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Peer reviewed

# **RESEARCH ARTICLE**

# A comparative analysis of chromatin accessibility in cattle, pig, and mouse tissues

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### Abstract

**Background:** Although considerable progress has been made towards annotating the noncoding portion of the human and mouse genomes, regulatory elements in other species, such as livestock, remain poorly characterized. This lack of functional annotation poses a substantial roadblock to agricultural research and diminishes the value of these species as model organisms. As active regulatory elements are typically characterized by chromatin accessibility, we implemented the Assay for Transposase Accessible Chromatin (ATAC-seq) to annotate and characterize regulatory elements in pigs and cattle, given a set of eight adult tissues.

**Results:** Overall, 306,304 and 273,594 active regulatory elements were identified in pig and cattle, respectively. 71, 478 porcine and 47,454 bovine regulatory elements were highly tissue-specific and were correspondingly enriched for binding motifs of known tissue-specific transcription factors. However, in every tissue the most prevalent accessible motif corresponded to the insulator CTCF, suggesting pervasive involvement in 3-D chromatin organization. Taking advantage of a similar dataset in mouse, open chromatin in pig, cattle, and mice were compared, revealing that the conservation of regulatory elements, in terms of sequence identity and accessibility, was consistent with evolutionary distance; whereas pig and cattle shared about 20% of accessible sites, mice and ungulates only had about 10% of accessible sites in common. Furthermore, conservation of accessibility was more prevalent at promoters than at intergenic regions.

**Conclusions:** The lack of conserved accessibility at distal elements is consistent with rapid evolution of enhancers, and further emphasizes the need to annotate regulatory elements in individual species, rather than inferring elements based on homology. This atlas of chromatin accessibility in cattle and pig constitutes a substantial step towards annotating livestock genomes and dissecting the regulatory link between genome and phenome.

Keywords: Chromatin accessibility, Functional annotation, Cattle, Pig, Comparative epigenomics

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### Background

Despite considerable progress to annotate protein-coding genes in livestock species, the vast majority of these genomes is noncoding and remains poorly characterized. Epigenomics techniques, such as chromatin immunoprecipitation followed by sequencing (ChIP-seq) and DNase I hypersensitive sites sequencing (DNase-seq), have been extensively employed to catalog functional elements in humans [1] and classical model organisms [2–6]. For instance, the international human epigenome consortium has profiled thousands of epigenomes and identified millions of regulatory elements in the human genome [1, 7, 8], yielding an atlas of functional elements that has been invaluable for subsequent research in a wide variety of biological processes, including disease [9–13], pluripotency [14–16], differentiation [17–19], and morphology [20, 21].

Ultimately, genome-wide patterns of chromatin accessibility and compaction determine which genomic regions are available to cellular machinery, and are thereby intimately connected to the cell-specific gene expression patterns that determine identity and function. Controlled exposure of specific sites provides opportunities for transcription factors to bind their recognition motifs and regulate gene expression through further recruitment of proteins, such as RNA polymerases [22, 23]. Consequently, profiling open chromatin has the potential to not only identify regulatory elements, but also profile their activities in different cell types. The increasing availability of nextgeneration sequencing-based techniques spurred development of several alternative epigenomics assays, such as the Assay for Transposase Accessible Chromatin (ATAC-seq), which was first reported by Buenrostro et al in 2013 [24]. Following its introduction, ATAC-seq quickly became one of the leading methods for identification of open chromatin, largely due to the simplicity of the technique and low input requirements, which made it possible to study chromatin structure in rare samples.

Here we implemented ATAC-seq to profile open chromatin in a set of cattle and pig tissues: subcutaneous adipose, brain (frontal brain cortex, hypothalamus, and cerebellum), liver, lung, skeletal muscle, and spleen. This set of prioritized tissues are associated with a large number of qualitative phenotypic traits relevant to animal production, such as disease resistance, growth, and feed efficiency. Overall, about 300,000 accessible regions were identified in each species, yielding an epigenomic resource that will benefit agricultural genomics research and enable cross-species comparisons that will enhance knowledge of comparative epigenomics and transcriptional regulation.

### Results

### ATAC-seq library quality control and preprocessing

Using a modified ATAC-seq protocol, genome-wide chromatin accessibility was profiled in eight tissues derived from two adult male Hereford cattle and two adult male Yorkshire pigs: three brain tissues (frontal brain cortex, hypothalamus, and cerebellum), liver, lung, spleen, subcutaneous adipose, and skeletal muscle (Fig. 1a). With the exception of one cattle cerebellum sample, which was lost during processing, ATAC-seq data were generated for two biological replicates per tissue (Table S1). In addition, two technical replicates were prepared for cattle cortex, pig cerebellum, and pig hypothalamus. ATAC-seq signal from technical replicates were highly correlated (Pearson R averaged 0.97), and principal components analysis (PCA) of genome-wide signal grouped biological and technical replicates together (Fig. S1).

Mapping of sequencing data resulted in  $58 \pm 4$  million (S.E.) informative reads (uniquely mapping, nonmitochondrial, monoclonal reads) per sample (Table 1). Normalized genome-wide ATAC-seq signal was highly reproducible between biological replicates, with the Pearson correlation coefficient averaging 0.97 (Fig. 1b; Fig. S2), and increased in intensity at transcription start sites (TSS) (Fig. 1c; Fig. S3). Genes with tissue-specific functions demonstrated open chromatin that was specific to the corresponding tissue; for instance, the gene STMN4, which is involved in neuron projection development, is specifically marked by open chromatin at two sites in both cattle (Fig. 1d) and pig (Fig. 1e) in all three brain tissues, which likely correspond to alternate TSS, based on the gene annotation in pig. PCA of normalized ATAC-seq signal separated samples by tissue, with brain tissues grouping together in both pig and cattle (Fig. 1f,g, Fig. S4).

