

Research Article

Identification of Differentially Expressed Genes in Different Types of Broiler Skeletal Muscle Fibers Using the RNA-seq Technique

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The difference in muscle fiber types is very important to the muscle development and meat quality of broilers. At present, the molecular regulation mechanisms of skeletal muscle fiber-type transformation in broilers are still unclear. In this study, differentially expressed genes between breast and leg muscles in broilers were analyzed using RNA-seq. A total of 767 DEGs were identified. Compared with leg muscle, there were 429 upregulated genes and 338 downregulated genes in breast muscle. Gene Ontology (GO) enrichment indicated that these DEGs were mainly involved in cellular processes, single organism processes, cells, and cellular components, as well as binding and catalytic activity. KEGG analysis shows that a total of 230 DEGs were mapped to 126 KEGG pathways and significantly enriched in the four pathways of glycolysis/gluconeogenesis, starch and sucrose metabolism, insulin signalling pathways, and the biosynthesis of amino acids. Quantitative real-time reverse transcription polymerase chain reaction (qRT-PCR) was used to verify the differential expression of 7 selected DEGs, and the results were consistent with RNA-seq data. In addition, the expression profile of *MyHC* isoforms in chicken skeletal muscle cells showed that with the extension of differentiation time, the expression of fast fiber subunits (types IIA and IIB) gradually increased, while slow muscle fiber subunits (type I) showed a downward trend after 4 days of differentiation. The differential genes screened in this study will provide some new ideas for further understanding the molecular mechanism of skeletal muscle fiber transformation in broilers.

1. Introduction

Chicken is widely welcomed by consumers because it is low in fat, is low in cholesterol, and has no religious restrictions. Therefore, chicken has become the second-largest consumed meat product after pork [1, 2]. In recent years, with the continuous improvement in people's living standards, their requirements for meat flavor have also become stricter. Age, sex, heredity, environment, nutrition, and intramuscular fat content (IMF) are all important factors affecting chicken quality [3–6]. In addition, studies have shown that muscle fiber properties were an essential factor affecting the meat quality of chicken [7, 8]. Skeletal muscle is one of the most important components of meat-producing animals,

accounting for approximately 45%–60% of the whole animal body [9]. According to the characteristics of contraction and metabolism, skeletal muscle fibers can be roughly divided into two types: slow-twitch (type I) and fast-twitch (type II) muscles. Type II muscle fibers can be categorized into a fast-twitch oxidation type (IIA), a fast-twitch glycolytic type (IIB), and a super-fast-twitch type (IIX) [10, 11]. Previous studies showed that muscle with a high proportion of fast-twitch muscle fibers appeared paler, while the muscle with more slow-twitch muscle fibers tended to be redder [8]. Therefore, skeletal muscle was usually divided into two kinds of muscle, red muscle and white muscle [12]. Moreover, it was found that muscles with a higher proportion of oxidized muscle fibers had better meat quality [13].

Currently, it has become a research hotspot to study the molecular mechanism of muscle fiber-type transformation in order to improve meat quality. It has been shown that *PGC-1* can promote the transformation of muscle into type I fibers in the skeletal muscle [14]. Cofilin2b (*CFL2b*) can affect the transformation pattern of myosin heavy chain (*MyHC*) subunits in the muscle of piglets [15]. Furthermore, noncoding RNA was also proven to play a role in slow-twitch muscle fiber formation and skeletal myogenesis [16]. At present, preliminary progress has been made on the molecular mechanism of chicken muscle fiber-type transformation. One study showed that the expression of *PGC-1 α* might be closely related to the slow muscle fiber content in chicken skeletal muscle [17]. In addition, miR-1611 was discovered to be highly expressed in slow-twitch muscle fibers and could drive the transformation from fast-twitch to slow-twitch muscle fibers in chicken [18]. However, the regulatory network of muscle development is complex, and studies on the formation and transformation of chicken skeletal muscle fiber types are still incomplete. Therefore, more relevant, important genes need to be further explored.

