Allele-specific expression reveals the phenotypic differences between thin- and fat-tailed sheep

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Abstract

Background: The thin-tailed sheep breeds from Europe and the fat-tailed sheep breeds from China exhibit distinct phenotypic differences in fat deposition and meat production traits. However, the molecular mechanisms underlying gene expression related to these phenotypic differences are not well understood. Allele-specific expression (ASE) refers to the significant imbalance of expression levels of two parental alleles. Characterization of such events in F1 hybrid offspring generated from these two groups of sheep breeds can minimize the external factors influencing gene expression and reveal the variants with a *cis*-regulatory effect on gene expression. The aim of the present study was to investigate the genetic factors that influence different fat-deposition and meat production traits between thin- and fat-tailed sheep.

Results: Fifteen F1 hybrids were generated from crosses between Texel and Kazakh sheep as the representative phenotypes of thin- and fat-tailed breeds, respectively. Totally, 33 whole genomes from F1 individuals and their parents were sequenced with an average depth of ~17.21× coverage per sample. ASE analysis results from 70 RNA-seq samples of adipose and skeleton muscle tissues showed 128 ASE candidate genes were related to the function of fat deposition and meat production traits. A genome-wide scan of selective sweeps was also conducted between these two groups of sheep breeds in an effort to identify genomic regions related to fat deposition and meat production, respectively. We detected signatures of selection in ASE genes associated with fat deposition (e.g., *PDGFD*) and meat production traits (e.g., *LRCC2*). Further analysis suggested that *PDGFD* and *LRCC2* genes were speculated to be causative
genes for fat deposition and meat production traits in sheep, respectively. Furthermore, AMPK signaling pathway was significantly enriched in ASE genes related to fatty acid biosynthesis in both adipose and skeleton muscle tissues, while PPAR signaling pathway was significantly enriched in ASE genes related to lipid metabolism in adipose tissue.

**Conclusions:** Our finding illustrates that the expression of identified ASE genes could potentially lead to the differences in traits of fat deposition and meat production between thin- and fat-tailed sheep.

**Keywords:** allele-specific expression, phenotypic difference, thin- and fat-tailed sheep, whole-genome sequencing, transcriptome
The sheep (*Ovis aries*) was one of the first grazing animals to be domesticated approximately 11,000 years ago in the Fertile Crescent [1]. Since domestication, a wide variety of sheep breeds with different production traits (e.g. meat, milk and fiber) and morphological characteristics (e.g. coat color and tail shape) have been shaped under the influence of natural and artificial selection [2]. According to the fat content in tail, sheep can be mainly classified into two groups: thin- and fat-tailed breeds. Fat tailed sheep breeds comprise approximately 25% of the world sheep population, which are first documented approximately 5,000 years ago [2]. They are generally well adapted to the dryland environments with seasonal poor nutrition [3, 4]. These sheep breeds are known for their ability to deposit up to 20% of the carcass weight as tail fat [5]. The tail fat plays as a valuable source of energy during migration and winter. However, nowadays with increasing attention to human health, fat is an unpopular constituent of meat for consumers. On the other side, the energy cost for producing fat is higher than an equivalent amount for lean tissue [6]. Therefore, a small size of fat is often favorable for both consumers and producers. Thus, identification of functional candidate genes and selective genomic regions associated with fat deposition and meat production traits is more important in sheep breeding programs.

Previous genome-wide association studies (GWAS) revealed candidate genes related to fat deposition and meat production traits in different sheep breeds. These studies have proposed candidate genes, such as *MSL1, PFKFB4, TRDN, FBF1, SETD7* and *GRM1* [7, 8]. On the other side, genomic scan for detecting regions associated with
fat deposition were also performed in several sheep breeds [9-17]. However, the molecular mechanisms underlying gene expression related to such economically important traits remained mostly unexplored. DNA sequence variation can lead to changes in gene expression levels, which is a main cause of phenotype diversity across individuals or populations [18]. Variation in the gene expression can be due to both genetic and non-genetic factors. Allele-specific expression (ASE) refers to two or more alleles at the same loci with imbalanced expression, which is one of the important genetic factors that lead to phenotypic variation in organisms [19-22]. ASE is essential for normal development, cellular programming and many other cellular processes [23]. Therefore, identification of loci involved in this phenomenon is important in developmental biology and genetics. The expression ratios of the parental alleles obtained from RNA-seq data of hybrid offspring can be used as a reliable proxy for ASE. Recently, high-throughput sequencing technologies allowed us to identify ASE genes at a transcriptome-wide level. Several studies in different species such as mouse [24], pig [25], cow [26], goat [27], sheep [28], mule [29] and dzo [29] have shown that the different expression levels of alleles due to ASE may lead to variation in phenotypes.