Several commonly used statistics were used to evaluate library quality (Table 2) [25, 26]. The non-redundant read fraction (NRF), which gauges library complexity by measuring the proportion of non-duplicate uniquely mapped reads out of all mapped reads, averaged  $0.62 \pm$ 0.03 (S.E.), indicating acceptable library complexity. The synthetic Jensen-Shannon distance (sJSD), which measures the divergence between the genome-wide ATACseq signal in a given sample versus a uniform distribution, averaged  $0.46 \pm 0.01$  (S.E.), suggesting a nonrandom ATAC-seq signal distribution throughout the genome. Finally, the Fraction of Reads in Peaks (FRiP) was calculated to evaluate the strength of signal over background. On average, 130,712 ± 10,994 (S.E.) ATAC-seq peaks (regions of enrichment) were called per sample (Supplementary Data 1-2), and the FRiP score averaged  $34.8 \pm 2.6\%$  (S.E). Notably, FRiP scores for adipose libraries were consistently lower (average  $8.5 \pm 2.5\%$ ) than for other tissues (average  $38.7 \pm 2.1\%$ ).

For each tissue, peaks called from biological replicates were compared to evaluate consistency between replicates and identify accessible regions with high confidence. On



at the *GAPDH* locus (a housekeeping gene), as well as several genes with tissue-specific activity. Pink bars indicate that signal exceeded the viewing range (up to 20 RPM). **e** Normalized ATAC-seq signal in pig tissues at the same genes. **f** PCA of normalized ATAC-seq signal in consensus open chromatin identified in cattle. G) PCA of normalized ATAC-seq signal in consensus open chromatin identified in pig

average,  $67.4 \pm 17.8\%$  (S.E.) of peaks from the replicate with fewer peaks called were also identified in the other replicate (Table 3). Regions that were enriched for ATACseq signal in both biological replicates of at least one tissue were collapsed to obtain a single comprehensive set of unique "consensus" peaks, accounting for accessibility in all eight tissues. Altogether, 306,304 and 273,594 consensus peaks were identified in pig and cattle, respectively (Table 4).

# Global characteristics of accessible chromatin in cattle, pig, and mouse

To infer the functional significance of accessible regions that were identified in pig and cattle tissues, consensus peaks were characterized by genomic localization (positioning relative to genes), sequence content, and tissuespecificity. In cattle, consensus peaks averaged 616 bp in width (Table 4) and covered 6.2% of the genome. Similarly, pig consensus peaks were 624 bp wide (Table 4)

Table 1 ATAC-seq data preprocessing. Per	library, total raw reads, r	napped reads (excluding r	nitochondrial DNA), duplicate reads,
informative reads (monoclonal and uniquel	y mapping), and percen	t of raw reads that were ir	nformative

Species	Tissue	Replicate	Raw reads	Mapped reads	Duplicate reads	Informative reads	% Raw
Cattle	Adipose	А	152,184,070	145,648,161	82,972,324	37,815,392	24.85
	Adipose	В	126,920,776	123,879,749	25,466,938	72,623,985	57.22
	Cerebellum	В	158,964,554	134,391,900	53,303,886	49,305,250	31.02
	Brain Cortex	А	216,202,716	189,763,599	122,567,574	30,997,045	14.34
	Brain Cortex	В	205,363,760	174,961,995	96,411,128	41,021,487	19.98
	Hypothalamus	А	146,439,048	102,771,471	50,044,176	35,090,140	23.96
	Hypothalamus	В	68,153,356	63,771,723	31,495,325	15,328,535	22.49
	Liver	А	175,124,072	165,112,503	95,001,248	40,031,560	22.86
	Liver	В	194,367,992	179,265,341	99,942,567	36,472,289	18.76
	Lung	А	163,469,102	159,710,295	27,682,063	99,691,291	60.98
	Lung	В	190,030,170	184,571,092	36,657,121	110,584,502	58.19
	Muscle	А	89,174,618	86,045,857	15,262,767	55,794,693	62.57
	Muscle	В	97,247,280	69,592,640	10,539,048	45,909,945	47.21
	Spleen	А	261,785,316	250,911,567	163,860,741	37,521,241	14.33
	Spleen	В	179,854,284	162,449,025	51,533,284	69,626,100	38.71
Pig	Adipose	А	93,118,998	87,025,520	15,814,107	57,651,677	61.91
	Adipose	В	71,639,956	66,597,412	12,368,404	42,960,583	59.97
	Cerebellum	А	186,388,542	175,280,070	90,013,906	63,538,783	34.09
	Cerebellum	В	125,817,350	116,536,743	41,557,703	57,997,395	46.10
	Brain Cortex	А	101,924,240	98,384,411	36,110,627	53,096,952	52.09
	Brain Cortex	В	160,871,726	155,783,257	77,377,043	66,403,810	41.28
	Hypothalamus	А	112,463,726	106,966,835	52,803,730	39,919,895	35.50
	Hypothalamus	В	170,163,006	162,907,052	92,303,710	56,125,160	32.98
	Liver	А	171,864,386	167,321,556	113,588,699	42,376,220	24.66
	Liver	В	169,952,062	164,205,920	85,279,200	64,117,458	37.73
	Lung	А	108,086,464	104,424,556	19,658,156	73,018,749	67.56
	Lung	В	110,690,180	106,275,981	17,791,030	74,869,852	67.64
	Muscle	А	168,211,474	165,225,305	41,747,860	113,058,649	67.21
	Muscle	В	141,069,686	138,785,466	39,577,315	91,780,486	65.06
	Spleen	A	91,549,652	88,698,199	13,254,769	64,352,893	70.29
	Spleen	В	93,747,292	90,558,895	12,817,045	67,241,934	71.73

and accounted for 7.2% of the genome. As expected, open chromatin was significantly enriched near TSS (p-value <1e-200), although most consensus peaks were intronic or intergenic (Fig. 2a). In addition, consensus peaks frequently occurred in only one tissue (54 and 58% of cattle and pig peaks, respectively) (Fig. 2b).

A comparable mouse ATAC-seq dataset, which included libraries from two male replicates for all tissues except hypothalamus, was downloaded from the CNGB Nucleotide Sequence Read Archive (Project ID CNP0000198) and processed in the same manner as the pig and cattle data. Similar to cattle and pig, 254,076 consensus peaks were identified from mouse tissues (Table 4). Mouse consensus peaks also covered a comparable portion of the genome (6.2%), were of similar width (average 668 bp), demonstrated enrichment at TSS (Fig. S5a), and were often tissue-specific (63% of peaks) (Fig. S5b). Surprisingly, a higher fraction of mouse consensus peaks localized to TSS (9.6%) than in cattle (5.1%) or pig (5.6%) (Fig. 2a, Fig. S5a). This discrepancy could be attributed either to the protocol, as pig and cattle ATAC-seq libraries were subjected to size-selection prior to sequencing and mouse libraries were not, or suboptimal genome annotations, as the pig and cattle annotations are relatively incomplete in comparison to the mouse. Nevertheless, the global characteristics of open chromatin were generally consistent across these three species.

