Comparative transcriptome analysis by deep RNA sequencing at early stage of skin pigmentation in goats (*Capra hircus*)

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ABSTRACT

Although specific genes have been found to be associated with skin pigmentation, the global gene expression profile for the early stage of skin pigmentation and development in mammals is still not well understood. Here we reported a rare natural group of goat (Youzhou dark goat) featuring the dark skin of body including the visible mucous membranes, which may be an exclusive kind of large mammalian species with this special phenotype so far. In the present study, we characterized the 100-day-old fetal skin transcriptome in hyperpigmented (dark-skinned) and wild-type (white-skinned) goats using deep RNA-sequencing. A total of 923,013,870 raw reads from 6 libraries were obtained, and a large number of alternative splicing events were identified in the transcriptome of fetal skin, including the well-known melanogenic genes ASIP, TYRP1, and DCT, which were differentially expressed in the skin between the dark-skinned and white-skinned goats. Further analysis demonstrated that differential genes including ASIP, TYRP1, DCT, WNT2, RAB27A, FZD4, and CREB3L1 were significantly overrepresented in the melanogenesis pathway and several biological process associated with pigmentation. On the other hand, we identified 1616 novel transcripts in goat skin based on the characteristics of their expression level and gene composition. These novel transcripts may represent two distinct groups of nucleic acid molecules. Our findings contribute to the understanding of the characteristics of global gene expression at early stages of skin pigmentation and development, as well as describe an animal model for human diseases associated with pigmentation.

INTRODUCTION

Skin pigmentation is a complex process that includes melanin biosynthesis in melanocytes, which
is transferred to keratinocytes (Wolff 1973; Seiberg 2001). Unlike the numerous variations of
coat color, there are only a few natural variants of skin color in other mammals compared to the
phenotypes of human skin color. As a result, most studies on the genetics of skin pigmentation
are conducted in humans (Baxter and Pavan; Sturm 2009; Quillen and Shriver 2011; Meng et al.
2012) and mice (Bennett and Lamoreux 2003; Garcia et al. 2008), in which 378 color
genes (171 cloned genes and 207 uncloned genes) have been identified
(http://www.espcr.org/micemut) to date. However, only a small portion of these color genes have
been identified as candidate causative genes for skin pigmentation in different populations of
humans, mice/rats, and sheep, including ASIP, TYRP1, DCT, TYR, HERC2, OCA2, MC1R,
SLC24A5, SLC45A2, IRF4, KITLG (Slominski et al. 2004; Yamaguchi et al. 2007; Ebanks et al.
2009; Yamaguchi and Hearing 2009; Garcia-Gamez et al. 2011; Kondo and Hearing 2011; Liu et
al. 2013; Raadsma et al. 2013; Han et al. 2015). In addition, few studies have characterized the
global gene expression profile of skin pigmentation and development. Although investigations of
the skin transcriptome have recently been conducted on coat color (Fan et al. 2013) and hair
follicles (Xu et al. 2013a; Xu et al. 2013b; Wang et al. 2015; Yue et al. 2015; Gao et al. 2016) in
sheep and goats, and skin color in the common carp (Jiang and Bikle 2014b; Wang et al. 2014),
red tilapia (Zhu et al. 2016), and chickens (Zhang et al. 2015), the genetic basis for skin
pigmentation is not well understood compared to that of hair or coat color.