In this study, to understand the genetic mechanisms underlying fat-deposition and meat production traits differences between thin- and fat-tailed sheep breeds, we chose Texel and Kazakh as the representative of two sheep breeds with thin- and fat-tailed, respectively. By integrating the findings from analysis of selective sweeps and gene expression data, several ASE genes associated with growth and development of fat deposition and meat production traits have been detected. Our results suggested that
identified ASE genes may contribute to phenotypic diversity between these two sheep breeds.

Methods

Sample collection and DNA Sequencing

Whole blood samples (10 ml) were collected from three Texel rams, fifteen Kazakh ewes and their fifteen F1 hybrids in Yining City of Xinjiang, China. DNA was isolated from whole blood using standard phenol-chloroform method [30]. The quality of genomic DNA was measured by NanoDrop 2000 (Thermo Fisher Scientific Inc., Wilmington, DE, USA). The total amount of genomic DNA from each sample was at least 8 ug and the concentrations of DNA was higher than 50 ng/ul. Paired-end sequencing libraries were conducted using the Illumina HiSeq2000 platform.

SNP genotyping and quality control

High-quality clean reads were obtained after removing adaptors and low-quality reads. Pair-end reads passing the filtering were aligned to the Ovis aries v4.0 reference genome using BWA software (v0.7.13, bio-bwa.sourceforge.net) [31, 32]. Heterozygous sites from F1 hybrid individuals were identified by applying the Genome analysis toolkit (GATK, v3.3-2) [33]. Low-quality variants were filtered using the parameter QUAL < 30, and the variants were annotated using ANNOVAR (Version: 2013-08-23) [34].
ASE analysis from RNA-seq data

A total of 70 transcriptome data generated from adipose tissue including subcutaneous fat \((n=14)\), tail fat \((n=15)\) and from skeleton muscle tissue including longissimus dorsi \((n=13)\), semimembranosus \((n=15)\), semitendinosus, \((n=13)\) of the F1 hybrid animals were used to identify the SNP with ASE \([35]\). All BAM files from these samples were then generated by mapping RNA-Seq data to the *Ovis aries* v4.0 reference genome, by using HISAT2 software \([36]\). Allele-specific reads at the heterozygous sites were obtained from these BAM files using the WASP software \([37]\). In order to eliminate the error caused by hetero-allele or repeat-sequence of the reference genome, loci of both alleles with \(< 10\) reads were filtered out, and genes with min 2 and max 100 heterozygous loci were selected \([28]\). Loci with both alleles \(< 100\) reads and each allele in heterozygous sites with \(< 3\) reads were filtered, which can remove loci exhibiting MAE and a relative lower expression \([38-40]\). Allelic read ratios were counted by a Python Script \([27]\), significant allelic imbalances were identified using the cut-off criteria of allele ratio \(\geq 0.65\) or \(\leq 0.35\) \([29]\), and statistical tests were conducted by Chi square test and Benjamini-Hochberg FDR correction. Imprinted genes were excluded according to the online database (http://www.geneimprint.com/site/home).

Functional enrichment analysis

We then employed KOBAS (v3.0) online tool (http://kobas.cbi.pku.edu.cn/) to conduct ASE genes enrichment analysis. Reference genes of *Ovis aries* v4.0 genome were used as background gene list, and corrected \(P\)-values were calculated by Chi-square test and
Benjamini-Hochberg methods [32]. Statistically, over-represented categories were determined by standard false discovery rate (adjust $p$-value < 0.05).

**Differential expression analysis**

The transcriptome data from longissimus dorsi of Texel and Kazakh sheep were downloaded and then trimmed using Trimmomatic to remove low quality reads [32, 35, 41]. BAM files were then obtained after mapping the high quality clean reads to the *Ovis aries* v4.0 reference genome [32], and DEGs were identified using R package DESeq 2 (version 1.16.1) [42].