**Table 2** Quality metrics of ATAC-seq libraries. Non-redundant read fraction (NRF) measures library complexity, Fraction of reads in peaks (FRiP) measures signal-to-noise ratio, and synthetic Jensen-Shannon distance (sJSD) measures divergence between ATAC-seq signal and a uniform distribution

Species	Tissue	Replicate	NRF	FRiP	sJSD
Cattle	Adipose	А	0.43	9.82	0.34
	Adipose	В	0.79	15.04	0.34
	Cerebellum	В	0.60	42.53	0.48
	Brain Cortex	А	0.35	30.16	0.48
	Brain Cortex	В	0.45	40.80	0.49
	Hypothalamus	А	0.51	43.67	0.50
	Hypothalamus	В	0.51	11.97	0.37
	Liver	А	0.42	37.99	0.50
	Liver	В	0.44	28.17	0.44
	Lung	А	0.83	37.63	0.45
	Lung	В	0.80	44.26	0.48
	Muscle	А	0.82	40.77	0.50
	Muscle	В	0.85	41.77	0.51
	Spleen	А	0.35	21.29	0.42
	Spleen	В	0.68	35.24	0.45
Pig	Adipose	А	0.82	5.21	0.40
	Adipose	В	0.81	4.12	0.40
	Cerebellum	А	0.49	43.05	0.46
	Cerebellum	В	0.64	41.49	0.47
	Brain Cortex	А	0.63	40.65	0.48
	Brain Cortex	В	0.50	44.53	0.46
	Hypothalamus	А	0.51	36.85	0.49
	Hypothalamus	В	0.43	37.76	0.44
	Liver	А	0.32	55.38	0.52
	Liver	В	0.48	49.25	0.49
	Lung	A	0.81	31.21	0.43
	Lung	В	0.83	36.87	0.46
	Muscle	A	0.75	58.16	0.62
	Muscle	В	0.71	61.94	0.65
	Spleen	А	0.85	29.41	0.43
	Spleen	В	0.86	22.70	0.38

To interrogate the potential function of accessible regions in cattle and pig, consensus peaks were subjected to motif enrichment analysis. Overall, consensus peaks were most significantly enriched for CTCF recognition sites, with about 8% of accessible regions harboring CTCF motifs in each species (Table 5). Of note, CTCF motifs were more prevalent in consensus peaks identified in 3 or more tissues (8% of peaks in cattle and pig) than in consensus peaks identified in only 1 or 2 tissues (3 and 2% of peaks in cattle and pig, respectively). Nevertheless, in both species 30% of consensus peaks containing a CTCF motif were only accessible in a single tissue (Fig. 2c), indicating that CTCF binding could play both tissue-specific and ubiquitous regulatory roles.

The unique open chromatin landscapes present in different cell types are crucial for regulation of transcription, the products of which ultimately confer cell identity and function. Most consensus peaks (54% in cattle, 58% in pig, and 63% in mouse) were only present in a single tissue (Fig. 2b), suggesting that these regions were involved in tissuespecific regulatory programs. These regions were of particular interest, considering that differentially accessible regions have been associated with higher density of transcription factor (TF) binding sites, hinting at interesting regulatory roles.

To stringently identify open chromatin that was specific to a given tissue, only consensus peaks that did not overlap any peaks called from either replicate in any other tissue were considered tissue-specific. In sum, 71,479 peaks in pig, 47,454 peaks in cattle, and 116,700 peaks in mouse (Fig. 3a) were identified as having highly tissue-specific ATAC-seq signal (Fig. 3b). Interestingly, tissues with the greatest number of tissue-specific peaks varied between species. Although cerebellum-specific peaks were numerous compared to the cortex in cattle and pig, the opposite trend was observed in the mouse. Of the remaining tissues, liver-specific peaks were particularly abundant in mouse, whereas lung-specific peaks were prevalent in cattle, and muscle-specific peaks were the most frequent in pig.

Whereas all consensus peaks were enriched at TSS (pvalue <1e-200) (Fig. 2a, Fig. S5a), tissue-specific peaks coincided with TSS less frequently (Fig. 3c). Although TSS annotation in these species is likely to be incomplete, the lack of tissue-specific peaks near annotated TSS suggests that tissue-specific open chromatin is more likely to delineate enhancers, which are known to demonstrate highly tissue-specific activity, both spatially and temporally. Tissue-specific peaks from cerebellum, cortex, liver, lung, muscle, and spleen were evaluated for motif enrichment in cattle, pig, and mouse. Adiposeand hypothalamus-specific peaks were excluded from this analysis, due to the low number of tissue-specific peaks detected, and lack of hypothalamus data in the mouse. Several TF families demonstrated consistent motif enrichment in particular tissues, such as forkhead box family members, which were enriched in liver- and lung-specific open chromatin in all three species (Fig. 3d). Homeobox motifs also demonstrated consistent enrichment patterns in tissue-specific open chromatin across species; HNF motifs were enriched in liver, NKX3.1 motifs in lung, and MEIS1 and SIX1 motifs in muscle. Brain-specific open chromatin was consistently enriched for motifs of brain-specific TFs (ATOH1, NEU-ROD1, and OLIG2). Nevertheless, several discrepancies

Species	Tissue	Rep. A Peaks	Rep. B Peaks	Rep. A Peaks overlapping Rep. B Peaks	% Rep. A Peaks	Rep B. Peaks overlapping Rep A. peaks	% Rep B. Peaks
Cattle	Adipose	59,612	133,768	38,647	64.8	38,471	28.8
	Brain Cortex	109,395	160,546	75,462	69.0	75,206	46.8
	Hypothalamus	59,966	37,036	19,970	33.3	19,999	54.0
	Liver	102,704	114,583	58,444	56.9	58,563	51.1
	Lung	221,576	248,844	167,311	75.5	166,777	67.0
	Muscle	107,208	113,502	76,780	71.6	76,801	67.7
	Spleen	110,852	200,323	79,355	71.6	79,007	39.4
Pig	Adipose	9192	7645	4778	52.0	4778	62.5
	Cerebellum	220,327	214,455	132,123	60.0	132,292	61.7
	Brain Cortex	142,658	156,069	103,581	72.6	103,382	66.2
	Hypothalamus	103,402	144,418	63,382	61.3	63,368	43.9
	Liver	112,202	136,085	78,395	69.9	78,274	57.5
	Lung	167,298	191,926	114,470	68.4	113,864	59.3
	Muscle	137,000	105,823	92,039	67.2	92,705	87.6
	Spleen	100,982	100,867	64,192	63.6	64,208	63.7