Here, we report a rare indigenous goat breed (Youzhou dark goat) that features dark skin of the
body including the visible mucous membranes (Fig.1A), which is a rare fibromelanosis that has
previously only been reported in the Silky fowl (Nozaki and Makita 1998; Muroya et al. 2000;
Dorshorst et al. 2011; Shinomiya et al. 2012). The pigmentation phenotype of the Youzhou dark
goat significantly differs from the piebald phenotype of the bovine (Weikard et al. 2013),
whereas it is similar to a recently reported case of dermal melanocytosis in humans (Lee et al. 2010). However, the underlying mechanism of this hyperpigmentation is yet to be explored. Therefore, the Youzhou dark goat can be used as a medical model to study human diseases associated with pigmentation, such as skin melanopathy, melanosis coli, and mucosal melanosis. Interestingly, based on our long-term observations (unpublished), skin pigmentation parallels skin development during the pregnant and postnatal period in the Youzhou dark goat. Consequently, to understand fibromelanosis in mammals, it is necessary to investigate the biology of skin pigmentation at early developmental stages in goats. In developmental embryology, the growth of the fetal skin peaks at approximately 100 days of gestation in sheep and goats (Wang et al. 1996; Qin 2001). In the present study, we used the Youzhou dark goat (hyperpigmented or dark-skinned) and Yudong white goat (wild-type or white-skinned) as models of skin pigmentation to characterize the skin transcriptome in 100-day-old fetal goats using deep RNA-sequencing. Our study not only contributes to the understanding of the biology of skin pigmentation and development but also provides valuable information towards understanding human melanocytosis.

MATERIALS AND METHODS

Ethical statement

All surgical procedures in goats were performed according to the Regulations for the Administration of Affairs Concerning Experimental Animals (Ministry of Science and Technology, China; revised in June 2004) and adhered to the Reporting Guidelines for Randomized Controlled Trials in Livestock and Food Safety (REFLECT).

Animals
Two goat groups with different skin pigmentation phenotypes were investigated in this study. The Yudong white goat (*Capra hircus*) is distributed in Southwest China (located at 31°14′-32°12′ N and 108°15′-109°58′ E) and features white color in the coat and skin. The Youzhou dark goat (*Capra hircus*) is an indigenous breed uniquely distributed in Youyang County in Chongqing, China (located at 26°54′ N and 108°57′ E) and features dark skin of the body including the visible mucous membranes but a generally white coat color. Briefly, three pregnant ewes from each breed were subjected to a caesarean section to collect the fetuses (n=3) at 100 days of gestation, and then the dorsal and ventral skins were collected from each fetus. The first sample (3 grams) was dissected and rapidly frozen whole in isopentane chilled over liquid nitrogen for histological examination. The second sample (3 grams) was snap-frozen in liquid nitrogen for RNA-sequencing and qPCR analysis.

**RNA isolation, library preparation and sequencing**

In the present study, a total of 6 libraries were generated for sequencing according to the sample size (n=6) in two breeds of goats. For each of the 6 fetal goats, total RNA was isolated using TRIzol reagent (Invitrogen, USA) according to the manufacturer’s instructions. RNA degradation and contamination was monitored on 1% agarose gels. RNA purity was checked using the NanoPhotometer spectrophotometer (Implen, USA). RNA concentration was measured using a Qubit RNA Assay Kit with a Qubit 2.0 Flurometer (Life Technologies, USA). RNA integrity was assessed using the RNA Nano 6000 Assay Kit with the Agilent 2100 Bioanalyzer System (Agilent Technologies, USA). A total of 3 μg of RNA per sample was used as the input material for the RNA sample preparations. First, ribosomal RNA was removed using the Epicentre Ribo-zero™ rRNA Removal Kit (Epicentre, USA), and rRNA free residue was cleaned up using ethanol precipitation. Subsequently, the highly strand-specific libraries were generated using the
rRNA-depleted RNA using the NEBNext Ultra™ Directional RNA Library Prep Kit for Illumina (NEB, USA) according to the manufacturer’s recommendations. Briefly, fragmentation was carried out using divalent cations under elevated temperature in NEBNext. First strand cDNA was synthesized using random hexamer primers and M-MuLV Reverse Transcriptase (RNaseH-).