**Selective sweeps analysis**

We downloaded a total of 52 sheep whole-genome sequencing data including 20 fat-tailed sheep of two breeds from Xinjiang, China, 15 improved thin-tailed sheep of seven breeds from Europe and 17 wild Mouflon sheep (*Ovis orientalis*) from the Middle East (Additional file 1: Table S1) [43, 44]. Selective sweeps between these thin- or fat-tailed sheep breeds and Mouflon sheep populations were scanned using a genome-wide sliding window strategy based on the SNPs with less than 10% missing data [45]. Population differentiation statistic ($F_{ST}$) between the selected thin- or fat-tailed sheep and wild Mouflon sheep were calculated using a 50 kb sliding window with 10 kb sliding step by vcftools v0.1.14, respectively [46]. To detect genomic regions under selection in thin- or fat-tailed sheep, we calculated the $\ln(\theta\pi_{\text{Mouflon}}/\theta\pi_{\text{thin-tailed sheep}})$ and $\ln(\theta\pi_{\text{Mouflon}}/\theta\pi_{\text{fat-tailed sheep}})$, respectively. Values of $\theta\pi$ in thin-tailed, fat-tailed sheep
and wild Mouflon sheep were calculated using the same parameters as calculating the $F_{ST}$ by vcftools v0.1.14 [46]. In each comparison, the genomic regions in the top 1% $F_{ST}$ values, 1% $\ln(\theta_\pi^{-\text{Mouflon}}/\theta_\pi^{-\text{thin-tailed sheep}})$ and top 1% $\ln(\theta_\pi^{-\text{Mouflon}}/\theta_\pi^{-\text{fat-tailed sheep}})$ values across the whole genome were considered to be putative selective sweeps. Combing the results of $F_{ST}$ values and $\theta_\pi$ ratio, the genes located in selective sweeps which were simply found in thin- or fat-tailed sheep used for subsequent analysis.

Results

The phenotypic differences of growth and meat production traits between Texel and Kazakh sheep breeds

Texel and Kazakh sheep were selected as parental breeds to establish F1 hybrid population. High phenotypic variations were observed between these two breeds: Texel sheep are characterized by well-developed skeleton, high mutton performance, wide and deep chest and relatively higher lean meat ratio, whereas Kazakh sheep have low meat production rate, slower growth rate and relatively higher content of fat in their carcass (Fig. 1a and 1b). In this study, we collected several phenotypic data related to fat deposition and meat production traits. From the above records, nine traits were concerned finally: weights of tail fat, longissimus dorsi, silverside, topside, rump, total fat and total meat; thickness of back fat and back muscle. As shown in Fig. 1 and Additional file 2: Table S2, the weights of tail fat, total fat and back fat thickness in Kazakh sheep are higher than those in Texel sheep. The average of tail fat weight estimated in Kazakh sheep (3252.0 g) was remarkably heavier than that of the Texel
sheep (46.71 g) (Fig. 1a, t-test, $P$ value < 0.05). However, the average weights of longissimus dorsi, back muscle thickness, silverside and topside from Texel sheep were significantly heavier than that from the Kazakh sheep (Fig. 1f-i, t-test, $P$ value < 0.05). The detailed differences of phenotype traits between Texel and Kazakh sheep breeds provide a rational framework in which to examine the genetic variation between these two breeds.

**Identification of ASE SNPs in the F1 hybrids of Texel and Kazakh sheep**

To eliminate the effects of differences in growth and development between Texel and Kazakh sheep, we compared the relative allelic expression in F1 hybrids of these two sheep breeds. Three Texel rams, fifteen Kazakh ewes and their fifteen F1 hybrid offspring were selected for whole genome re-sequencing analysis. Samples were sequenced with an average depth of $\sim 17.21 \times$ coverage (Additional file 3: Table S3). We then estimated that each F1 hybrid contained an average of 7.55 Mb heterozygous SNPs, by which the two parental alleles can be distinguished (Additional file 4: Table S4). Among those heterozygous sites, about an average of 1.83% were found in exon regions of genes. In addition, we detected an average of 59.05%, 34.03% and 1.55% heterozygous sites in intergenic, intron and gene up/downstream regions of genome, respectively (Fig. 2a, Additional file 4: Table S4). A total of 810 ASE SNPs were identified from the samples of adipose or skeleton muscle tissue, 60 (7.41%) of the total ASE SNPs were located on upstream and downstream regions of genes (Fig. 2b, Additional file 4: TableS4). Totally, 605 (74.69%) of the total ASE SNPs were found
on exon regions of genes, in which 303 (50.08%) were located on the untranslated regions (UTR) of genes, 155 (25.62%) were synonymous mutation, 73 (12.07%) were non-synonymous mutation and 74 (12.23%) were unknown mutation (Fig. 2b, Additional file 5: Table S5). It has been suggested that these ASE SNPs are located on untranslated regions of genes, which can act as cis-regulatory elements (CREs) to regulate the transcription of a neighboring gene [47]. On the other side, the proportion of transition (77.59%) was three times more than that of the transversion (22.41%) (Fig. 2c, Additional file 4: Table S4). And the ratio of transition and transversion was 3:1, which is similar to the previous studies [48].