At present, transcriptome sequencing technology has been widely used in biological research, medical diagnostics, and therapeutic studies [19]. Meanwhile, this technique also plays an important role in farm animal muscle development. Based on the transcriptome analysis of biceps between Small-tail Han sheep and Duper sheep, a total of 1300 differentially expressed genes were identified, which could help further elucidate the mechanisms of muscle development as they are affected by breed [20]. Forty-nine differentially expressed genes were screened from the longissimus dorsi of Nelore cattle among different grades of marbling by RNA-seq, in an effort to find strategies to select animals with greater marbling [21]. Moreover, another study used RNA-seq to identify genes related to pig muscle development and meat quality [22]. RNA-seq technology was also used in broiler chicken groups to uncover 68 genes that might participate in the tenderization process [23]. Thus, it can be seen that whole-genome sequencing technology exerts an immense influence on the discovery of new functional genes.

To better explore the molecular mechanism of skeletal muscle fiber transformation in broilers, this study used RNA-seq technology to screen differentially expressed genes between breast and leg muscles of Ross 308 broiler chickens. Subsequently, through GO and KEGG pathway analysis, genes and pathways related to skeletal muscle fiber-type conversion were discovered. In addition, to further explore the growth characteristics of broiler muscle fibers, a skeletal muscle cell differentiation model of broilers was constructed *in vitro*. Subsequently, the expression of *MyHC* subunits during the differentiation of chicken skeletal muscle satellite cells was studied. These results will provide a theoretical basis for the genetic improvement in muscle quality in broilers in the future.

2. Materials and Methods

2.1. Experimental Animals and Tissues. The chickens used in this study were Ross 308 broiler chickens from a farm in Fuyang District, Hangzhou, Zhejiang Province, China. Five

42-day-old Ross 308 broilers with similar body weights were selected for slaughter. The leg and breast muscles were collected and frozen at -80°C for RNA extraction.

At the same time, we bought 15 fertilized eggs of the same breed and incubated them for 10 days to collect chicken skeletal muscle satellite cells. All animal procedures used in this study were approved by the Ethics Committee for Animal Experiments of Zhejiang A&F University and were performed in accordance with the Guidelines for Animal Experimentation of University (Hangzhou, China).

2.2. Total RNA Extraction, Library Construction, and Sequencing. Total RNA was extracted from the gastrocnemius of legs and the pectoralis major of breasts using the TRIzol method [24]. The quality inspection was carried out through the Sample Testing Center of Biomarker Technologies Corporation (Beijing, China) to ensure that the samples were up to the standard, and the test quality report of the RNA samples is presented in Table S1. All RNA samples with an OD 260/280 1.8-2.0 qualified for further analysis. Sequencing libraries were generated using a NEBNext Ultra™ RNA Library Prep Kit (NEB, USA). Briefly, mRNA was purified from 1 μg of total RNA per sample using poly-T oligoattached magnetic beads. Then, first-strand cDNA was synthesized using a random hexamer primer and M-MuLV Reverse Transcriptase. Second-strand cDNA synthesis was subsequently performed using DNA polymerase I and RNase H. The cDNA fragments of 240 bp were preferentially selected with an AMPure XP system (Beckman Coulter, USA) for cDNA adaptor ligating. At last, polymerase chain reaction (PCR) amplification was carried out and the products were purified (AMPure XP system). The library quality was assessed on the Agilent Bioanalyzer 2100 system. The libraries were sequenced by paired-end (PE 150 bp) sequencing on an Illumina HiSeq™ platform (Illumina, USA), and the raw reads were obtained. The leg muscle and breast muscle tissues of broilers were numbered PT1-PT5 and PX1-PX5, respectively, for subsequent analysis.

2.3. Sequencing Quality Assessment and DEG Screening. Clean data (clean reads) was obtained by removing reads containing adapters or poly-N; low-quality reads were also removed from the raw data. At the same time, Q20, Q30, GC-content, and sequence duplication levels of the clean data were calculated. Ultimately, all downstream analyses were based on high-quality clean data. The chicken genome sequence (GRCg6a) was downloaded from the NCBI database (<https://www.ncbi.nlm.nih.gov/genome/?term=chicken>). Then, Hisat2 software was used to map these clean reads to the chicken reference genome [25]. Transcript assembly was performed using StringTie software [26]. StringTie is an algorithm based on optimization theory, which uses comparison information to construct a multivariable clipping graph and traffic network to assemble and evaluate the expression of reads according to the maximum flow algorithm. Compared with other software such as Cufflinks, it can build a more complete transcript and better evaluate expression [26]. Quantification of gene expression levels was estimated by the fragments per