Table 3 Replicability of ATAC-seq peaks. Overlap of peak sets derived from biological replicates

between species were noted in motif enrichment of tissue-specific open chromatin. For instance, bHLH factors MYF5, MYOD, and MYOG motifs were consistently enriched in muscle-specific open chromatin, but only demonstrated enrichment in cortex-specific open chromatin in the mouse. Across species, spleenspecific open chromatin demonstrated little motif enrichment, with the notable exception of ETS, IRF, and RUNT TF family motifs, which were almost exclusively enriched in the pig. Overall, liver-, lung-, and muscle-specific regulatory circuitry appeared to be the most highly conserved across cattle, pig and mouse, whereas brain-specific regulation was more varied between species.

### Conservation of chromatin accessibility across mammals

Although extensive epigenetic divergence is expected between species, sequence similarity among cattle, pig, and mouse genomes well above the coding sequence fraction suggests that a significant portion of epigenetic control of transcription is likely under evolutionary constraint. Having observed similarities in motif enrichment in tissue-specific open chromatin between species, it was suspected that the sequence and accessibility of regulatory elements would also be constrained.

Reasonably, portions of regulatory elements, such as the motifs that facilitate TF-DNA contacts, would be under selective pressure, especially those that connect tissue-specific transcription factors to conserved tissue-

**Table 4** Regions identified as ATAC-seq peaks in both biological replicates of pig, cattle, and mouse tissues. To obtain a single comprehensive set of "consensus" ATAC-seq peaks for each species, regions that were identified as peaks in both biological replicates for each tissue were collapsed, such that any overlapping peaks were merged into a single unique interval

	Pig		Cattle		Mouse	
Tissue	Peaks in both replicates	Average size (bp)	Peaks in both replicates	Average size (bp)	Peaks in both replicates	Average size (bp)
Adipose	4785	373.583	38,745	501.575	64,008	633.770
Cerebellum	134,086	555.921	93,927	535.284	57,771	618.660
Brain Cortex	104,272	515.209	75,762	465.855	149,223	604.979
Hypothalamus	63,736	514.036	20,045	415.089	-	-
Liver	78,841	498.528	58,853	482.561	90,228	679.330
Lung	115,491	569.537	169,734	619.450	76,168	652.975
Muscle	93,550	698.708	77,378	566.262	22,003	514.762
Spleen	64,667	560.972	79,960	542.832	35,278	601.650
Consensus (collapsed set)	306,304	623.608	273,594	616.064	254,076	668.062



from tissue-specific (accessible in only one tissue) to ubiquitous (accessible in all sampled tissues). Consensus peaks that were accessible in a single tissue were further broken down by tissue. **c** Distribution of consensus peak activity for regions containing CTCF motifs

specific expression programs. To identify homologous regions that correspond to regulatory elements, the coordinates of consensus ATAC-seq peaks from each species were projected to the other two species using Ensembl Compara [27], which is largely based on wholegenome pairwise and multiple sequence alignments. Unsurprisingly, considering the smaller size of the mouse genome and the larger relative evolutionary distance between mice and ungulates, more consensus peaks could be mapped between pig and cattle, as opposed to between pig and mouse or cattle and mouse. Overall, about half of peaks were conserved at the sequence level between cattle and pig, whereas only about a third of peaks were conserved at the sequence level between ungulates and mice (Fig. 4a). Moreover, about 40% of accessible regions that could be mapped between pig to cattle were accessible in at least one tissue in both species, whereas only about 30% of accessible regions mapped between mouse and pig, or between mouse and cattle, demonstrated conserved accessibility in at least one tissue (Fig. 4a; Fig. S6). Overall, conservation of open chromatin at specific loci was in line with evolutionary distance. Comparing cattle and pig, which are separated by about 62 million years, about 20% of

Cattle consensus peaks			Pig consensus peaks			
Motif	P-value	Peaks with motif (%)	Motif	P-value	Peaks with motif (%)	
CTCF (Zf)	1e-2891	8.66	CTCF (Zf)	1e-2587	8.16	
BORIS (Zf)	1e-1695	11.56	BORIS (Zf)	1e-1580	10.90	
NF1(CTF)	1e-505	16.51	Jun-AP1 (bZIP)	1e-784	8.25	
Sp1 (Zf)	1e-389	8.60	Fosl2 (bZIP)	1e-769	11.25	
CEBP (bZIP)	1e-334	14.00	Fra1 (bZIP)	1e-718	16.30	
ETS (ETS)	1e-278	9.70	Sp1 (Zf)	1e-639	8.16	
NRF1 (NRF)	1e-276	3.09	BATF (bZIP)	1e-624	18.32	
RFX (HTH)	1e-272	2.75	Mef2d (MADS)	1e-583	6.00	
Rfx2 (HTH)	1e-264	3.07	Atf3 (bZIP)	1e-576	19.02	
Mef2d (MADS)	1e-257	4.82	Mef2c (MADS)	1e-502	11.63	

**Table 5** Motif enrichment in consensus open chromatin. Top ten enriched known binding motifs identified from the merged set of consensus peaks in each species

consensus peaks were conserved in terms of sequence and accessibility, whereas mouse, separated from ungulates by about 96 million years, only shared about 10% of consensus peaks, in terms of sequences and accessibility, with either species (Fig. S7a,b). Additionally, promoter accessibility at homologous regions was considerably more conserved than enhancer accessibility at homologous regions, with almost half of promoter open chromatin in the pig detected in cattle, while only a fifth of all open chromatin in pig was conserved in cattle (Fig. S7a,b).