**Analysis of ASE genes in adipose tissue**

A total number of 106 ASE genes were identified in the samples of adipose tissues, in which 48 (45.28%) common ASE genes were found in both tail fat and subcutaneous fat samples. We identified 42 (39.62%) and 16 (15.09%) unique ASE genes from tail fat and subcutaneous fat samples, respectively (Fig. 3a, Additional file 5: Table S5). Additionally, we found 89 (83.96%) of 106 ASE genes have less than 10 ASE SNPs, and 17 (16.04%) of these ASE genes have more than 10 but less than 51 ASE SNPs (Fig. 3b, Additional file 5: Table S5). According to the previous report, in which the authors identified differentially expressed genes (DEGs) in samples of tail fat between thin- and fat-tailed sheep breeds, here we also found two up-regulated (*PDGFD* and *IRF2BP2*) and one down-regulated ASE genes (*TEN1*) in samples of tail fat, by comparing these two breeds [15]. *IRF2BP2* gene is located in the quantitative trait loci
QTL which is related to tail fat deposition, and TENI gene was considered to be associated with tail fat weight trait in Hulun Buir sheep [49]. A total of two and three ASE SNPs were found in IRF2BP2 and TENI genes, respectively. The expression of ASE SNPs identified in IRF2BP2 showing consistent in both tail fat and subcutaneous fat samples, and also for TENI gene, we found the SNPs showing consistent ASE in tail fat samples (Additional file 5: Table S5).

In order to further investigate the potential functional and metabolic process of these identified ASE genes and DEGs, we performed Kyoto Encyclopedia of Genes and Genomes (KEGG) enrichment pathway analysis and found Chemokine signaling, AMPK signaling and PPAR signaling pathways were significantly enriched (Fig. 3c). Four ASE genes (PPARG, PCK1, LPL and SCD genes) and two DEGs (LOC101115115 and FABP5) were significantly annotated in PPAR signaling pathway (Fig. 3d). In addition, six ASE genes (PCK1, PFKFB1, PFKFB4, PPARG, SCD and SLC2A4 genes) and two DEGs (RAB2A and CREB3L1) were significantly annotated in AMPK signaling pathway. In terms of directions of ASE in both tail fat and subcutaneous fat samples, we found PFKFB1 gene showing consistent in samples from both tail fat and subcutaneous fat, PCK1 and SCD genes showing consistent in subcutaneous fat samples, while PPARG and SLC2A4 genes showing inconsistent in tail fat samples. Besides this, PFKFB4 and LPL genes were also found showing consistent in some specific subcutaneous fat and tail fat samples (Additional file 5: Table S5). The above results elucidated that DEGs and ASE genes which enriched in PPAR signaling and AMPK pathways may contribute to the difference of lipid metabolism in adipose tissue.
between thin- and fat-tailed sheep.

**Analysis of ASE genes in skeleton muscle tissue**

The analysis identified a total of 47 ASE genes, in which 15 (31.91%) of them were shared by longissimus dorsi, semitendinosus and semimembranosus samples. Also, 13 (27.66%), 4 (8.51%) and 4 (8.51%) ASE genes were found in semimembranosus, semitendinosus and longissimus dorsi samples, respectively (Fig. 4a, Additional file 4: Table S4). We found that 40 (85.11%) of 45 ASE genes have less than 10 SNPs and the other 7 (14.89%) ASE genes have more than 10 but less than 30 SNPs (Fig. 4b, Additional file 5: Table S5). Compared with ASE genes identified in adipose tissue, 21 (17.07%) ASE genes were private to skeleton muscle tissue and 24 ASE genes were common in both of these two tissues (Fig. 4c, Additional file 5: Table S5). It can be seen that majority of ASE genes were more inclined towards tissue-specific. Among the identified ASE genes in skeleton muscle tissue, *EPM2A*, *LRRC2*, *PFKFB4* genes were thought to be associated with meat production traits in sheep [7]. *EPM2A* gene showing consistent in samples from semitendinosus and longissimus dorsi. *LRRC2* and *PFKFB4* genes showing consistent in some specific samples of semimembranosus, while *PFKFB4* genes showing inconsistent in any semitendinosus samples. The above analysis illustrated that some identified ASE genes showed inconsistent ASE in different tissues or different samples from the same tissue. It is suggested that one of the alleles may function better in one specific tissue or one sample type. The F1 hybrids of Texel × Kazakh sheep may able to use the better allele in growth and development,
eventually leading to higher performance in their meat production traits.