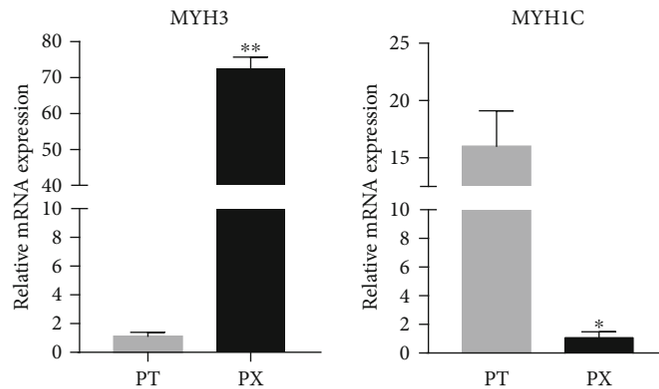


FIGURE 1: Expression of *MyHC* genes in different muscle tissues of broilers. Fast muscle marker gene *MYH3* and slow muscle marker gene *MYH1C*. **An extreme significant difference between the two groups ($P < 0.01$); *a significant difference between the two groups ($P < 0.05$); PT represents leg muscle; PX represents breast muscle, $n = 5$.

kilobase of transcript per million fragments (FPKM) method [27]. Based on raw counts, the reads mapped to the reference genome were quantified by StringTie software to obtain FPKM [28]. Then, DESeq R package (1.39.0), a statistical program for determining differential expression in digital gene expression data based on a model of negative binomial distribution, was used to analyze the differential gene expression between samples with biological repetition [29]. Differentially expressed genes (DEGs) were identified using the following filter criteria: fold change ≥ 2 and FDR < 0.05 . Fold change indicates the ratio of expression quantity between two samples (groups). The false discovery rate (FDR) was obtained by correcting the significant difference P value.

2.4. Gene Ontology (GO) and KEGG Pathway Enrichment Analysis. Gene Ontology (GO) enrichment analysis of the DEGs was implemented by Goseq R packages (<http://www.geneontology.org/>) based on Wallenius noncentral hypergeometric distribution [30], which could adjust for gene length bias in DEGs. Pathway significant enrichment analysis was run based on the hypergeometric test in the KEGG database (<http://www.genome.jp/kegg/>) to uncover the most significantly enriched pathways among the DEGs. KOBAS software [31] was used to test the statistical enrichment of DEGs in KEGG pathways.

2.5. Quantitative Real-Time RT-PCR Verification. Utilizing the NCBI gene bank, primers of 7 randomly selected DEGs, three different isoforms of *MyHC* genes (*MyHC* I, IIA, and IIB) and the house-keeping gene *GAPDH* were designed by Primer Premier 5 software (Table S1). Primers were synthesized by Hangzhou Youkang Biotechnology Co., Ltd. (Zhejiang, China). Total RNA from cells or tissues was used to synthesize cDNA by a 5X All-In-One MasterMix Kit (abm, China). The cDNA was used as a template for quantitative PCR by using the EvaGreen 2X qPCR MasterMix Kit (abm, China). There were three replicates for each sample. The relative expression level of each related gene was calculated using the $2^{-\Delta\Delta CT}$ statistical analysis method.

TABLE 1: Quality assessment of sequencing data.

Sample name	Clean reads	Clean bases	GC (%)	Q20 (%)	Q30 (%)
PT1	29135911	8705716622	51.03	97.65	93.63
PT2	32518744	9699863244	51.19	97.74	93.87
PT3	32658211	9752147744	51.27	97.73	93.85
PT4	37753032	11278877720	52.40	97.92	94.24
PT5	34146017	10159903756	52.05	97.86	94.13
PX1	25454197	7590698526	51.36	97.11	92.33
PX2	33709265	10026742412	51.19	97.61	93.31
PX3	29558343	8794379722	51.21	97.29	92.71
PX4	25035478	7478569054	51.89	97.64	93.46
PX5	29461523	8807133318	51.89	97.57	93.37

