Comprehensive analyses of 723 transcriptomes enhance genetic and biological interpretations for complex traits in cattle

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By uniformly analyzing 723 RNA-seq data from 91 tissues and cell types, we built a comprehensive gene atlas and studied tissue specificity of genes in cattle. We demonstrated that tissue-specific genes significantly reflected the tissue-relevant biology, showing distinct promoter methylation and evolution patterns (e.g., brain-specific genes evolve slowest, whereas testis-specific genes evolve fastest). Through integrative analyses of those tissue-specific genes with large-scale genome-wide association studies, we detected relevant tissues/cell types and candidate genes for 45 economically important traits in cattle, including blood/immune system (e.g., CCDC88C) for male fertility, brain (e.g., TRIM46 and RAB6A) for milk production, and multiple growth-related tissues (e.g., FGF6 and CCND2) for body conformation. We validated these findings by using epigenomic data across major somatic tissues and sperm. Collectively, our findings provided novel insights into the genetic and biological mechanisms underlying complex traits in cattle, and our transcriptome atlas can serve as a primary source for biological interpretation, functional validation, studies of adaptive evolution, and genomic improvement in livestock.

Over the last decade, genome-wide association studies (GWAS) have been successful at discovering trait-/disease-associated genomic variants (Visscher et al. 2012, 2017). However, such studies provided limited information about novel molecular mechanisms underlying complex traits and diseases, partly due to the lack of knowledge of in what tissues or cell types those genomic variants would act. Recently, researchers have been actively pursuing a comprehensive map of functional elements, aiming to identify which genes and regulatory factors (e.g., promoters and enhancers) are functional or active in a large range of tissues and cell types—for example, Roadmap Epigenomics (Roadmap Epigenomics Consortium et al. 2015), GETx (The GTEx Consortium 2017), and Cell Atlas (Regev et al. 2017) projects in human, as well as the Functional Annotation of Animal Genomes (FAANG) project in livestock (Andersson et al. 2015). Integrative analyses of functional genome information with large-scale GWAS data provide unprecedented potential to discover trait-/disease-relevant tissues or cell types, which is crucial for understanding the molecular underpinnings of complex traits and diseases (Finucane et al. 2018; Hormozdiari et al. 2018). For instance, the Roadmap Epigenomics Consortium (2015) showed that GWAS hits of many traits and diseases are significantly enriched in epigenomic marks (e.g., H3K4me1) of trait-/disease-relevant tissues and cell types in humans. Finucane et al. (2018) recently explored disease-relevant tissues and cell types for various human diseases by examining their heritability enrichments among diverse tissues and cell types, such as inhibitory neurons for bipolar disorder (Finucane et al. 2018).

In livestock, due to the limited amount of functional genome data available (Fang et al. 2019), to our knowledge no previous publication has systematically reported the causal tissues or cell types for complex traits and diseases of economic importance. A comprehensive map linking complex traits with their specifically
relevant tissues will offer valuable information for fine-mapping causal genes/variants, for functionally validating of GWAS hits (i.e., selecting the “right” tissues and cell types), and for understanding of adaptive evolution (Quiver and Lachance 2018), as well as for the design of genome editing experiments (Ruan et al. 2017). Additionally, a better understanding of the genetic architecture underlying complex traits may make a contribution to the genetic improvement programs among livestock species (Goddard and Hayes 2009; Georges et al. 2019). For instance, Fang et al. (2017a,b) reported improved genomic prediction accuracy for mastitis and milk production traits in cattle by incorporating biological priors and gene expression information relevant to bacterial infection into genomic prediction models (Fang et al. 2017a,b).

Here, we uniformly assembled and analyzed 723 (156 newly generated and 567 existing) RNA-seq data sets to build a new gene atlas in cattle (Supplemental Code), which included 91 tissues and cell types from 447 individuals (http://cattlegeneatlas.roslin.ed.ac.uk). We summarized the global design of this study in Supplemental Figure S1. We first detected genes that were highly and specifically expressed in each tissue or cell type and then explored their biological characteristics in terms of biological function, DNA methylation, and evolution. We detected relevant tissues/cell types and candidate genes for 45 complex traits of economic importance in cattle, including 18 body conformation, six milk production, 12 reproduction, eight health, and one feed efficiency traits, by integrating those tissue-specific genes with large-scale (n = 27,214 bulls) GWAS data. We validated our findings by analyzing whole-genome DNA methylation data across major somatic tissues and sperm in cattle. In addition, we tested whether the tissue-specificity information of genes can improve genomic prediction. Our results, for the first time, systematically establish connections at the RNA level between tissue/cell types and complex traits in livestock and provide an important starting point for post-GWAS functional experiments to explore genotype-phenotype relationships in livestock.