Second strand cDNA synthesis was subsequently performed using DNA Polymerase I and RNase H. In the reaction buffer, dNTPs with dTTP were replaced by dUTP. The remaining overhangs were converted into blunt ends via exonuclease/polymerase activities. After adenylation of the 3’ ends of the DNA fragments, NEBNext Adaptor with hairpin loop structure were ligated to prepare for hybridization. To select cDNA fragments of preferentially 150-200 bp in length, the library fragments were purified using the AMPure XP system (Beckman Coulter, USA). Then, 3 μl of USER Enzyme (NEB, USA) was used with size-selected, adaptor-ligated cDNA at 37°C for 15 min followed by 5 min at 95°C before PCR. PCR was performed with Phusion High-Fidelity DNA polymerase, Universal PCR primers and Index (X) Primer. Finally, the products were purified (AMPure XP system) and the library quality was assessed on the Agilent 2100 Bioanalyzer System. The clustering of the index-coded samples was performed on a cBot Cluster Generation System using TruSeq PE Cluster Kit v3-cBot-HS (Illumina) according to the manufacturer’s instructions. After cluster generation, the libraries were sequenced on an Illumina Hiseq 2000 platform and 100 bp paired-end reads were generated.

**Quality control**

The raw data were firstly processed through in-house perl scripts. In this step, clean data were obtained by removing the reads containing adapters, reads containing over 10% of poly-N sequences, and low quality reads (more than 50% of bases with Phred scores less than 5%) from the raw data. The Phred score (Q20, Q30) and GC content of the clean data were calculated. All
the subsequent analyses were based on the high quality data. The sequencing data were submitted to the Genome Expression Omnibus (Accession Numbers GSE69812) in NCBI.

Mapping to the reference genome

The reference genome and gene model annotation files were downloaded directly from the genome website (http://goat.kiz.ac.cn). The index of the reference genome was built using Bowtie v2.0.6 (Langmead et al. 2009; Langmead and Salzberg 2012) and paired-end clean reads were aligned to the reference genome using TopHat v2.0.9 (Trapnell et al. 2012; Kim et al. 2013).

Transcriptome assembly

The goat reference genome and gene model annotation files were downloaded directly from the genome website (http://goat.kiz.ac.cn). The index of the reference genome was built using Bowtie v2.0.6 (Langmead et al. 2009; Langmead and Salzberg 2012) and paired-end clean reads were aligned to the reference genome using TopHat v2.0.9 (Trapnell et al. 2012; Kim et al. 2013). The mapped reads of each sample were assembled using both Scripture (beta2) (Guttman et al. 2010) and Cufflinks (v2.1.1) (Trapnell et al. 2010) in a reference-based approach. Scripture was run with default parameters. Cufflinks was run with ‘min-frags-per-transfrag=0’ and ‘--library-type fr-firststrand’, with all other parameters set as default.

Quantification of gene expression level

Cuffdiff (v2.1.1) was used to calculate the FPKM (fragments per kb for a million reads) of both lncRNAs and coding genes in each sample (Trapnell et al. 2010). For biological replicates (n=3), transcripts or genes with a $P$-adjust<0.05 were considered differentially expressed between the two groups of goats (dark-skinned and white-skinned).
Alternative splicing analysis

Alternative splicing (AS) events were classified into 12 basic types using the software Asprofile v1.0 (Florea et al. 2013). The number of AS events in each of the 6 samples was estimated separately. We used DEXSeq software (Anders et al. 2012) for the differential exon usage analysis of the AS transcripts, in which a general linear model was employed for the differential analysis of exon expression and a \( P \)-adjust < 0.05 indicated a significant result.

Identification of novel transcripts

To identify the novel transcripts from clean data, we first distinguished the novel transcripts and the candidate long noncoding RNAs using four tools including CNCI (v2) (Sun et al. 2013), CPC (0.9-r2) (Kong et al. 2007), Pfam-scan (v1.3) (Punta et al. 2012), and PhyloCSF (v20121028) (Lin et al. 2011). Transcripts predicted to be without coding potential by all of four tools above were filtered away (which are considered as the candidate long noncoding RNAs), and those with coding potential (which are selected by either of four tools above) but lacking of any known annotation were kept and considered as the novel transcripts in the present analysis. Quantification of gene expression level were estimated by calculating FPKMs of the novel transcripts.