2.6. Isolation and Culture of Chicken Skeletal Muscle Satellite Cells In Vitro. The 10-day fertilized eggs were disinfected with ethanol. The eggshell on the upper part of the air chamber was removed using tweezers on the egg receptacle. A new pair of tweezers was used to clip out the chicken embryo. The leg muscle of the chicken embryo was isolated and placed in a new petri dish. Muscle samples were washed 3 times in phosphate buffer solution (PBS) (Logan, USA) containing penicillin-streptomycin liquid (Solarbio, China), and skin, blood vessels, adipose tissue, and connective tissue were removed. The sterile muscle tissue was cut into meat paste and digested with 0.25% trypsin-EDTA (Thermo Fisher, China) at 37°C for 20 min. DMEM/F12 (Logan, USA) containing 10% fetal bovine serum (FBS, Thermo Fisher, China) was added to terminate the digestion. The suspension was filtered through a 70 μm mesh sieve and centrifuged at 1200 r/min for 8 min at room temperature. As the supernatant was discarded, cells were resuspended with DMEM/F12 containing 15% FBS and cultured in a 5% CO₂ incubator at 37°C. One hour later, the cell suspension was transferred into a new cell petri dish; this was repeated twice to enrich for muscle satellite cells and eliminate fibroblasts. Cell differentiation was induced by replacing 15% FBS with 2% equine serum (Logan, USA). Once the isolated skeletal muscle

TABLE 2: Statistical results of the comparison with the reference genome.

Sample name	Total reads	Mapped reads	Mapped ratio (%)	Uniquely map	Uniquely mapped ratio (%)	Reads mapped to "+" strand	Reads mapped to "+" strand ratio (%)	Reads mapped to "-" strand	Reads mapped to "-" strand ratio (%)
PT1	58271822	51290093	88.02%	48720157	83.61%	25470179	43.71%	25437454	43.65%
PT2	65037488	55445945	85.25%	52738564	81.09%	27563636	42.38%	27476279	43.65%
PT3	65316422	56957569	87.2%	54089388	82.81%	28293609	43.32%	28228884	43.22%
PT4	75506064	64269152	85.12%	61021292	80.82%	31904145	42.25%	31888064	42.23%
PT5	68292034	53465050	78.29%	50826310	74.42%	26552333	42.25%	26493568	38.79%
PX1	50908394	43024726	84.51%	40845785	80.23%	21424253	42.08%	21316946	41.87%
PX2	67418530	58984363	87.49%	55702668	82.62%	29322172	43.49%	29258327	43.40%
PX3	59116686	51374316	86.90%	48593438	82.20%	25530131	43.19%	25474898	43.09%
PX4	50070956	42965914	85.81%	40786889	81.46%	21351653	42.64%	21342061	42.62%
PX5	58923046	51046402	86.63%	48489963	82.29%	25375162	43.06%	25318363	42.97%

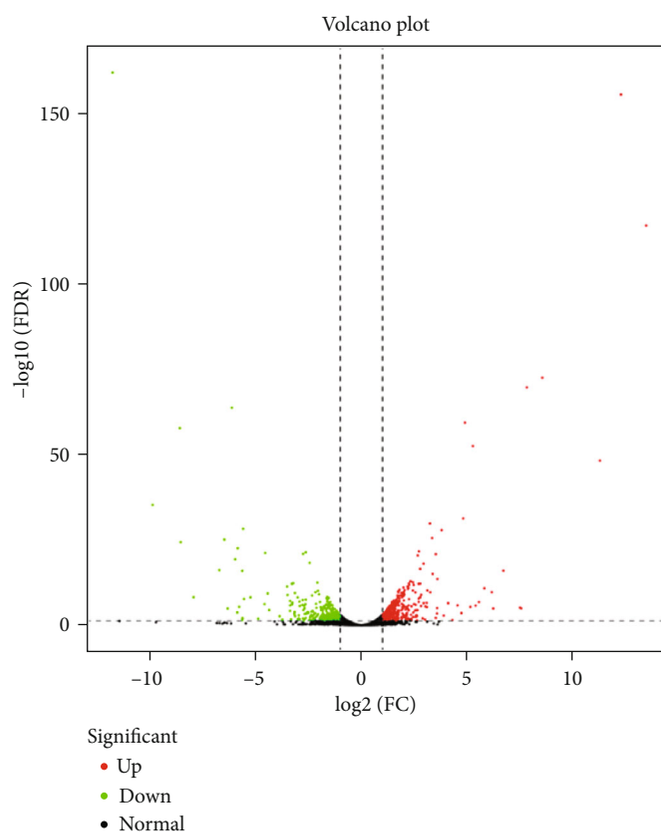


FIGURE 2: Volcano plot for differential gene expression.

satellite cells reached 70%-80% fusion, the growth medium was replaced with differentiation medium and the cells were cultured for 2, 4, and 6 days. Cells at 70%-80% fusion indicated day 0.