Among the consensus peaks identified in cattle, pig and mouse, 145,801 regions could be mapped to all three species. Of these, 13,735 were consistently accessible in the same tissue (at least one) in all three species, and 30,215 were accessible in at least one tissue in both cattle and pig (Fig. 4b). Regions with conserved accessibility in all three species tended to be ubiquitously present in all sampled tissues. Whereas only 2-4% of consensus peaks were accessible in all tissues when considering a single species, 7% of regions with conserved accessibility in all species were accessible in all tissues (Fig. S7c). Furthermore, regions with conserved accessibility in all three species were heavily enriched around TSS (32% of regions), especially those which were accessible in all tissues (97% of regions), which marked TSS of housekeeping genes (Fig. 4c; Table S2). In contrast, regions that demonstrated conserved accessibility in cattle and pig, but not in mouse, were very rarely accessible in all tissues (only 23 out of 14, 543 regions, or 0.2%) (Fig. S7d), and only occurred at TSS 4% of the time (Fig. S7e).

Intriguingly, most regions with conserved accessibility in cattle, pig and mouse were only open in one or two tissues (62%) and were predominantly intronic and intergenic (Fig. S7e). For instance, several regions upstream of the *MEF2A* locus demonstrated musclespecific accessibility in all three species (Fig. 4d). These regions could represent conserved enhancers, suggesting that even distal regulatory elements are subject to some level of evolutionary constraint. In all, 3105 intergenic loci (6%) demonstrated conserved open chromatin signatures in all three species, and further examination of the genes that were closest to these sites (within 100 kb) revealed functional enrichment for developmental processes, such as regionalization and organogenesis (Table S3).

Notably, this small subset of "conserved" enhancers may underrepresent conserved activity at distal loci. For instance, accessible regions around the *FOXG1* locus appear to be syntenically conserved across all three species; however, only one region – corresponding to the TSS of a human long non-coding RNA – could be mapped to all three species (Fig. S8). The remaining loci could not be mapped between two species, suggesting a pervasive lack of overall sequence identity, despite apparent functional conservation.

### Discussion

Despite the intimate connection between chromatin structure and regulation of transcription, an atlas of chromatin accessibility in livestock tissues had not yet been reported. To address this gap in knowledge, ATAC-seq was used to identify regions of open chromatin in a prioritized set of pig and cattle tissues, yielding a first glimpse at the landscape of active regulatory elements in these genomes. In all, 6 to 7% of the cattle and pig genomes demonstrated accessibility in at least one tissue, which was consistent with a comparable dataset in the mouse [28]. Notably, about half of these accessible sites were intergenic, accounting for about 3% of each genome. The identification of these regulatory elements is a crucial first step towards a comprehensive annotation of the non-coding genome, which has been severely lacking in livestock species [29].

Although efforts are currently underway to further characterize these regulatory elements as enhancers, silencers, insulators, promoters, etc. [29, 30], motif enrichment analysis highlighted some of their potential regulatory



bp upstream and downstream. **c** Distribution of tissue-specific open chromatin relative to gene annotations. **d** Enrichment of known TF binding motifs in tissue-specific open chromatin. Motifs sorted by TF family. Sets of tissue-specific open chromatin for each species are grouped by tissue. Increasing size and color intensity indicate increasing enrichment for a given motif

roles. By far, the most enriched sequences in cattle and pig open chromatin were CTCF motifs, suggesting pervasive involvement of accessible regions in higher order chromatin organization. In particular, convergently oriented CTCF sites are known to demarcate topologically associated domain (TAD) boundaries [31], which are largely invariable across cell types [32–34] and even across species [32, 33, 35, 36]. Indeed, regions that were globally accessible in all pig and cattle tissues were particularly enriched for CTCF recognition motifs, suggesting that these regions may delineate TAD boundaries, although direct profiling of chromatin interactions will be necessary to provide more definitive annotations of 3D chromatin structure.

Interestingly, out of all accessible CTCF motifs, almost a third were only accessible in a single tissue, which is consistent with CTCF binding being highly variable in different cell types [37, 38], even though TAD structure is largely consistent [32–34]. Only a fraction of CTCF



three species and in ungulates laid over a phylogenetic tree reflective of evolutionary distance in millions of years (MYA). **c** Genomic distribution of regions with conserved accessibility in all three species, relative to the mouse gene annotation. Brief summary of enriched gene ontology (GO) terms for genes marked by conserved open chromatin at their TSS in every tissue. **d** Consensus peaks, consensus peaks with conserved sequence (that could be mapped to all three species), and consensus peaks with conserved accessibility in all three species at the *MEF2A* locus. Tracks show normalized ATAC-seq signal. Conserved promoter open chromatin highlighted in green. Conserved distal (putative enhancer) open chromatin highlighted in yellow

binding sites (15%) actually localize to TAD boundaries [39], and most CTCF binding sites are interspersed with enhancers, stabilizing enhancer-promoter interactions [40], and forming cell type-specific chromatin loops linked to differential gene expression [40–42]. Therefore,

differentially accessible CTCF motifs may reflect tissuespecific chromatin looping. Taken together, the presence of both tissue-specific and globally accessible CTCF motifs suggests a multi-tiered 3D structure that participates in both fundamental and tissue-specific regulation.

Tissue-specific open chromatin was widespread and conspicuously lacking near TSS. Motif enrichment analysis revealed that tissue-specific open chromatin demonstrated conserved enrichment for tissue-specific TF binding motifs, which was expected, as expression programs in vertebrate tissues are thought to be controlled by a highly conserved set of tissue-specific TFs [43]. However, not all TF families demonstrated consistent motif enrichment in tissue-specific open chromatin. The motifs of the RUNX family, highly conserved TFs involved in cell fate determination [44], were only enriched in mouse lung-, pig cerebellum-, and spleenspecific open chromatin. Whether this points to a divergence in tissue-specific regulation remains unclear, as motif enrichment analyses rely heavily on known binding motifs in human and mouse, failing to account for any species-specific differences in TF recognition sites.