RepeatMasker (http://www.repeatmasker.org/cgi-bin/WEB RepeatMasker) was used with the default parameters to identify various TE components in goat. To identify the position bias of TEs in the novel transcript, we searched the TEs in the 2,000 bp upstream of TSS (Transcription Start Site) of each transcript identified in the goat genome (http://goat.kiz.ac.cn) and plotted read coverage at TSSs with the ggplot2 package in R (Wickham 2009).

Differential expression and functional enrichment analysis
Cuffdiff provides statistical routines to determine differential expression in digital transcript or gene expression data using a model based on the negative binomial distribution (Trapnell et al. 2010). For biological replicates, transcripts or genes with a $P$-adjust<0.05 were considered differentially expressed. To identify the molecular events or cascades involved, the differentially expressed genes or IncRNA target genes were analyzed using the DAVID platform (Huang Da et al. 2009b; Huang Da et al. 2009a). Significance was expressed as a $P$-value, which was calculated using the EASE score (a $P$-value of 0.05 was considered significant).

**Validation of gene expression in RNA-seq using quantitative PCR**

To validate the gene expression in RNA-seq, the total RNA from the RNA-seq analysis was used for qPCR (Fig. 4). Briefly, first strain cDNA was obtained using a One Step cDNA Synthesis Kit (Bio-Rad, USA), and the mRNA was then quantified using a standard SYBR with GAPDH as an endogenous control. Quantitative PCR was performed under the following conditions: 95°C for 30 sec, 40 cycles of 95°C for 5 sec, and the optimized annealing temperature for 30 sec. The primers and annealing temperatures for the 8 genes are listed in Table S1. All reactions were performed in triplicate for each sample. Gene expression was quantified relative to GAPDH expression using the $2^{(-\Delta\Delta C_t)}$ method. Corrections for multiple comparisons were performed using the Holm-Sidak method.

**Statistical analysis**

Data analyses were performed using the R statistical package.

**Data availability**

Figure S1 contains classification of the raw reads by RNA-seq for each of the libraries (samples) in the hyperpigmented and normal fetal goats. Figure S2 contains a heatmap of the cluster
analysis for the differentially expressed genes. Table S1 contains statistics of the mapping reads to the reference genome. Table S2 contains annotation of novel transcripts in Swiss-Prot database. Table S3 contains differential exon usage (DEU) analysis of the mRNAs in goat fetal skin. Table S4 contains the differentially expressed genes between the wild-type and hyperpigmented goat skins. Table S5 contains results of GO and KEGG analysis of the up-regulated DE genes using DAVID. Table S6 contains results of functional annotation clustering of the DE genes using DAVID.

RESULTS

RNA sequencing of skin in fetal goats

As for phenotype, there are obvious differences in skin tissue pigmentation between the two breeds of fetal goat (Fig. 1A and B). Then we examined the phenotypic differences at the histological level using H & E staining. The results demonstrated that there were more melanin granules in the 100-day fetal epidermal and dermal layers in the Youzhou dark goat than in the Yudong white goat (Fig. 1C and D). To further explore the mechanism underpinning the differences in skin pigmentation, we characterized the skin transcriptome using RNA-seq performed with the Illumina HiSeq 2000 platform. In the present study, we obtained a total of 923,013,870 raw reads from 6 libraries (samples), of which 841,895,634 clean reads remained for further analysis after discarding the raw reads that contained adapter sequences, N sequences and low quality sequences. The percentage of clean reads among the raw tags in each library ranged from 88.39% to 93.02% (Fig. S1). We then mapped the clean reads to the goat reference genome
Of the total reads from each library, more than 83% matched to a unique genomic location, whereas only 3% matched to multiple genomic locations (Table S1). The Phred score (Q20) for each library was greater than 95.4%, which indicates our RNA-seq data were high quality and suitable for the subsequent analyses.