2.7. Immunofluorescence Staining. The isolated skeletal muscle satellite cells at 70%-80% fusion and the cells that were induced to differentiate after transfection were used for immunofluorescence detection following the method of Luo

et al. [32]. Briefly, cells grown in 12-well plates were washed three times with precooled PBS and fixed with 4% paraformaldehyde for 15 min. Thereafter, the cells were permeabilized with 0.25% Triton X-100 per well for 10 min and blocked at 4°C overnight. Afterwards, cells were incubated with 1:100 diluted primary anti-*Desmin* (Abcam, China), anti-*Pax7* (Abcam, China), or anti-*MyHC* (Santa, USA) antibodies for 1 hour at room temperature. Then, 1:100 diluted fluorescent secondary antibody (Thermo Fisher, China) was

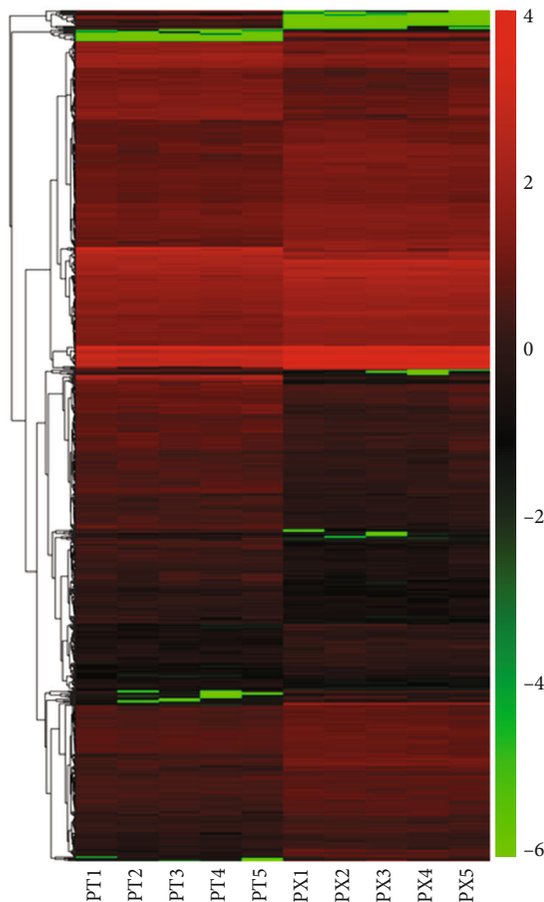


FIGURE 3: The heat map of differential gene expression (PT represents broiler leg muscle; PX represents breast muscle).

incubated with the cells for 1 hour at room temperature. After being washed by PBS three times, DAPI (Invitrogen, USA) was added to the cells and then incubated for 15 min at room temperature to stain the cell nuclei. Lastly, samples were captured using a fluorescence microscope (OLYMPUS, Japan).

3. Results

3.1. Expression of MyHC Genes in Different Muscle Tissues of Broilers. In some previous studies, compared with histochemical methods, qRT-PCR was a fast and sensitive approach to distinguish different muscle types by identifying the expression of *MyHC* subunits [33, 34]. This method was also applied to identify the muscle types of chickens [35, 36]. Therefore, in order to compare the fiber types of different muscles in broilers, the expression of *MYH3*, a fast muscle fiber marker gene [37], and *MYH1C*, a slow muscle fiber marker gene [38], were detected by qRT-PCR between breast and leg muscles of 10 randomly selected broilers. The results showed that the expression of *MYH3* in the breast muscle was significantly higher than that in the leg muscle, while the expression of *MYH1C* showed the opposite result (Figure 1). The results indicated that the breast and leg muscles of broilers were two different types of skeletal

muscle. The former was mainly composed of slow muscle fibers, while the latter was mainly composed of fast muscle fibers.