Certainly, divergent chromatin structure has implications for differential transcriptional regulation, a phenomenon that has been long recognized as a significant contributor to phenotypic diversity [45-49]. As expected, the proportion of open chromatin that was conserved between species was consistent with evolutionary distance - higher concordance was observed between cattle and pig, than between mouse and either cattle or pig. By classifying loci as either proximal or distal based on their closeness to annotated genes, it was also apparent that accessibility at proximal elements, such as promoters, was significantly more conserved than accessibility at distal elements. This discrepancy in functional conservation was not altogether surprising; whereas promoters are fundamental for gene expression in any context, modulation of enhancer activity can subtly alter phenotypes without compromising viability [50, 51]. In fact, enhancers are known to evolve rapidly, and several studies have demonstrated how changes to enhancer sequences can lead to differing phenotypes between species [52-55]. Indeed, only 17% of intergenic open chromatin in cattle was also accessible in pig, and a meager 6% was accessible in mice, indicating that enhancers are largely species-specific, as has been previously demonstrated [3, 46]. Nevertheless, more than 3000 intergenic loci, relative to gene annotations in the mouse, had a conserved open chromatin signature in at least one tissue in all three species. Considering some highly conserved enhancers have been implicated in core biological processes, such as embryonic development [56], these intergenic loci are suspected to be involved in fundamental biological processes in adult tissues, which would account for their abnormal sequence constraint and functional conservation.

Intriguingly, several loci appeared to share open chromatin signatures based on synteny, despite lack of sequence conservation. Several studies have demonstrated that enhancer function can be conserved even when overall sequence is not [57–59]. Instead, selective pressure may only operate on the functional components of regulatory elements: TF binding sites, which are typically short and degenerate sequences [45, 46, 52]. Although inferring sequence conservation is possible with sequences as short as 36 bp [45], detecting homologous regulatory regions based only on conserved TF binding sites (6–12 bp [60]) is problematic. This begs a pragmatic question in the field of comparative epigenomics: if orthologous regions cannot be determined based on sequence, how then can we determine whether function is conserved? If TF binding sites are all that is required for enhancer function, then most enhancers would not be conserved in the canonical sense of sequence constraint, but instead through TF binding and relative positioning.

### Conclusions

To our knowledge, these data constitute the first atlas of chromatin accessibility in a common set of livestock tissues, and consequently a first look at the distribution across multiple tissues of active regulatory elements in the pig and cattle genomes. Moreover, this initial annotation of the non-coding genome will help to inform the identification of causal variants for disease and production traits. From the standpoint of comparative epigenomics, these data contribute to the ever-growing wealth of epigenomic information; the comprehensive analysis of which will undoubtedly help bridge the gap between genome and phenome, providing crucial insight into transcriptional regulation and its connection to evolution.

### Methods

### Tissue collection and cryopreservation

All necessary permissions were obtained for collection of tissues relevant to this study, following the Protocol for Animal Care and Use #18464, as per the University of California Davis Animal Care and Use Committee (IACUC). As described previously [61], two intact male Line 1 Herefords, provided by Fort Keogh Livestock and Range Research lab, were euthanized by captive bolt under USDA inspection at the University of California, Davis. Both cattle were 14 months of age and shared the same sire. Two castrated male Yorkshire pigs were humanely euthanized by animal electrocution followed by exsanguination, which is the standard method of euthanasia at pig slaughterhouses, under USDA inspection at Michigan State University Meat Lab. Pigs were littermates aged 6 months old, and sourced from the Michigan State University Swine Teaching and Research Center. From each animal, subcutaneous adipose, frontal cortex, cerebellum, hypothalamus, liver, lung (left lobe), longissimus dorsi muscle, and spleen were collected and promptly processed for cryopreservation. For each sample, roughly one gram of fresh tissue was minced and transferred to 10 mL of ice-cold sucrose buffer (250 mM

D-Sucrose, 10 mM Tris-HCl (pH 7.5), 1 mM MgCl<sub>2</sub>; 1 protease inhibitor tablet per 50 mL solution just prior to use). Minced tissue was twice homogenized using the gentleMACS dissociator "E.01c Tube" program. Homogenate was filtered with the 100  $\mu$ m Steriflip vacuum filter system, volume was brought up to 9.9 mL with sucrose buffer, and 1.1 mL DMSO was added to achieve a 10% final concentration. Preparations were aliquoted into cryovials and frozen at – 80 °C overnight in Nalgene Cryo 1 °C freezing containers, then stored at – 80 °C long-term.

### ATAC-seq library construction and sequencing

A modified ATAC-seq protocol compatible with cryopreserved tissue samples was employed [62]. Cryopreserved tissue samples were thawed on ice, then centrifuged for 5 min at 500 rcf and 4 °C in a centrifuge with a swinging bucket rotor. Pellets were resuspended in 1 mL ice-cold PBS, and centrifuged again for 5 min at 500 rcf and 4 °C. Pellets were then resuspended in 1 mL ice-cold freshly-made ATAC-seq cell lysis buffer (10 mM Tris-HCl pH = 7.4, 10 mM NaCl,  $3 \text{ mM MgCl}_2$ , 0.1% (v/v) IGEPAL CA-630), and centrifuged for 10 min at 500 rcf and 4 °C. Pellets were then resuspended again in ice-cold PBS for cell counting on a hemocytometer. Between 50,000 and 1,000,000 cells were aliquoted for library preparation, depending on tissue, success of previous library preparation attempts, and cell abundance in a given preparation (Table S1). Aliquoted cells were centrifuged once more for 5 min at 500 rcf and 4 °C, and pellets were resuspended in 50 µL transposition mix  $(22.5 \,\mu\text{L} \text{ nuclease-free H}_2\text{O}, 25 \,\mu\text{L} \text{ TD buffer and } 2.5 \,\mu\text{L}$ TDE1 enzyme from Nextera DNA Library Prep Kit (Illumina, cat. no. FC-121-1030)). Nuclear pellets were incubated with transposition mix for 60 min at 37 °C, shaking at 300 rpm. Transposed DNA was purified with the MinElute PCR Purification Kit (Qiagen, cat. no. 28004) and eluted in 10 µL Buffer EB. Eluted DNA was added to 40 µL PCR master mix (25.4 µL SsoFast<sup>™</sup> Eva-Green<sup>°</sup> Supermix,  $13 \,\mu\text{L}$  nuclease-free H<sub>2</sub>O,  $0.8 \,\mu\text{L}$ 25 µM Primer 1, 0.8 25 µM Primer 2 (see Table S4 for sequences)) to  $10 \,\mu\text{L}$  eluted DNA and PCR cycled (1 x [5 min at 72 °C, 30 s at 98 °C], 10-13x [10 s at 98 °C, 30 s at 63 °C, 1 min at 72 °C]). Libraries were then purified again with the MinElute PCR Purification Kit, and eluted in 10 µL Buffer EB. Libraries were quantified by Qubit (Thermo Fisher Scientific, Inc., Waltham, MA), and checked for nucleosomal laddering using a Bioanalyzer High Sensitivity DNA Chip (Agilent Technologies, Santa Clara, CA) (Figs. S9 and S10). Details for individual library preparations, including cell input, PCR cycles, and concentrations can be found in Table S1. Finally, libraries were size selected for subnucleosomal length fragments (150–250 bp) on the PippinHT system using a 3% agarose cassette (Sage Science, Beverly, MA). Size selection and DNA concentration were evaluated with a Bioanalyzer High Sensitivity DNA Chip, pooled, and submitted for sequencing on the NextSeq 500 platform to generate 40 bp paired end reads.