Results

Summary of cattle gene atlas

Using a uniform pipeline of bioinformatics analysis, we obtained 18,468,126,120 clean reads from 723 RNA-seq data sets with an averaged uniquely mapping rate of 94.18%. We summarized details of sample information in Supplemental Table S1. We determined the normalized expression levels (i.e., fragments per kilobase per million mapped reads, FPKM) for all 24,616 Ensembl genes among 723 samples. In general, we found an average of 15,864 genes (median = 16,086, ranging from 7807 to 18,258) expressed (FPKM > 0) across 91 tissues and cell types, of which the majority (n = 14,682 on average) were protein-coding genes (Supplemental Fig. S2). Despite differences in experimental conditions and sample characteristics, samples from similar tissues and cell types clustered together based on their gene expression profiles (Fig. 1A), validating the potential of our data for studying the specificity of tissue expression. For instance, we found that samples from 14 adult brain regions (central neural system, CNS) clustered together with those from fetus brain and four other brain endocrine tissues (stark median eminence [SME], anterior pituitary, posterior pituitary, and pineal gland). All samples from seven blood/immune tissues and cell types clustered together, including CD4 cells, CD8 cells, white-blood cells, lymphocyte, spleen, thymus, and lymph nodes (Fig. 1A).

Detection and functional characterization of tissue-/cell type–specific genes

We calculated a t-statistic to measure the specific expression of a gene in a given tissue/cell type (Methods). We found that tissues and cell types within the same system highly positively correlated based on these t-statistics (Supplemental Fig. S3), indicating the high similarity of their tissue-specific expression. Of special interest, we found that mammary gland highly negatively correlated with corpus luteum and endometrium (Pearson’s r = −0.88 and −0.85, respectively) (Supplemental Fig. S3). This may reflect the well-known, antagonistic relationship between milk yield and fertility in dairy animals (Veerkamp et al. 2001; Berry et al. 2003). Additionally, liver and rumen epithelial cells negatively correlated with several immune tissues and cell types, including CD4 cells, CD8 cells, white blood cells, and thymus (the averaged Pearson’s r = −0.62) (Supplemental Fig. S3). This may support the observed connections between feed efficiency and immune responses in cattle (Hou et al. 2012). However, the underlying molecular mechanisms of these negative correlations are largely unknown and require further investigations.

We detected tissue-specific genes for each tested tissue based on the rank of t-statistics (i.e., top 5%). We showed the top tissue-specific genes in brain (GRM5), liver (SLC22A9), white blood cell (FCRL3), uterus (TDFG1), and testis (TRIM69) as examples in Figure 1B. The functional annotation of tissue-specific genes validated the known tissue-relevant biology (Fig. 1C; Supplemental Table S2). For instance, brain-specific genes significantly enriched for nervous system development (FDR = 1.67 × 10−48, enrichment fold = 3.24), liver for organic acid metabolism process (FDR = 1.33 × 10−51, enrichment fold = 4.92), white blood cell for regulation of immune system (FDR = 7.81 × 10−48, enrichment fold = 3.85), uterus for embryonic morphogenesis (FDR = 1.97 × 10−202, enrichment fold = 3.63), and testis for male gamete generation (FDR = 2.94 × 10−28, enrichment fold = 5.08) (Fig. 1C). In addition, we confirmed that promoters of tissue-specific genes in liver and muscle had specifically low DNA methylation in the corresponding tissues (Supplemental Fig. S4), consistent with promoter methylation being negatively correlated with gene expression (Smith and Meissner 2013). For instance, the promoter methylation of SLC22A9 (liver-specific expression), which is an important hepatic transport protein (Riedmaier et al. 2016), was significantly lower in liver when compared to other tissues (Supplemental Fig. S5). Moreover, our motif enrichment analysis of tissue-specific genes revealed potential master regulators (transcriptional factors) (Supplemental Fig. S6; Supplemental Table S3), which could contribute to regulation of gene activity and differentiations of cell types and tissues (Spitz and Furlong 2012). As shown in Supplemental Figure S6, we found that STAT1 was significantly (FDR<0.05) enriched in CD4 cells and lymph nodes, which have crucial roles in multiple immune responses (Shuai and Ivashkiv 2009), whereas ZFX, which participates in neuronal differentiation, was significantly enriched in hippocampus and cerebral cortex (Harel et al. 2012; Burney et al. 2013).