**Identification of ASE genes for directly comparison between Texel and Kazakh sheep**

We also downloaded the transcriptome data of longissimus dorsi of Texel and Kazakh sheep and identified DEGs between these two sheep breeds. As we expected, we found a total of 4,446 genes showing significant differential expression between skeleton muscle tissues from Texel and Kazakh breeds; including 1,988 down-regulated and 2,458 up-regulated (adjust p-value ≤ 0.05, log2 RPKM ≥ 1 and log2 Fold change ≥ 1) (Additional file 6: Table S6). In order to investigate pathways associated with meat production traits between thin- and fat-tailed sheep breeds, we then conducted an integrated functional analysis of ASE genes identified in skeleton muscle tissues of F1 hybrids and DEGs as mentioned above. Several signaling pathways such as MAPK signaling, Rap1 signaling, AMPK signaling and Notch signaling were significantly enriched for those ASE genes and DEGs (Fig. 4d). *FASN* and *PFKFB4* ASE genes were annotated in 5′-adenosine monophosphate-activated protein kinase (AMPK) pathway, which plays a key role in the regulation of muscle regeneration and size [50]. *FASN* gene could promote the synthesis of long chain saturated fatty acids as the rate-limiting enzyme, SNPs in this gene are associated with fatty acids content in meat and fat tissues [51]. Two ASE SNP were found in *FASN* gene, which were located in the exonic region of the genes and the directions of these two ASE SNPs were consistent (Additional file 5: Table S5). Moreover, we found *FASN* gene and other eight ASE genes were down-
regulated in longissimus dorsi samples of Texel sheep, as compared to those of in
Kazakh sheep (Fig. 4e). These results implied that AMPK signaling pathway is an
important factor in regulation of different energy metabolism in skeleton muscle tissue
between thin- and fat-tailed sheep breeds.

**Investigation of ASE genes located in the selected regions**

In order to investigate whether the identified ASE genes were under selection in thin-
or fat-tailed sheep, we performed whole genome sequences analysis between 17 wild
Mouflons sheep and 20 fat-tailed sheep from Xinjiang and 15 improved thin-tailed
sheep from Europe, respectively. By combing the results from population-
differentiation statistic ($F_{ST}$) and $\ln(\theta\pi^{\text{Mouflon}} / \theta\pi^{\text{fat-tailed sheep}})$ tests, a total of 176
putative selective genomic regions with an average size of 100.97 kb and comprised
with 254 genes were found in the fat-tailed sheep (Additional file 7: Table S7). We
further found six ASE genes including *PDGFD, PPDPF, PM20D1, TUBA8, PPM1K*
and *PPA2* genes were selected in the fat tailed sheep breeds (Fig. 5a). As a member of
the platelet derived growth factor (PDGF), *PDGFD* gene plays an important role in
regulation of adipocyte function and its expression is closely related to the fat
deposition in sheep tail [15, 52]. From eight samples of adipose tissue, seven ASE SNPs
were found in *PDGFD* gene. Five SNPs were located in the 3′-UTR region and the
remaining two SNPs were located in the exon regions. In addition, the third and fourth
ASE SNP were sample-specific, but the other ones were at least with two samples (Fig.
5b, Additional file 4: Table S4). Notably, the genotype patterns located in the upstream
region of PDGFD gene were quite different between the wild Mouflon sheep and fat-tailed sheep (Fig. 5b).

In addition, a total number of 194 putative selective sweep regions which with an average size of 86.80 kb and comprised 256 genes were identified in the thin-tailed sheep according to the values of $F_{ST}$ and $\ln(\theta_{\pi}^{-\text{Mouflon}} / \theta_{\pi}^{-\text{thin-tailed sheep}})$ (Additional file 8: Table S8). Out of these selected genes, we found seven genes ($LRRC2$, $BIN1$, $LOC101118164$, $IRF2BP2$, $SFRP1$, $LOC1011045430$ and $BCO2$) showing ASE (Fig. 6a). $LRRC2$ gene is mainly expressed in samples of longissimus dorsi, semitendinosus and semimembranosus from sheep and is located in the QTL that is associated with muscle density [49, 53]. A total of nine ASE SNPs were identified in the 3’-UTR region of $LRRC2$ gene, four of these ASE SNPs were shared by at least two samples and the remaining ASE SNPs were found with sample-specific (Fig. 6b, Additional file 4: Table S4). The genotype pattern of selection sweep region located in $LRRC2$ gene were different between wild Mouflon sheep and thin-tailed sheep (Fig. 6c) These results suggested that those ASE genes which were selected in the thin- or fat-tailed sheep may contribute the different fat deposition and meat production traits between thin- and fat-tailed sheep.