**Novel transcripts in goat genome**

We identified a total of 27,947 mRNAs and 1616 novel transcripts in fetal goat skin in the present study. To examine their homology with proteins in state-of-the-art database, we made a BLAST alignment for these novel transcripts in Swiss-Prot database (http://www.uniprot.org/). Results showed that 660 of them had a similarity in various degree, but with a relative lower coverage to the known proteins (Table S2). Another 956 novel transcripts were found without annotation in the state-of-the-art database presently. Then we examined the differences in gene length and expression levels (FPKM) between the two groups of novel transcripts, and found that there were significant differences in gene length (Kolmogorov-Smirnov test, \( P = 0.011 \); Fig 2A) and expression levels (Kolmogorov-Smirnov test, \( P = 0.000 \); Fig 2B) between them. To further characterize the two groups of novel transcripts, we examined the composition of transposable elements (TEs) harboring in their gene sequences. Our results demonstrated that there were considerable differences in density of various TE components between the two groups of novel transcripts, as well as between the novel transcripts and mRNAs (Fig 2C). Further investigation of position bias for TEs relative to TSS (transcription start site) revealed that the LINE/RTE-BovB were higher enriched in novel transcripts without annotation than the other two groups (Fig 2D). These above findings suggest that the identified transcripts may represent different types of novel transcripts in goat genome.

**Alternative splicing of the fetal skin transcriptome**
To ascertain the alternative splicing (AS) events of the skin transcriptome, we examined the data from six samples using ASprofile software. Our results showed that TSSs (alternative 5' first exon, transcription start sites), TTSs (alternative 3' last exon, transcription terminal sites), and SKIPS (skipped exons) were the three most frequently observed AS events among the 12 AS subtypes in each sample (Fig. 3A). Among the over 66,000 AS events, we observed AS events in two well-known genes involved in pigmentation, \textit{ASIP} and \textit{TYRP1}, which were found in both goat breeds. To further examine the AS events of the differentially expressed genes between the dark- and white-skinned goat breeds, we subsequently investigated differential exon usage of the AS events using DEXSeq software. The results demonstrated that 360 AS exons belonging to 253 known and 28 unknown genes were significantly differentially expressed between the dark- and white-skinned groups ($P$-adjust$<0.05$) (Table S3). Importantly, we found that some AS exons belonging to well-known genes involved in pigmentation, such as \textit{ASIP}, \textit{DCT}, and \textit{RAB27A}, were significantly differentially expressed between the two different skin colors (Fig. 3B). This indicates that alternative splicing may be an important mechanism by which expression and function of \textit{ASIP}, \textit{DCT}, and \textit{RAB27A} is regulated in melanocytes.

**Identification and functional clustering analysis of the differentially expressed genes**

In our data, we obtained 448 differentially expressed (DE) genes ($P$-adjusted$<0.05$ and fold change$\geq1.5$) using Cuffdiff between the dark- and white-skinned goats. Of the 448 DE genes, 101 genes were up-regulated and 347 were down-regulated in the dark-skinned goats compared with the white-skinned goats (Fig. 4A, Table S4, Fig. S2). Here, to elucidate the biology of skin pigmentation in goats, we specially focused on the up-regulated DE genes in the dark-skinned...
goats. Gene functional classification using DAVID demonstrated that a family of keratin (KRT) was significantly enriched in the up-regulated DE genes, including KRT-1, KRT81, KRT83, KRTAP13-1, KRTAP11-1, and KRTAP7-1, which are constitutive components of skin produced by keratinocytes. Gene ontology analysis showed that the top six biological processes were response to inorganic substance, response to extracellular stimulus, response to reactive oxygen species, pigmentation during development, pigment biosynthetic process, and response to cAMP (Table S5). KEGG analysis showed that genes involved in melanogenesis and the MAPK signaling pathway were significantly enriched. These results suggest that the up-regulated genes play important roles in the dark skin phenotype of the Youzhou dark goats.