### ATAC-seq data preprocessing and quality evaluation

Low quality bases and residual adapter sequences were trimmed from raw sequencing data using Trim Galore! (v0.4.0), a wrapper around Cutadapt (v1.12) [63], with the options "-a CTGTCTCTTATA" and "-length 10" to retain trimmed reads at least 10 bp in length. Trimmed reads were then aligned to either the susScrofa11 (pig), ARS-UCD1.2 (cattle), or GRCm38 (mouse) genome assemblies using BWA mem with default settings (v0.7.17) [64]. Duplicate alignments were removed with Picard-Tools (v2 .9.1), and mitochondrial and low quality (q < 15) alignments were removed using SAMtools (v1.9) [65]. Finally, broad peaks were called using MACS2 (v2.1.1) [66] with options "-q 0.05 -B –broad –nomodel –shift -100 –extsize 200."

For all reported statistics, standard error is also reported. Quality metrics were calculated as follows. The non-redundant read fraction (NRF) was calculated by dividing the number of nonduplicate uniquely mapping reads out of all mapped reads. The Fraction of Reads in Peaks (FRiP) score for each sample was calculated using the plotEnrichment function from the deepTools suite (v.3.2.0), given the broad peaks identified by MACS2. Finally, the synthetic Jensen-Shannon distance (sJSD) was calculated using the deepTools plotFingerprint function.

For visualization, genome-wide ATAC-seq signal was normalized by RPKM in 50 bp windows using the bam-Coverage function from the deepTools suite. Other deepTools functions were used along with the resulting bigwig files to generate [1] pairwise scatter plots of genome-wide signal, including Spearman correlation coefficients (plotCorrelation -log1p -removeOutliers), [2] principal components analyses of signal in consensus peaks (plotPCA -transpose -log2), and signal at loci of interest (plotHeatmap), such as TSS (computeMatrix reference-point -beforeRegionStartLength 2000 -after-RegionStartLength 2000 -skipZeros) and peaks (computeMatrix scale-regions -beforeRegionStartLength 500 regionBodyLength 500 -afterRegionStartLength 500 skipZeros). Normalized signal from bigWig files was visualized at specific loci with the UCSC Genome Browser, limiting the y-axis range to 20 and displaying the mean in smoothing windows of 4 bases.

# Identification of consensus and tissue-specific open chromatin

To obtain a single comprehensive set of consensus ATAC-seq peaks for each species, regions that were identified as peaks in both biological replicates of a tissue were first identified, and then collapsed. Specifically, for each tissue peaks from biological replicates were compared with BEDtools intersect (v2.26.0) [67] to identify intersecting regions that were consistently identified as peaks. In the case of cattle cerebellum, for which only one biological replicate was available, robust peaks were identified by calling broad peaks more stringently with MACS2 (-q 0.01 -B -broad nomodel -shift - 100 -extsize 200). Regions that were called as ATAC-seq peaks in both biological replicates (or which were especially robust, in the case of cattle cerebellum) were then collapsed with BEDtools merge [67] (v2.26.0) to obtain a single comprehensive set of "consensus" peaks that accounted for accessibility in any of the eight tissues. Tissue-specific peaks were defined as consensus peaks that were [1] identified as peaks in both biological replicates of the tissue in question, and [2] not detected as accessible in either biological replicate of any other tissue. These comparisons were conducted using BEDtools intersect to compare consensus peaks with broad peaks called from individual biological replicates.

# Categorization of peaks by location relative to gene annotations

Peaks were categorized by position relative to features in the Ensembl annotations (v96) for each species using BEDtools intersect with default settings. Because many peaks overlap multiple features, peaks were first classified as TSS (within 50 bp), then as promoters (within 2 kb upstream of TSS), as transcription termination sites (TTS; within 50 bp), as overlapping a 5' untranslated region (UTR), as overlapping a 3' UTR, as exonic, as intronic, and finally, if peaks did not overlap any of these features, they were considered to be intergenic.

To determine if peaks were enriched near genomic features, peak locations were iteratively randomized 100 times using BEDtools shuffle, excluding "unmappable" genomic regions to avoid bias. Unmappable regions were empirically defined as any 500 bp window to which no read mapped in any library. Randomized peaks were also categorized by position relative to gene annotations, and compared to the localization of the actual peak set using a one-sample T-test (one-tailed).

#### Motif enrichment analysis

Regions of interest were evaluated for motif enrichment using the HOMER findMotifs.pl function (v4.8) [68], and the top ten enriched known motifs, based on pvalues, were reported.

### Conservation of open chromatin

All interspecies comparisons were based on the 46mammalian Enredo-Pecan-Ortheus (EPO) multiple sequence alignment (MSA) available through Ensembl Compara (v99) [27]. Regions of consensus open chromatin in each species were projected onto the other two species using the Ensembl Compara Application Programming Interface (API). For simplicity, regions that mapped to multiple loci in another species were discarded prior to evaluating whether accessibility was conserved. Chromatin accessibility was considered to be conserved at homologous regions if they overlapped (by at least 1 bp) consensus open chromatin in the same tissue (at least one tissue) in all species in question.

### Functional annotation enrichment analysis

Ensembl IDs were converted to external gene names using the BiomaRt package, and these were submitted to DAVID (v6.8) [69, 70] for functional annotation clustering. *Mus musculus* was used as background, and functional annotation clustering was conducted on medium stringency for the following terms: GOTERM\_BP\_5, GOTERM\_CC\_DIRECT, GOTERM\_MF\_DIRECT, BIO-CARTA, and KEGG\_PATHWAY. For each gene set, the top four clusters were reported.