To explore the evolutionary conservation of tissue-specific genes among mammals, we first compared the cattle tissue-specific genes with human tissue-specific genes among 10 major tissues. We found that tissue-specific genes significantly overlapped in the matched tissues between cattle and human (Fig. 1D). We further explored dN/dS ratios of orthologous genes between cattle and five other mammals (i.e., human, mouse, dog, pig, and sheep). We consistently observed that genes specific for brain regions had
the significantly lowest $d_{SNP}/d_{DD}$ ratios, whereas genes specific for male reproductive tissues (e.g., testes and sperm) and blood/immune system (e.g., lymph nodes) had the significantly highest $d_{SNP}/d_{DD}$ ratios (Fig. 2; Supplemental Fig. S7). We then correlated the expression of all orthologous genes among major tissues in both cattle versus human and cattle versus sheep comparisons and confirmed that testes had the lowest correlation, whereas brain showed a relatively higher one (Supplemental Fig. S8). Our findings demonstrated that, in constrained tissues (e.g., brain), tissue-specific genes tended to evolve slowly, whereas in the relaxed tissues (e.g., testes), tissue-specific genes evolved more rapidly, revealing the importance of tissue-driven evolution.

Detection of tissues and cell types relevant with 45 agronomic traits

By integrating tissue-specific genes with large-scale GWAS, we revealed a comprehensive genetic relationship between 91 tissues/cell types and 45 complex traits of economic importance in cattle, providing novel insights into the molecular underpinnings of such economically important traits (Fig. 3). To validate our findings, we repeated GWAS signal enrichment analyses using tissue-specific DNA methylation regions instead of tissue-specific genes. We found GWAS enrichments from DNA methylation highly correlated with those from gene expression across all 45 traits among multiple tissues, for example, sperm (Pearson’s $r = 0.67$; $P = 4.41 \times 10^{-7}$) and lung (Pearson’s $r = 0.65$; $P = 1.60 \times 10^{-6}$) (Fig. 4). We summarized details of all 288 significant (FDR < 0.1) associations between traits and tissues in Supplemental Table S4. In addition, we summarized the top three expressed tissues of all 525 fine-mapped genes across all complex traits in Supplemental Table S5. The details of fine-mapped genes were described previously (Jiang et al. 2019).

Milk production traits

Generally, we observed that milk production traits were significantly associated with a few tissues and cell types (Fig. 3), indirectly supporting their highly polygenic architecture (Cole et al. 2009; Kemper and Goddard 2012). Of note, we found that mammary gland was the most significant ($P = 2 \times 10^{-4}$) tissue for protein yield (Supplemental Fig. S9A) and validated this by demonstrating that two of its fine-mapped genes (i.e., CSN1S1 with the posterior probability of causality (PPC) = 1, and PAEP with PPC = 0.84) were highly specifically expressed in mammary gland and milk cells (Supplemental Fig. S9B). Mammary gland was also the top significant tissue ($P = 1.5 \times 10^{-3}$) for lifetime net merit, which is an economic index including multiple traits that is used to rank animals for selection, suggesting the importance of a “good”
mammary gland in the dairy industry. Moreover, we found two fine-mapped genes, MRTFA (previously known as MKL1) (PPC = 1) and NCF4 (PPC = 0.55), for milk/protein yields and protein percentage, respectively, which specifically expressed in blood/immune system (Supplemental Fig. S10). This provides evidence of the underlying genetic correlations between milk production and immune disorders (e.g., mastitis) in cattle. Although brain tissues showed no significant enrichments for milk production (Fig. 3), we noticed that they indeed exhibited a significantly higher enrichment for milk production as compared to other types of traits, except for feed efficiency (i.e., residual feed intake [RFI]) (Fig. 5A). We found two fine-mapped genes, TRIM46 (PPC = 0.59) and RAB6A (PPC = 0.79), for protein percentage and milk yield, respectively, which highly specifically expressed in brain regions (Fig. 5B). By examining quantitative trait loci (QTL) of 19 milk-relevant traits in cattle QTLdb, we confirmed that brain-specific genes were significantly enriched for genes (i.e., closest genes to the lead SNPs) associated with milk production traits (e.g., milk yield and fat/protein percentage) but not for certain milk content traits (e.g., such as milk iron and zinc contents) (Supplemental Fig. S11). To further explore which brain regions were relevant to milk production traits, we pinpointed tissue-specific genes within 11 brain regions and another four brain endocrine tissues. We observed that anterior or pituitary, cerebellum, and temporal cortex were significantly (FDR < 0.05) associated with protein yield (Fig. 5C). To our knowledge, no previous publication has reported such relationships between the brain and milk production by an integrative analysis of genomic and transcriptomic data.

Body conformation traits

Body conformation (type) traits were significantly associated with many tissues and cell types, except for brain regions (Fig. 3), similar to findings in human height (Finucane et al. 2018), reflecting their highly polygenetic architectures. We used cattle stature as an example and found that three of its fine-mapped genes with PPC = 1, FG66, CCND2, and TCP11, were highly specifically expressed in fetal muscle, rumen epithelial cell, and testes, respectively (Supplemental Fig. S9C,D). By examining heritability enrichments of human height among 33 tissues (Finucane et al. 2018), we observed that these tissues significantly positively correlated (Pearson’s r = 0.64; P = 6.82 × 10⁻⁵) between human height and cattle stature in terms of GWAS signal enrichment (i.e., −log₁₀P) (Supplemental Fig. S12). Uterus and aorta were the top significantly enriched tissues for both human height and cattle stature, and they significantly associated with many other body type traits in cattle as well, such as rump width and rump angle (Fig. 3). All these findings support the view that mammals shared similar molecular mechanisms underlying body size.