3.2. Sequencing Quality Evaluation. To verify the authenticity of the sequencing data, samples were tested for quality. In Table 1, it could be found that the Q20 (a base quality > 20 and error rate < 0.01) and Q30 (a base quality > 30 and error rate < 0.001) levels were both above 90%. Meanwhile, the GC content was between 49% and 53%. The results showed that the sequencing data had a small probability of error.

It was also found that more than 75% of reads aligned to the reference genome and that the number of reads aligned to a unique location of the reference genome was between 74% and 92%, which indicated that these samples were highly available (Table 2). To summarize, the sequencing data were good and the subsequent analysis could be carried out.

3.3. Screening of DEGs. To show the distribution of gene expression, a volcano plot of the differentially expressed genes was constructed, with $-\log_{10}(\text{FDR})$ as the ordinate and $\log_2(\text{FC})$ as the abscissa. Because the differential expression analysis of transcriptome sequencing was an independent statistical hypothesis test for a large number of gene expression values, there was a false-positive problem. Therefore, in the process of differential expression analysis, the recognized Benjamini-Hochberg correction method was used to correct the significant *P* values obtained by the original hypothesis test. Finally, FDR was used as the key index for differentially expressed gene screening. In our study, $\text{FDR} < 0.05$ was used as a screening condition for DEGs. The results showed that there were a large number of differentially expressed genes. A total of 767 DEGs were identified. Compared with leg muscle, there were 429 upregulated genes and 338 downregulated genes in the breast muscle (Figure 2 and Table S2).

Meanwhile, in order to ensure the accuracy of the analysis, the screened differentially expressed genes were analyzed by hierarchical cluster analysis where genes with the same or similar expression patterns clustered. The abscissa represented the sample clustering. Among them, one column represents a sample, and clustering is based on the similarity of gene expression between samples. The more similar the gene expression between samples, the closer the distance. This result suggested that these DEGs could significantly separate the samples into the leg muscle group and breast muscle group (Figure 3), indicating the reliability of our analysis.

3.4. Gene Ontology (GO) Annotation and Enrichment Analysis. To further reveal the molecular characterization of the DEGs, GO annotation was used to analyze the DEGs. The GO database contains three main branches: biological process, molecular function, and cellular component. The results showed that a total of 468 DEGs were annotated into 61 GO terms, including 24 biological processes, 19 cellular components, and 18 molecular function annotations. In biological processes, the significantly enriched GO terms mainly

