant	352,763	804,593	474,751	1,026,316
TF binding site-related features				
Footprint-matching motifs	125,500	1,352,480	253,476	1,098,796
Partial motif-containing footprints	198,272	1,569,613	374,664	680,374
Motifs in significant peaks	1,354,404	24,540,298	15,242,424	2,962,352
TF binding sites	43,007	182,643	136,538	135,362
Total non-redundant	451,212	2,783,299	2,443,102	428,545

NFR, nucleosome-free regions; OCR, open chromatin regions.

Table 2. Genomic coverage of regulatory features.

Regulatory feature	pig	cattle	sheep	chicken
Basic regulatory feature (bp)				
NFR	60,641,955	95,978,174	9,085,698	33,571,036
OCR	300,233,553	324,314,161	186,787,025	309,414,106
Narrow H3K27ac peaks	700,812,854	608,126,611	117,598,663	229,512,492
Footprints in OCR	93,287,631	127,134,565	24,974,332	151,709,670
Total non-redundant	787,386,263	794,764,318	287,920,733	441,600,747
TF binding site-related features (bp)				
Footprint-matching motifs	3,295,980	22,900,815	4,146,394	18,530,843
Partial motif-containing footprints	7,771,203	46,493,819	9,943,498	30,423,419
Motifs in significant peaks	20,463,073	135,948,968	53,057,694	50,400,477
TF binding sites	31,158,919	332,861,617	55,897,125	100,867,511
Total non-redundant	51,963,305	391,146,808	104,501,330	107,662,441

NFR, nucleosome-free regions; OCR, open chromatin regions.

Table 3. Variant classification scheme for scoring system in IFmut database.

Categorization Scheme	
Category	Description
	High possibility of affecting transcription factor binding
1a	QTL + OCR + NFR + Footprints in OCR + Footprint-matching motifs
1b	OCR + NFR + Footprints in OCR + Footprint-matching motifs
1c	QTL + OCR + NFR + Part motif-containing footprints
1d	OCR + NFR + Part motif-containing footprints
	Moderate possibility of affecting TF binding
2a	QTL + OCR + Footprints in OCR + Footprint-matching motifs
2b	OCR + footprint in OCR+ Footprint-matching motifs
2c	QTL + OCR + Part motif-containing footprints
2d	OCR + Part motif-containing footprints
	Low possibility of affecting TF binding
3a	OCR + NFR + Footprints in OCR
3b	OCR / NFR + Motifs in significant peaks
	Minimal possibility of affecting TF binding
4	OCR / NFR / Footprints in OCR / TF binding significant peaks
	Likely to be associated with gene expression
5	H3K27ac significant peaks

QTL, quantitative trait loci; NFR, nucleosome-free regions; OCR, open chromatin regions

Table 4. Predictive reliability of EBV for traits ADG and BF.

Scenarios	SNP numbers	ADG	BF
11k random	11,000	0.268 (0.014)	0.265 (0.015)
muscle	10,544	0.319	0.316
liver	6,049	0.348	0.346
adipose	3,801	0.380	0.378

Numbers within the parentheses are the standard errors of the reliability.