Discussion

The great variations in different phenotypic traits of two selected sheep breeds provide an opportunity for studying molecular mechanisms underlying gene expression related to phenotypic difference between thin- and fat-tailed sheep breeds. In this study, the
candidate causative genes were detected by ASE analysis, which avoid the effect of
trans-acting and has become research focus in recent years [24-28]. We divided all
sheep individuals into 15 family trios. Each family consists of one Texel ram, one
Kazakh ewe and their F1 hybrid offspring. All these individual genomes were then re-
sequenced and used to find heterozygous loci. The unbiased mapping BAM files from
allele-specific reads of RNA-seq data were produced by the aids of WASP software,
which can map the reads to the correct location of reference genome with a low error
rate [37]. A total of 128 ASE genes were identified from the collected samples of
adipose and skeleton muscle tissues, the expression profile of majority ASE genes in
F1 hybrid tissues were tended to be tissue-specific. In a recent study on analysis of ASE
in immune-related tissues, the authors reported individual-specific and tissue-specific
gene expression for several genes in a crossbred sheep breed (Texel × Scottish
Blackface) [28]. In mouse, Pinter et al. (2015) showed that 95% and 82% of the
identified ASE genes in F1 hybrids and in a reciprocal cross were also with tissue
specificity, respectively [54]. Two alleles in rice hybrids might play different role in
different environmental conditions and developmental stages, a higher proportion of
ASE genes in those hybrids showed inconsistent patterns [55]. In the current study, 22,
81 and 25 ASE genes were identified in samples from skeleton muscle, adipose and
both sharing tissues, respectively. Many of these genes showed inconsistent patterns in
terms of directions of ASE bias across different samples. It can be speculated that one
allele may prefer to one tissue, while the other one prefer to other tissues in one case.
And in the other case, one allele might be inclined to one sample, while the other one
might be inclined to other samples in the same tissue.

Comparative transcriptome analysis is another efficient way used for identifying candidate causative genes that associated with distinct phenotypic differences between thin- and fat-tailed sheep breeds [56-58]. However, it presented little power to identify genetic factors that contribute to these phenotypic differences because the identified DEGs always reflect both cis- and trans-acting regulatory variations [59]. In addition, the different growing conditions between these two groups of sheep breeds can also influence gene expression. Among the identified DEGs in the skeleton muscle tissue between thin- and fat-tailed sheep breeds, we found that three (TEN1, PDGFD and IRF2BP2) and nine (FASN, CD68, CIR1, EEF1A1, EHD1, GAA, RGMB, SMTNL1 and GLUL) DEGs showing ASE in selected samples of adipose and skeleton muscle tissues between thin- and fat-tailed sheep breeds, respectively. All of the nine ASE genes were down-regulated in longissimus dorsi samples of Texel sheep, in which the expression level of FASN gene was negatively correlated with intramuscular fat (IMF) content in the longissimus of Kazakh sheep [60]. TEN1 gene showing ASE in samples collected from both adipose and skeleton muscle tissues, and all of the ASE SNPs identified in this gene were located in 5'-UTR region. Promoters are typically located in 5'-UTR region of the gene and mutations in promoters often cause disease in human [61]. Overexpression of PDGFD gene in sheep pre-adipocytes could significantly increase the expression levels of PPARG and LPL genes [52]. Three ASE SNPs were found in PPARG gene, which can control the expression of genes related to adipocyte differentiation and lipid metabolism as an important transcription factors [62]. It was
also involved in the intramuscular adipocyte differentiation in the early growing stage
of cattle [62]. A total of 15 ASE SNPs were identified in LPL gene from eight samples
of adipose tissues, and its expression in the longissimus dorsi of sucking lambs could
be affected by fatty acid indicators including polyunsaturated fatty acid (PUFA),
saturated fatty acid (SFA), conjugated linoleic acid (CLA) and n-6/n-3 according to the
difference of forage type and lamb sex [63]. Results of KEGG pathway enrichment
analysis from identified ASE genes and DEGs in adipose and skeleton muscle tissues
revealed enriched categories related with AMPK signaling pathway. Activating AMPK
could inhibit fatty acid synthesis while promote fatty acid oxidation and regulate brown
adipogenesis [64, 65]. It also plays a crucial role in the regulation of skeletal muscle
development [50]. Besides this, PPAR signaling pathway was significantly enriched for
ASE genes identified in adipose tissue of F1 hybrids and DEGs that was found in tail
fat samples of between thin- and fat-tailed sheep. Genetic variants of genes in the PPAR
signaling pathway were associated with the traits of porcine meat quality (PMQ) [66].
It is illustrated that PPAR signaling and AMPK signaling pathways may be the key
pathways regulating differential fatty acid profile and lipid metabolism between thin-
and fat-tailed sheep breeds.