To gain a comprehensive insight of the gene expression between the two breeds, we performed Gene ontology and pathway analyses of all DE genes between the normal and dark-skinned goats using DAVID. Functional clustering analysis of all DE genes showed that 65 annotation clusters were significantly enriched, particularly cluster 41, which comprises 7 terms related to hyperpigmentation including pigment biosynthetic process, melanin biosynthetic process, pigment metabolic process, melanin metabolic process, secondary metabolic process, melanogenesis, and pigmentation during development. Regarding the pathways involved, we identified 14 significantly overrepresented pathways (Table S6) including the melanogenesis pathway (Fig. 4B). Selected DE genes from the RNA-seq analysis were validated using quantitative PCR (Fig. 5). These findings provide additional evidence of the underlying diversity of dermal colors in goats on the transcriptome level. Moreover, we explored the enrichment of genes involved in human pigmentation diseases using DAVID to determine whether there are diseases with similar variations in dermal colors in goats as are observed in humans. As we expected, ASIP and TYRP1 were significantly enriched in one type of OMIM disease (Table 1).
This suggests that ASIP and TYRP1 are the two likely candidate genes responsible for dermal hyperpigmentation in Youzhou dark goats.

**DISCUSSION**

Although transcriptomic investigations on variations in skin color have been conducted in the common carp (Jiang and Bikle 2014b; Wang et al. 2014), red tilapia (Zhu et al. 2016), and chickens (Zhang et al. 2015), few studies have investigated this topic in mammals to date. We first characterized the skin transcriptome in two breeds of fetal goats that feature different skin colors using deep RNA-seq methods. The present study produced a large amount of data (over 84.2 Gb of clean reads for six samples) and more than 86% of the total reads were mapped to the goat reference genome (http://goat.kiz.ac.cn), which indicated that the RNA-seq data were of high quality and valuable for further analysis. First, we identified a significant number of AS events (Fig. 3A) in the skin transcriptome of fetal goats, which suggests that gene transcription and regulation is complex in the skin during this developmental stage in goats. AS events are tissue specific and widespread in the genome (Xu et al. 2002). AS produces variety in the proteins translated from a limited number of genes through significant alterations in protein conformations that modulate cell functions (Yura et al. 2006). Our findings (Fig. 3; Table S3) indicate that alternative splicing may be an important mechanism by which the function of ASIP, TYRP1, DCT, and RAB27A are regulated in melanocytes. Therefore, it is necessary to identify the isoforms of these genes and their roles in pigmentation in the future. Interestingly, in the present study, we did not observe alternative splicing of the MITF gene in the skin, whereas multiple splice variants of MITF were previously found in other species such as sheep (Saravanaperumal et al. 2014), mice (Bismuth et al. 2005), and humans (Kuiper et al. 2004). There might be
differences in the regulatory mechanisms of skin pigmentation between goats and the species mentioned above. Another valuable finding of the present study is the DE genes identified in the fetal skin between the two goat groups. In particular, members of the keratin family were highly differentially expressed in the fetal skin between the two goat groups, including \textit{KRT1}, \textit{KRT81}, \textit{KRT83}, \textit{KRTAP13-1}, \textit{KRTAP11-1}, and \textit{KRTAP7-1} (Table S5). Keratins make up the largest subgroup of intermediate filament proteins and represent the most abundant proteins in epithelial cells. These proteins are essential to sustain normal epidermal function and play a role in signaling (Porter and Lane 2003). Mutations of a member of the KRT family resulted in an unusual skin pigmentation in humans (Irvine \textit{et al.} 1997; Horiguchi \textit{et al.} 2005; Pascucci \textit{et al.} 2006; Geller \textit{et al.} 2013). Therefore, these DE keratins might participate in the signaling transduction involved in melanin synthesis in melanocytes or the transportation of melanin from melanocytes to keratinocytes through cellular interactions (Nakazawa \textit{et al.} 1995; Seiberg 2001; Joshi \textit{et al.} 2007).