Reproduction and health traits

We noticed that immune/blood system was significantly associated with multiple reproduction traits (Fig. 3). Overall, reproduction traits showed a significantly higher enrichment in immune/blood system when compared to other types of traits (Fig. 6A). The lymph node was the most significant tissue for both sire conception rate (P < 10⁻⁵) and sire still birth (P < 10⁻⁵) (Fig. 6B; Supplemental Table S4). We found a fine-mapped gene (i.e., CCDC88C with PPC = 1) of DFB (days to first breeding, a measurement of fertility ability), which highly specifically expressed in both blood/immune system and infundibulum (Fig. 6C). CCDC88C plays important roles in the regulation of T cells maturation during bacterial inflammation (Kennedy et al. 2014). Additionally, we found that immune/blood system was significantly associated with several health traits (Fig. 3). For instance, thymus was the top relevant (P = 2.70 × 10⁻³) tissue for ketosis (KETO) (Fig. 6B). We also found C6, a fine-mapped gene (PPC = 1) for somatic cell score (SCS) in milk, highly specifically expressed in liver and duodenum (Fig. 6C). SCS is an important indicator of mastitis in dairy cattle (Heringstad et al. 2006). Because the small intestine system exhibited immune functions (Santaolalla and Abreu 2012) and showed significant associations with multiple reproduction traits and health traits (Fig. 3), we further pinpointed tissue-specific genes within blood/immune system and four intestine parts, including ileum, duodenum, jejenum, and caecum. We found that thymus showed the highest and most significant enrichments for multiple health and reproduction traits, including daughter still birth,
daughter calving ease, SCS, KETO, DFB, and displaced abomasum (DSAB). CD8 cells were significantly associated with daughter calving ease and daughter still birth, whereas CD4 cells were significantly associated with cow conception rate and SCS (Fig. 6D).

Feed efficiency

We observed that the alimentary canal was the top relevant system for feed efficiency (i.e., RFI), among which rumen was the most significant ($P = 5.70 \times 10^{-3}$) tissue (Supplemental Fig. S13). We also found that brain pons was associated ($P = 2.19 \times 10^{-2}$) with RFI, which suggests the important role of the gut-brain axis in feed intake (Konturek et al. 2004). Nasal mucosa was another tissue associated ($P = 1.06 \times 10^{-2}$) with RFI, in line with the fact that olfactory receptors are known to be associated with RFI in cattle (Seabury et al. 2017).

A further application of this gene atlas is to explore whether the tissue specificity of genes could enhance genomic improvement in dairy cattle. We focused on three milk production traits (milk, fat, and protein yields). To reduce the redundancy and computational burdens, we clustered 91 tissues and cell types into 20 categories (Supplemental Fig. S14A). For each category, we then fitted SNPs within tissue-specific genes of this category and those in the remaining genome into a two-component Bayesian prediction model. By comparing with a two-component model (i.e., all genes vs. the remaining genome), whose prediction accuracy was similar to that of a single-component model (including all SNPs), we found that there is no improvement in genomic prediction accuracy on average (Supplemental Fig. S14B). We showed that the number of SNPs within tissue-specific genes did not bias prediction accuracy of models (Supplemental Fig. S15). However, we found the category consisting of immune/blood system and liver resulted in an increase of 0.041, 0.032, and 0.015 in prediction accuracy for fat yield, milk yield, and protein yield, respectively (Supplemental Fig. S14B). Another category consisting of salivary gland, larynx cartilage, tongue, chorid plexus, and muscle also increased the prediction accuracy across the three milk traits, that is, 0.044, 0.028, and 0.003 for fat, milk, and protein yields, respectively. We also

Figure 3. The relationships between 45 complex traits and 91 tissues and cell types. The color corresponds to enrichment degrees (i.e., $-\log_{10} p$) that are computed using a sum-based GWAS signal enrichment analysis based on the top 5% tissue-specific genes and a 50-kb extension. (* Corrected-$p$ (FDR) < 0.1.)
observed that brain regions led to an increase of 0.033 and 0.016 in prediction accuracy for fat yield and milk yield, respectively, but not for protein yield (Supplemental Fig. S14B). Of note was that DGAT1, a well-known milk and fat gene of large effect (Grisart et al. 2002), was not in those categories, implying that multiple loci of small effects are enriched in these tissue-specific genes.