Candidate causative genes that contribute to phenotypic differences could also be
identified by selection signals or GWAS, and genomic regions associated with tail fat
deposition and meat production traits have been widely studied in thin- and fat-tailed
sheep populations [7-10, 12, 14, 15, 67]. However, it is hard to discriminate causative
genes as there are many other genes located in genomic region for these traits. Beside
this, the identified selection signals often comprise a higher proportion of false positive
signals. We then chose the overlapping genes between the ASE genes and those that
were selected in the improved thin-tailed sheep from Europe or the fat-tailed sheep from
Xinjiang in order to understand the molecular mechanisms underlying gene expression
associated with different fat deposition and meat production traits between thin- and
fat-tailed sheep. Seven (BCO2, BIN1, IRF2BP2, LRRC2, LOC101118164, LOC101104530 and SFRP1 genes) and six (PDGFD, PPDPF, PM20D1, PPM1K, PPA2 and TUBA8 genes) ASE genes were selected in thin- and fat-tailed sheep, respectively. Among these ASE genes, LRRC2 gene was identified as the important
gene which were thought to be associated with meat production traits at chromosome-wise significance level in sheep [49], while PDGFD gene was considered to be a
primary causal gene for fat deposition in the tails of sheep [15]. A total of nine and
seven ASE SNPs were found in LRRC2 and PDGFD genes, respectively, majority of
these ASE SNPs located in 3'-UTR region of their respective genes. It is widely
accepted that miRNA binds to 3'-UTR region of genes and then regulate gene
expression. The small RNA derived from dominant allele can repress recessive allele
via trans action, and the target-site SNPs of miRNA is associated with ASE SNP of
miRNA target genes [68-70]. Both of these two genes exhibiting ASE may be the result
of miRNA binds to those ASE SNPs in 3'-UTR region. Enhancers were typically
located on downstream (3’) and upstream (5’) of the genes, promoters and enhancers
belong to CREs, and mutation of CREs can generate many phenotypic variants [71-73].
The distinct genotype pattern in the upstream region of LRRC2 and PDGFD genes
maybe another important cause for the gene showing ASE. In addition, we found BCO2 and SFRP1 genes showing ASE in the samples from adipose tissue, while BIN, PM20D1, PPM1K genes showing ASE in the samples from both skeleton muscle and adipose tissues. BCO2 gene plays an important role in the process of β-carotene metabolism [74, 75]. Seven ASE SNPs were identified in BCO2 gene from the samples of subcutaneous fat and tail fat, six of these ASE SNPs were located in exon region of the gene and the remaining one ASE SNP was located in its 3’-UTR region. Knocking out the biallelic BCO2 gene could lead to yellow fat in Tan sheep and a nonsense mutation in this gene is strongly related to the yellow fat phenotype in Norwegian sheep [76, 77]. A total of seven ASE SNPs were found in PM20D1 gene, which involved in the cellular lipid process. Six of these ASE SNPs were located in 3’-UTR region of PM20D1 gene, and the other one was located in exon region of the gene. The miR-324-5p could target the 3’-UTR region of PM20D1 gene, overexpression of miR-324-5p could suppressed the expression of it [78]. PPM1K gene showing ASE in the samples of adipose and skeleton muscle tissues, and the gene located in the QTL that is related to fat and muscle density [49]. Collectively, our findings will expand our knowledge about the molecular mechanisms of gene expression related to fat deposition and meat production variations between thin- and fat-tailed sheep breeds.

Conclusions

The results from this study fill research gap about the genetic basis underlying the different fat deposition and meat production traits between thin- and fat-tailed sheep.
breeds. The combination results of ASE and DEGs provide several candidate genes putatively involved in economically important traits. Overall, the identification of ASE genes will contribute to explain the genetic mechanisms of phenotypic difference between thin- and fat-tailed sheep, especially for tail fat deposition and meat production traits, and lay the research foundation for sheep breeding.

**Supplementary information**

Additional file 1: Table S1 Sample information used for identifying signals of selection in thin- or fat-tailed sheep breeds populations.

Additional file 2: Table S2 The data of fat deposition and meat production traits about Texel sheep and Kazakh sheep.

Additional file 3: Table S3 Samples information used for in this study.

Additional file 4: Table S4 Heterozygous loci identified in each F1 hybrids.

Additional file 5: Table S5 ASE genes identified in this study.

Additional file 6: Table S6 Differentially expressed genes in longissimus dorsi between Texel sheep and Kazakh sheep.