Further functional clustering analysis of the DE genes showed that the terms involved in pigmentation and melanogenesis were highly overrepresented in our findings (Table S5 and S6), which is similar to the findings from studies on coat color in sheep (Fan \textit{et al.} 2013) and skin color in the common carp (Jiang and Bikle 2014a). However, the specific DE genes of the skin differed in this study compared with the studies above. Specifically, the melanogenic \textit{ASIP} (fold change $>$31) and \textit{TYRP1} (fold change $>$87) were not only among the most significantly DE genes between the dark- and white-skinned goats, but were also associated with human pigmentation (Table 1). Previous studies have demonstrated that polymorphisms of \textit{ASIP} are associated with darker skin color in African Americans (Bonilla \textit{et al.} 2005) and fair skin color in Caucasians (Nan \textit{et al.} 2009). \textit{TYRP1}, a key regulator of melanin biosynthesis in melanocytes, is reported to
be involved in the diversity of skin color in Europeans (Lao et al. 2007) and is a candidate gene for skin color (such as mouth, nose, and ear) in sheep (Raadsma et al. 2013). In addition, DCT is also associated with variations of skin pigmentation in Asians (Myles et al. 2007). However, the results of the present study suggest that ASIP is more likely the candidate gene for skin color in goats than TYRP1 and DCT. We believe that TYRP1 and DCT are two downstream genes whose expression levels are affected by ASIP in the KEGG pathway melanogenesis (Fig. 4B). However, efforts still be taken to ascertain the candidate genes underlying variations of skin color in goats using gene mapping techniques such as genome wide association studies (GWAS), next generation sequencing (NGS), or quantitative trait loci (QTL) analyses.

Our another valuable finding is the novel transcripts identified by RNA-seq (Table S1). Cabili, et al (2011) characterized a class of novel transcripts that were excluded by their coding potential criteria (a Pfam domain, a positive PhyloCSF score, or previously annotated as pseudogenes), and firstly termed them as TUCP (transcripts of uncertain coding potential) (Cabili et al. 2011). Since they merely focused the lincRNAs in human genome under a certain classification strategy, other subtypes of IncRNAs such as intronic IncRNAs and antisense IncRNAs were retained and thus grouped into the catalog of TUCP in their findings. However, in the present study, the intronic IncRNAs and antisense IncRNAs were absolutely excluded from the novel transcripts based on our recent study (Ren et al. 2016). Thus there are certain differences in classification of novel transcripts between the study from Cabili et al and ours. In view of subsequent analyses, we are certain that the identified transcripts with annotation and without annotation are two different groups of novel transcripts. Especially for those without annotation, the characteristics of a relatively low expression level and a highly enrichment of TE component (LINE/RTE-BovB) around TSS (transcription start site) (Fig 2) which are similar with that of the lincRNAs in recent
studies (Cabili et al. 2011; Derrien et al. 2012; Kelley and Rinn 2012; Li et al. 2012; Nam and Bartel 2012; Billerey et al. 2014; Ren et al. 2016), strongly suggest this group of novel transcripts could be long nocoding RNAs. However, we are not certain that the novel transcripts with some annotation should be protein coding genes or long nocoding RNAs based on the current knowledge. It still need further efforts to identify their identity. Generally, this study provides a valuable resource for the genetic mechanisms involved in pigmentation diseases and contributes to the understanding of the biology of skin pigmentation and development in mammals.

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Figure legends

Fig. 1 Histomorphological examination of the hyperpigmented and normal skin in fetal goats.