Discussion

In this study, we built a cattle gene atlas by analyzing 723 RNA-seq data across seven tissues. Each dot represents a trait. The y-axis is for GWAS signal enrichments (−log₁₀P) obtained using tissue-specific DNA methylated regions, whereas the x-axis is for GWAS signal enrichments obtained using tissue-specific expressed genes. The r is for Pearson’s correlation.

more genes (n = 22,243) than before (n = 16,517). To increase the statistical power for detecting tissue-specific genes, we generated another 51 new RNA-seq data from 14 major somatic tissues and sperm in Holstein, as well as uniformly analyzed other 567 public RNA-seq data of high quality. Using this newly built gene atlas, we identified relevant tissues/cell types and candidate genes for 45 complex traits of economic importance and further applied it in genomic prediction. Of interest, we observed that brain was associated with milk production traits, and two brain-specific genes, TRIM46 and RAB6A, were fine-mapped genes for protein percentage and milk yield, respectively. TRIM46 plays key roles in neuronal polarity and axon specification (van Beuningen et al. 2015), whereas RAB6A is a key regulator of membrane traffic from the Golgi apparatus toward the endoplasmic reticulum (Matsuto et al. 2015). PheWAS based on both Gene atlas (http://geneatlas.roslin.ed.ac.uk/region-phewas/) (Canela-Xandri et al. 2018) and GWAS atlas (https://atlas.ctglab.nl/PheWAS) (Watanabe et al. 2019) showed that TRIM46 was significantly associated with many metabolic traits (e.g., blood urea nitrogen and impedance), whereas RAB6A was significantly associated with both neurological and metabolic traits (e.g., cingulum axial dissipivities and whole body fat-free mass). Our cattle gene atlas will serve as a valuable source for the livestock science community to interpret GWAS findings, to design follow-up validation experiments through choosing the “right” tissues and cell types, as well as to enhance genomic improvement in livestock. With more molecular phenotypes becoming available across diverse tissues in livestock in the near future, for instance, from the on-going FAANG project (Andersson et al. 2015), our current research strategy will help gain more novel insights into the genetic and biological mechanisms underpinning agronomic traits and thus enhance genomic improvement programs.

We noticed some limitations in our current study. Our basic assumption here was that genomic variants ultimately regulated complex traits by altering gene expression in the relevant tissues and cell types. Previous studies showed that the majority of expression quantitative trait loci (eQTL) were cis-variants (The GTEx Consortium 2017). We therefore focused on cis-regulators of tissue-specific genes by extending certain distances (i.e., 10 kb, 20 kb, and 50 kb) around such genes. In order to study trans-eQTLs, we need a large amount of samples for each tissue and cell type due to their relatively small effects (Grundberg et al. 2012). The cell type composition of tissues could confound our interpretation of results. As we showed in Figure 6D, CD4 cells and CD8 cells had distinct enrichments across 19 reproduction and health traits. Therefore, pure bulk cells and/or single-cell expression data may help further detect which cell types are causal in a trait-relevant tissue. Additionally, tissues sharing similar expression patterns with causal tissues could hinder us from detecting the “drivers” among multiple “passengers,” which is similar to the situation with GWAS results, wherein we can only interpret the significant tissues and cell types as the “best proxy” for the causative one. We are also limited by the availability of transcriptomic data, thus potentially ignoring trait-relevant tissues and cell types, which are only biological important for the given traits in certain physiological stages or environmental conditions.

Due to the large amount of linkage disequilibrium (LD) among genomic markers within a single cattle breed (e.g., Holstein), traditional single-component prediction models (e.g., GBLUP and BayesA), which assume that all markers are drawn from the same prior distribution, work quite well within breed (Meuwissen et al. 2001). Such high LD within breed and the highly
polygenic architecture of economic traits also make it hard to partition genomic variance into distinct components accurately in a linear mixed model framework, due to the potential high correlations among components. When incorporating tissue-specific genes into the extended prediction models, we thus observed a limited increase in prediction accuracy within Holstein compared to the traditional model, consistent with our previous findings (Fang et al. 2017a,b). However, this functional information may contribute much more to genomic prediction in other scenarios where reduced relatedness is observed between reference and target populations, such as multiple breeds and over generations (Liu et al. 2015; MacLeod et al. 2016; Fang et al. 2017a,b). In addition, when a large range of biological priori information is available in the future, we may use GWAS enrichment analysis as a guide to choose the most ‘relevant’ biological priori information for genomic prediction, as genomic prediction is often computationally intensive (Fang et al. 2017a).