### Supplementary information

Supplementary information accompanies this paper at https://doi.org/10. 1186/s12864-020-07078-9.

Additional file 1 Figure S1. Correlation of ATAC-seg signal in select technical replicates ATAC-seqlibraries. A) Pearson correlation of genomewide signal (RPKM) in 500 bp windows. B) PCA of Cortex A technical replicate libraries alongside all biological replicates. C) PCA of pig technical replicate libraries alongside all biological replicates. D) Signal of cattle cortex technical and biological replicates at the STMN4 locus. E) Signal of pig cerebellum and hypothalamus technical and biological replicates at the STMN4 locus. Figure S2. Correlation of ATAC-seq signal in biological replicates. Scatterplots showing Pearson correlation of normalized genome-wide signal in 500 bp windows between biological replicates for cattle and pig tissues. Figure S3. ATAC-seq signal at TSS. Heatmaps depicting normalized ATAC-seq signal in the proximity of TSS, including 2 kb upstream and downstream, with TSS sorted by signal intensity. Figure S4. PCA of normalized ATAC-seq signal in consensus open chromatin identified in pig and cattle tissues tissues. Principal components 1, 2 and 3 are included to better visualize clustering of tissues. Figure S5. Mouse open chromatin localization and differential accessibility. A) Distribution of mouse consensus open chromatin relative to the Ensembl gene annotation (v96). B) Distribution of consensus peak activity, ranging from tissue-specific (accessible in only one tissue) to ubiquitous (accessible in all sampled tissues). Consensus peaks that were accessible in a single tissue were further broken down by tissue. Figure S6. Conservation of open chromatin in individual tissues. Titles above bar plots indicate the species that consensus peaks were identified in, followed by the species to which the consensus peak coordinates were projected to evaluate accessibility conservation in the corresponding tissue. Figure S7. Characterization of conserved open chromatin. Proportion of all consensus peaks, promoter consensus peaks (within 2 kb upstream and 50 bp downstream of TSS), and intergenic consensus peaks that were identified in (A) cattle or (B) pig that demonstrated both conserved sequence and accessibility in the other two species. Number of tissues in which consensus peaks demonstrated conserved accessibility in (C) all three species or (D) only in cattle and pig. E) Distribution of consensus peaks with conserved accessibility in cattle, pig, and mouse, relative to the mouse gene annotation (Ensembl v96). Figure S8. Positional conservation of chromatin accessibility at the FOXG1 locus. For each species, consensus peaks, consensus peaks with conserved sequence (that could be mapped to all three species), and consensus peaks with conserved accessibility are

shown. Tracks show normalized ATAC-seq signal for each sample. Conserved promoter open chromatin is highlighted in green. Consensus peaks that appear to be syntenically conserved, relative to FOXG1, but which could not be mapped between species, are highlighted in grey. Figure S9. Bioanalyzer traces of cattle ATAC-seg libraries prior to size selection. Bioanalyzer traces were used to check for nucleosomal laddering. Size selection removed excess primer and fragments > 250 bp. Figure **\$10.** Bioanalyzer traces of pig ATAC-seq libraries prior to size selection. Bioanalyzer traces were used to check for nucleosomal laddering. Size selection removed excess primer and fragments > 250 bp. Table S1. ATACseq library construction details. For each library constructed, rounds of PCR amplification, number of cells used as input, and concentration in the 150–250 bp range prior to size selection are indicated. Table S2. Functional annotation clustering of genes with conserved and global TSS accessibility. Genes with accessible TSS (± 50 bp) in all profiled tissues in all species were subjected to functional annotation clustering to identify enriched cellular functions. Top four clusters reported. Table S3. Functional annotation clustering of genes near conserved intergenic open chromatin. Genes that were closest (within 100 kb) to intergenic open chromatin that was conserved in all three species were subjected to functional annotation clustering to identify enriched cellular functions. Top four clusters reported. Table S4. ATAC-seg oligos used for PCR. Sequences have been previously described by Buenrostro et al., 2013. Primers 2A-2X contain variable barcodes which permit library pooling prior to sequencing, and which were used to demultiplex sequencing data.

**Additional file 2.** Cattle ATAC-seq peaks. Genomic locations of ATAC-seq peaks that were called for each tissue in each replicate.

**Additional file 3.** Pig ATAC-seq peaks. Genomic locations of ATAC-seq peaks that were called for each tissue in each replicate.

### Abbreviations

ATAC-seq: Assay for Transposase Accessible Chromatin using sequencing; CDS: Coding sequence; ChIP-seq: Chromatin immunoprecipitation followed by sequencing; DNase-seq: DNase I hypersensitive sites sequencing; FRIP: Fraction of reads in peaks; GO: Gene ontology; MYA: Million years ago; NRF: Non-redundant read fraction; PCA : Principal components analysis; RPM: Reads per million; sJSD: Synthetic Jensen-Shannon distance; TAD: Topologically associated domain; TF: Transcription factor; TSS: Transcription start site; TTS: Transcription termination site; UTR: Untranslated region

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Not applicable.

### Authors' contributions

MMH, HZ and PJR designed the study. JM, AVE, IK, CKT, CWE contributed to the experimental design. MMH, PS, YW, and GC collected samples and constructed ATAC-seq libraries. MMH conducted bioinformatics analyses with guidance from CK. MMH, HZ and PJR wrote the manuscript. All authors have read and approved the manuscript.

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### Availability of data and materials

ATAC-seq data generated by this study is available from the European Nucleotide Archive under project ID PRJEB14330 (https://www.ebi.ac.uk/ena/ data/view/PRJEB14330). Raw ATAC-seq data for male mesenteric fat, cerebrum, cerebellum, liver, lung, skeletal muscle and spleen from mice [28] were downloaded from the CNGB Nucleotide Sequence Read Archive, under Project ID CNP0000198. Genome assemblies and annotations (v96) can be obtained from ENSEMBL (ftp://ftp.ensembl.org/pub/release-96/).

### Ethics approval and consent to participate

This study was conducted following ethics guidelines of the University of California, Davis. All necessary permissions were obtained for collection of tissues relevant to this study following the Protocol for Animal Care and Use #18464, as per the University of California Davis Animal Care and Use Committee (IACUC).

### **Consent for publication**

Not applicable.

### **Competing interests**

The authors declare that they have no competing interests.

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