Additional file 7: Table S7 Selection signals identified in the fat-tailed sheep.

Additional file 8: Table S8 Selection signals identified in the thin-tailed sheep.

**Abbreviations**

ASE: allele-specific expression; SNP: Single nucleotide polymorphism; BAM: Binary Alignment MAP; BWA: Burrows-Wheeler Aligner program; GATK: Genome Analysis
Toolkit, GO: Gene ontology; KEGG: Kyoto Encyclopedia of Genes and Genomes; GWAS: Genome-wide association studies; DEGs: Differential expressed genes; RPKM: Reads Per Kilobase per Million mapped reads.

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Authors’ contributions

Y. J. and M. J. L. designed the experiment. S. G. H. and X. Y. P. collected sample and conducted DNA extraction. J. J. S., L. C., Z. R. Y., X. H. and S. G. conducted bioinformatics analysis. J. J. S., X. Y. P., H. A. N., R. L. and Y. W. wrote the manuscript. Y. J. and M. J. L. revised the manuscript. All authors reviewed and approved the final manuscript.

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Availability of data and materials

The whole genome sequence data has been deposited in NCBI short read archive under study PRJNA 623062.

Ethics approval and consent to participate

All animal experiments in this study were carried out in accordance with the regulations of Instructive Notions with Respect to Caring for Experimental Animals, Ministry of Science and Technology of China. All protocols of animal handing and sampling in this study was approved by Experimental Animal Management Committee of the Northwest A&F University (No. 2014ZX08008002). All efforts were made to minimize animal suffering.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no conflict of interests.

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**Figure legend and Tables**

**Fig. 1 Phenotypic difference between Texel and Kazakh sheep.** a-b, Photos from Texel and Kazakh sheep, respectively. c-k, Comparison of fat deposition and meat production traits between Texel (n = 5) and Kazakh sheep (n = 29). Groups with
significant differences were analyzed by two-tailed unpaired \( t \)-test, \( * p < 0.05, ** p < 0.01. \)

Fig. 2 Heterozygous sites identified from F1 hybrids of Texel \( \times \) Kazakh sheep. a, Annotation of the whole genome SNPs; b, Annotation of ASE SNPs identified from the adipose and skeleton muscle tissues; c, Percentage of different substitution types in the ASE SNP.

Fig. 3 ASE genes identified in adipose tissue. a, Venn diagram showing the number of ASE genes identified in between subcutaneous fat and tai fat samples; b, Distribution of ASE SNPs in per ASE gene identified in adipose tissue; c, Results of KEGG pathway enrichment for ASE genes identified in adipose tissue of F1 hybrids and DEGs identified in skeleton muscle tissue between thin- and Kazakh sheep breeds d, ASE genes were annotated in PPAR signaling pathway.

Fig. 4 ASE genes identified in skeleton muscle tissue. a, Venn diagram showing the number of ASE genes identified in samples from longissimus dorsi, semitendinosus and semimembranosus; b, Distribution of ASE SNPs in ASE genes identified in adipose tissue; c, Comparison of ASE genes identified between skeleton muscle and adipose tissues; d, Results of KEGG pathway enrichment for ASE genes identified in skeleton muscle tissue of F1 hybrids and DEGs identified in skeleton muscle tissue between Texel and Kazakh sheep breeds; e, ASE genes were differential expressed in skeleton
muscle tissue between thin- and fat-tailed sheep, * $p < 0.05$, ** $p < 0.01$. Data are showed as mean ± standard error.

Fig.5 Genomic regions with selection sweep signals identified between Mouflon sheep and fat-tailed sheep populations. a, ASE genes were significantly selected by $F_{ST}$ & ln($\theta\pi$-Mouflon / $\theta\pi$-fat-tailed sheep); b, Genotype patterns for the upstream region of $PDGF$ gene between Mouflon sheep (n=17) and fat-tailed sheep (n=20). Detailed information of ASE SNPs identified in $PDGF$ gene (Samples 1-4 from subcutaneous fat, Samples 5-8 from tail fat).

Fig.6 Genomic regions with selection sweep signals identified between Mouflon sheep and thin-tailed sheep populations. a, ASE genes were significantly selected by $F_{ST}$ & ln($\theta\pi$-Mouflon / $\theta\pi$-thin-tailed sheep); b, Detailed information of ASE SNPs identified in $LRCC2$ gene (Sample 1 from longissimus dorsi, Samples 2-3 from semimembranosus); c, Genotype patterns of the $LRRC2$ gene region between Mouflon sheep (n=17) and thin-tailed sheep (n=15).