Methods

Bioinformatics analysis of second-generation sequencing data

In this study, we collected all 156 samples under the approval of the U.S. Department of Agriculture Agricultural Research Services Institutional Animal Care and Use Committee under the Protocol 16-016. We provided references where RNA-seq data were retrieved (i.e., SRP042639, PRJNA177791, PRJNA379574, PRJNA416150, PRJNA305942, PRJNA3922196, PRJNA428884, PRJNA298914, PRJEB27455, PRJNA268096, and PRJNA446068) and summarized details of all 723 analyzed RNA-seq samples in Supplemental Table S1. Among the newly generated data, we collected 51 from six Holstein cows (GSE137943, GSE148707), 94 from the sequenced Hereford cow (L1 Dominette 01449) and its relatives (GSE129423) (Fang et al. 2019). Briefly, we extracted the total RNA from snap-frozen tissues using TRIzol (Thermo Fisher Scientific) according to the manufacturer’s instructions. We measured the quantity and purity of RNA using a NanoDrop 8000 Spectrophotometer (NanoDrop Technologies) and Agilent 2100 Bioanalyzer System (Agilent). We sequenced these RNA samples using the Illumina HiSeq 2000 platform (Illumina) with paired-end (100- to 150-bp) reads for most of them and single-end reads for the rest (Supplemental Table S1).

We analyzed all 723 RNA-seq data uniformly using the following bioinformatics pipeline. First, we removed contaminating adapter molecules, reads containing poly(N), and low-quality reads using Trimmomatic (version 0.38) (Bolger et al. 2014), obtaining a total of 18,468,126,120 clean reads. We then mapped clean reads to the cattle reference genome UMD3.1.1 using HISAT2 (version 2.1.0) (Kim et al. 2015), resulting in an averaged uniquely mapping rate of 94.18% (Supplemental Table S1). We used Ensembl genes (release 94) as the gene annotation file, including 24,616 genes. We determined gene expression levels (i.e., FPKM) using StringTie (version 1.3.4) (Pertea et al. 2015), while accounting for differences in sequence depth and gene length across samples.

Based on known biology (Harhay et al. 2010), we classified 91 tissues and cell types into 17 biological categories. In order to detect tissue-specific expression signatures, we computed a t-statistic for each gene in a given tissue using the following approach (Finucane et al. 2018), by excluding tissues and cell types in the same biological category while accounting for known covariates (i.e., age, sex, and study) (Supplemental Table S1). We scaled the log_2-transformed expression (i.e., log2FPKM) of genes to have a mean of zero and variance of one within each tissue and cell type.

Figure 5. Relationships between milk production traits and brain regions. (A) Milk production traits have a significantly higher GWAS signal enrichments (−log10P) than other types of traits in 14 brain regions (CNS), except for feed efficiency (i.e., residual feed intake [RFI]). We calculate P-values between groups using Student’s t-test. (B) Two fine-mapped genes, TRIM46 (top; posterior probability of causality [PPC] = 0.59) and RAB6A (bottom; PPC = 0.79), for protein percentage and milk yield, respectively, are specifically highly expressed in CNS compared with all other tissues and cell types. (C) The associations of milk production traits with brain regions and four brain endocrine tissues (i.e., stalk median eminence [SME], anterior pituitary, posterior pituitary, and pineal gland) based on the GWAS signal enrichments of tissue-specific genes detected within these brain-relevant tissues. (*) Corrected-P (FDR) < 0.1.
\[ y = \mu + Xb + Zc + e, \]  

where \( y \) is the scaled \log_{2} \text{FPKM}, \( \mu \) is the intercept, \( X \) is the dummy variable for tissue, \( b \) is the corresponding tissue effect, \( Z \) is the matrix for covariables, \( c \) is the corresponding covariable effects, and \( e \) is the residual effect. We fitted this model for each gene in each tissue using the ordinary least-squares approach, as implemented in R (R Core Team 2018), and then obtained the \( t \)-statistic (i.e., the coefficient, \( b \), divided by its standard error) for each gene to measure its expression specificity in the corresponding tissue. We employed the same approach to pinpoint tissue-specific genes within a biological category of interest (e.g., brain-regions and blood/immune system). We ranked genes in each tissue according to their \( t \)-statistic and chose the top 3\%, 5\%, and 10\% of genes as tissue-specific genes, respectively. We conducted all subsequent analyses using these three cut-offs and obtained similar results. Therefore, we only presented results from the top 5\% in Results.