To explore the histomorphological differences between the two breeds of goats at the early stage of skin pigmentation and development, the 100-day-old fetal skin from the Youzhou dark goat (A
and C) and Yudong white goat (B and D) were examined using H & E staining. The scale bar for the images in (C) and (D) is 10 μm.

**Fig. 2** Characteristic differences between two groups of novel transcripts in goat genome

The transcript length (A) and expression level (B) of two classes of novel transcripts were compared using the Kolmogorov-Smirnov test, and a P value of 0.05 indicates significance between two groups. In two box plots, the circle indicates the outlier, and the asterisk labels the extreme. (C) The proportion of the main TE families in two groups of novel transcripts and mRNAs in the goat genome (http://goat.kiz.ac.cn). Differences in TE components between them were measured by using the Fisher Exact test. (D) The position bias of TE components in the 2,000 bp upstream of TSS in transcripts above mentioned.

**Fig. 3** Alternative splicing (AS) events in the goat genome.

Following the analysis using Cufflinks, the classification of AS events of transcripts were performed for each library (sample) using the ASprofile tool. The 12 main AS events (TSS, TTS, SKIP, XSKIP, MSKIP, XMSKIP, IR, XIR, MIR, XMIR, AE, and XAE) in the goat genome are summarized in (A). Y1, Y2, and Y3 indicate the dark skin goats, whereas B1, B2, and B3 indicate the white skin goats. (B) Differential exon usage of the AS events in the ASIP and DCT genes between the dark-skinned (Y, blue) and white-skinned (B, red) goats.

**Fig. 4** The differentially expressed genes between the dark- and white-skinned goats.

(A) The differential expression of genes was identified based on the FPKM values (corrected P-value≤0.05 and fold change≥1.5) in the two groups. The volcano plot was generated based on the values above. Y indicates the dark-skinned goats, and B refers to the white-skinned goats. (B) Enrichment of the DE genes was conducted under KEGG pathways. Red indicates the
up-regulated DE genes and green refers to the down-regulated genes in the dark-skinned goats compared to the control goats (white-skinned).

**Fig. 5 Validation of gene expression in the dark- and white-skinned goats using qPCR.**

Some of the identified melanogenic genes were examined in the dark- and white-skinned fetal goats using qPCR. Gene expression was quantified relative to \textit{GAPDH} expression using the $2^{(-\Delta C_t)}$ method. Corrections for multiple comparisons were performed using the Holm-Sidak method. The data are shown as the mean ± 1 SE (n = 3). *$P<0.05$, **$P<0.01$.

**Additional files**

- Fig. S1. Classification of the raw reads by RNA-seq for each of the libraries (samples) in the hyperpigmented and normal fetal goats.
- Fig. S2. Heatmap of the cluster analysis for the differentially expressed genes.
- Table S1. Statistics of the mapping reads to the reference genome.
- Table S2. Annotation of novel transcripts in Swiss-Prot database.
- Table S3. Differential exon usage (DEU) analysis of the mRNAs in goat fetal skin.
- Table S4. Differentially expressed genes between the wild-type and hyperpigmented goat skins.
- Table S5. GO and KEGG analysis of the up-regulated DE genes using DAVID.
- Table S6. Functional annotation clustering of the DE genes using DAVID.
Statistics of AS events

- XSKIP
- XMSKIP
- XMIR
- XIR
- XAE
- TTS
- TSS
- SKIP
- MSKIP
- MIR
- IR
- AE

AS category

Number of AS

Y1

Y2

Y3

B1

B2

B3

0 5000 10000 15000 20000 25000
The image shows a volcano plot with the title "Y_vs_B". The plot displays the negative log of the q-value on the y-axis and the log2 of the fold change on the x-axis. The plot includes red and green points indicating upregulated and downregulated genes, respectively. The number of upregulated genes is 101, and the number of downregulated genes is 347.