We conducted the functional enrichment analyses for tissue-specific genes using a hypergeometric test with GO database, as implemented in PANTHER 14.0 (Mi et al. 2012). We obtained the tissue-specific genes of 10 major tissues in humans (https://www.proteinatlas.org/humanproteome/tissue/tissue-specific) and then tested their enrichments with cattle tissue-specific genes among the matched tissues using a hypergeometric test. We conducted the motif enrichment analyses for the promoter regions (i.e., 1500 bp upstream of and 500 bp downstream from the transcriptional start sites [TSSs]) of tissue-specific genes using MEME software (Bailey et al. 2009). For enrichment analyses with cattle QTLdb (Release 36, Aug. 22, 2018) (Hu et al. 2012), we chose QTLs for 19 milk-relevant traits and then arbitrarily considered the gene closest to the lead SNP in each corresponding QTL as the “causal” gene. We thus obtained a list of “causal” genes for each of the 19 milk-relevant traits. We conducted the QTL enrichment analysis for tissue-specific genes using the same hypergeometric test like the GO enrichment analysis. For DNA methylation data (Zhou et al. 2018), we also mapped them to the cattle reference genome UMD3.1.1 using Bismark v0.19.0 (Krueger and Andrews 2011). We only kept CpG sites with at least fivefold coverage for subsequent analyses. We employed an entropy-based framework to determine tissue-specific DNA methylation regions, as implemented in the SMART2 software (Liu et al. 2015). We only considered tissue-specific hypomethylated regions to validate our results of trait-relevant tissues obtained by using tissue-specific genes, because hypomethylation is generally related to gene activation (Jones 2012). We determined the promoter methylation level for each gene as the average methylation of CpG sites within its promoter region as defined above. We then obtained the adjusted promoter methylation in a tissue by adjusting for the averaged methylation over the entire genome in this particular tissue. For comparing gene expression among cattle, sheep, and human, we retrieved multi-tissue gene expression for human from GTEx v6 https://gtexportal.org/home/datasets, and for sheep from https://doi.org/10.1371/journal.pgen.1006997.s004. We obtained the ortholog genes among mammals from Ensembl v94 (https://www.ensembl.org/info/website/archives/index.html).
Single-marker GWAS and fine-mapping results

We previously reported details of the single-marker GWAS and fine-mapping analyses for body type, reproduction, production, and health traits from 27,214 U.S. Holstein bulls (Jiang et al. 2019; Freebern et al. 2020) and for feed efficiency (i.e., RFI) from 3947 Holstein cows (Li et al. 2019). Briefly, we used de-regressed breeding values (predicted transmitting abilities [PTA]) of Holstein bulls as phenotypes. We have adjusted such phenotypes for all known systematic effects, including herd, year, season, and parity (Norman et al. 2009). For feed efficiency, we corrected for the dry matter intake for milk yield, metabolic body weight, body weight change, and several environmental effects to obtain RFI (Lu et al. 2015). We used the high-density genotypes (777K) and imputed sequence markers (n = 2,619,418) with an imputation accuracy of 96.7% (Vanraden et al. 2017), minor allele frequency (MAF) > 0.01, and Hardy-Weinberg Equilibrium (HWE) test (P > 10^-6) to conduct GWAS analyses for RFI and the remaining traits, respectively. We employed the following linear mixed model, implemented in MMAP software (https://mmap.github.io/), to test for association of genomic variants with all complex traits except for RFI:

\[ y = \mu + Xb + g + e, \]  
\[ \text{(2)} \]

where \( y \) is the de-regressed PTA, \( \mu \) is the overall mean, \( X \) is the genotype of a genomic marker (coded as 0, 1, or 2), \( b \) is the marker effect, \( g \sim N(0, \sigma_g^2 G) \) is the polygenic effect accounting for familial relationship and population structure, and \( e \sim N(0, \sigma_e^2 R) \) is the residual. \( G \) is the genomic relationship matrix (Vanraden 2008), built using HD markers with MAF=0.01. \( R \) is a diagonal matrix with \( R_{ii} = 1/t_i^2 - 1 \), where \( t_i^2 \) is the reliability of phenotype for the \( i \)-th individual. For RFI, we used a single-step method to conduct GWAS analysis, which was implemented in the BLUPF90 (version 2018) (Wang et al. 2012; Li et al. 2019).

GWAS signal enrichment analysis

Because complex traits being studied here are highly polygenic (Cole et al. 2009; Kemper and Goddard 2012; Boyle et al. 2017), we applied the following sum-based marker-set test approach, as implemented in the QGG package (Rohde et al. 2019), to determine whether GWAS signals were enriched in tissue-specific genes. We added 10-kb, 20-kb, and 50-kb windows around gene regions to include the potential cis-regulatory variants. Previous studies showed that this approach had at least equal power when compared to other commonly used GWAS signal enrichment methods in humans (Rohde et al. 2016), Drosophila melanogaster (Sørensen et al. 2017), and livestock (Sarup et al. 2016; Fang et al. 2017a,c), especially for the highly polygenic traits.

\[ T_{\text{sum}} = \sum_{i=1}^{m} b_i^2, \]  
\[ \text{(3)} \]

where \( m \) is the number of genomic markers within a list of tissue-specific genes, and \( b \) is the marker effect from single-marker GWAS. We controlled marker-set sizes and LD patterns among markers through applying the following genotype cyclical permutation strategy (Rohde et al. 2016; Sørensen et al. 2017). Briefly, we first ordered marker effects (i.e., \( b_i^2 \)) using their chromosome positions (i.e., \( b_1^2, b_2^2, \ldots, b_{m-1}^2, b_m^2 \)). We then randomly selected one marker (i.e., \( b_k^2 \)) from this vector as the first place and shifted the remaining ones to new positions, while retaining their original orders (i.e., \( b_1^2, b_2^2, \ldots, b_{i-1}^2, b_i^2, b_{i+1}^2, \ldots, b_{m-1}^2, b_m^2 \)). We calculated a new summary statistic for given tissue-specific genes using their original chromosome locations. To obtain an empirical \( P \)-value for a list of tissue-specific genes, we repeated this permutation procedure 10,000 times and employed a one-tailed test of the proportion of random summary statistics greater than that observed.

For comparison, we also employed the following count-based approach that focused on the top variants passing a certain genome-wide significance level:

\[ T_{\text{count}} = \sum_{i=1}^{m} I(p_i < p_0), \]  
\[ \text{(4)} \]

where \( m \) is the number of markers in the tested tissue-specific gene list, \( p_i \) is the \( P \)-value for the \( i \)-th marker from single-marker GWAS, \( p_0 \) is an arbitrarily selected significant threshold, and \( I \) is an indicator function that takes value one when \( p_i < p_0 \) and value zero otherwise. Here, we chose \( p_0 = 0.01 \) as the significant cut-off. Under the null hypothesis, we assumed that \( T_{\text{count}} \) follows a hypergeometric distribution (Sørensen et al. 2017). We observed that results from these two GWAS enrichment approaches showed a positive correlation of 0.68 (Supplemental Fig. S16A). We here focused on results of a sum-based method. As results of 10-kb and 20-kb extensions were similar to those of 50-kb (Supplemental Fig. S16B), we only showed results of the 50-kb extension.

SNP-set-based genomic prediction analysis

We divided the entire Holstein cattle population into the reference population (n = 19,575) and the validation population (n = 3983) according to the year of birth (Vanraden et al. 2017). We applied the SNP-set-based genomic prediction (SSGP) software to incorporate tissue-specific genes into genomic prediction (https://sites.google.com/view/ssgp), which allows us to split genomic markers into different groups with group-specific effect variance.

\[ y = \mu + \sum_{k=0}^{p} K_{kh} \epsilon_k + e, \]  
\[ \text{(5)} \]

where \( y \) is the phenotype vector (i.e., PTA), \( \mu \) is the population mean, and \( p \) denotes the number of genetic components in the model. Here, we choose \( p = 2 \), corresponding to random effects for markers within the category-specific genes and the remaining genome, respectively. The random effects within the \( h \)-th component are assumed to follow a multivariate normal distribution: \( \epsilon_h \sim \text{MVN}(0, \Sigma_h \sigma^2) \); \( h = 0 \) or 1, where \( \Sigma_h \) is a predefined diagonal matrix to weight each of the random effects, and \( \sigma^2_e \) is assumed to follow an inverse-gamma distribution \( \sigma^2_e \sim \text{Inv} - \text{Gamma}(a_{\sigma}, b_{\sigma}) \), where \( a_{\sigma} \) and \( b_{\sigma} \) are scale and shape parameters, respectively. \( K_h \) is the corresponding design matrix. \( e \) is the residual effect following normal distribution \( e \sim \text{MVN}(0, \Sigma_e) \), where \( \Sigma_e \) is the diagonal matrix with a predefined weight for error variance. To make models comparable (i.e., the same number of parameters to be estimated), we fitted another model with two genomic components as a null model, where all 24,616 genes as the first component and the rest of the genome as the second component. We also fitted a single-component model that was equivalent to the GBLUP model. We determined the prediction accuracy as the correlation between predicted PTAs and true PTAs in the validation population.

Data access

All raw and processed sequencing data generated in this study have been submitted to the NCBI Gene Expression Omnibus (GEO; https://www.ncbi.nlm.nih.gov/geo/ under accession numbers GSE128075, GSE137943, GSE147087, GSE147184, and GSE148707. The GWAS summary statistics for all complex traits
have been submitted to Figshare, that is, body type, production, and reproduction traits under https://figshare.com/s/ea726fa95 a5bacc158ac1, and the remaining ones under https://figshare .com/s/94540148512d1d7ed32. All scripts and source codes can be found as Supplemental Code, as well as at the Cattle Gene Atlas (http://cattlegenatlas.roslin.ed.ac.uk) and GitHub (https://github.com/LingzhaoFang1/Cattle-GeneAtlas).

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