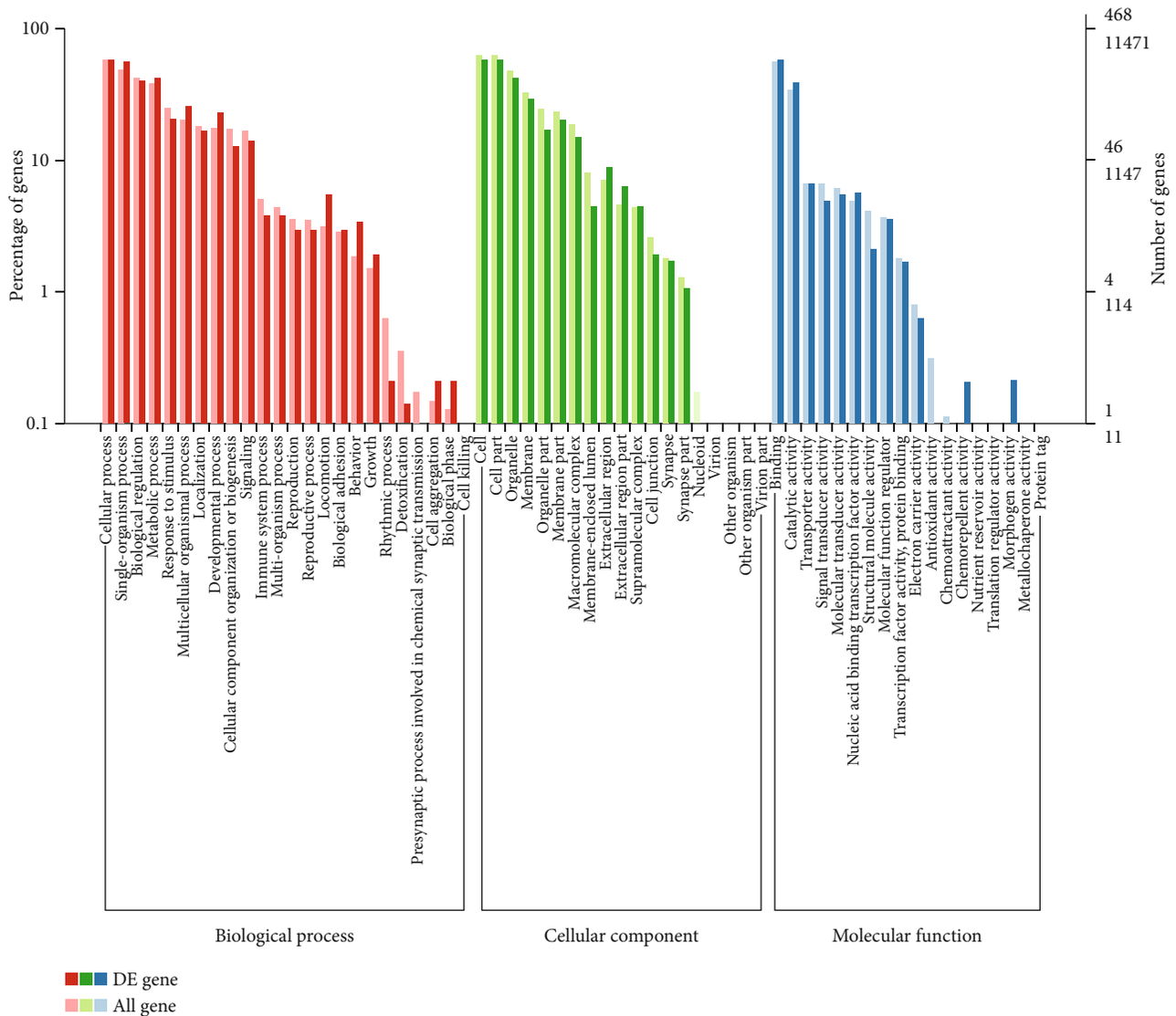


FIGURE 4: GO analysis of the DEGs.

contained cellular process, single organism process, biological regulation, metabolic process, response to stimulus, multicellular organismal process, localization, developmental process, cellular component organization, biogenesis signaling, etc. Cells, cellular parts, and organelles were significantly enriched in cellular components. Simultaneously, the main functional terms of molecular function were binding and catalytic activity, which were enriched with 273 and 183 genes, respectively (Figure 4).

In addition, some important genes ($FDR < 0.01$) that may be involved in the transformation of muscle fiber types are enriched in these GO terms, such as mitotic cell cycle process (*FHL1*, *PDS5A*) and calcium ion binding (*NKD1*, *RYR3*) (Table S3).

3.5. KEGG Pathway Enrichment Analysis. To know the signal pathways where the genes were distributed, a KEGG pathway analysis was performed. A total of 230 DEGs were mapped to 126 KEGG pathways, of which 20 pathways were significantly

enriched ($P \leq 0.05$) (Table S4). There were several extremely significantly enriched pathways (Figure 5), including glycolysis/gluconeogenesis, starch and sucrose metabolism, insulin signalling, and biosynthesis of amino acids. Some important genes were found to be enriched in these pathways: *PKM*, *LDHA*, *LDHA*, *PYGM*, and *HRAS* ($FDR < 0.01$) (Table S4).

3.6. Validation of DEGs by qRT-PCR. The expression of 7 randomly selected DEGs, including 4 downregulated genes (*FHL1*, *MYBPC1*, *ADPRHL*, and *TPM2*) and 3 upregulated genes (*HOMER3*, *MYH3*, and *BPGM*), was verified through quantitative real-time PCR. The results illustrated that the expression of *FHL1*, *MYBPC1*, *ADPRHL*, and *TPM2* in the breast muscle was significantly lower than that in the leg muscle, while the expression of *HOMER3*, *MYH3*, and *BPGM* was exactly the opposite (Figure 6(a)). As shown in Figure 6, the expression trend of these 7 genes was consistent

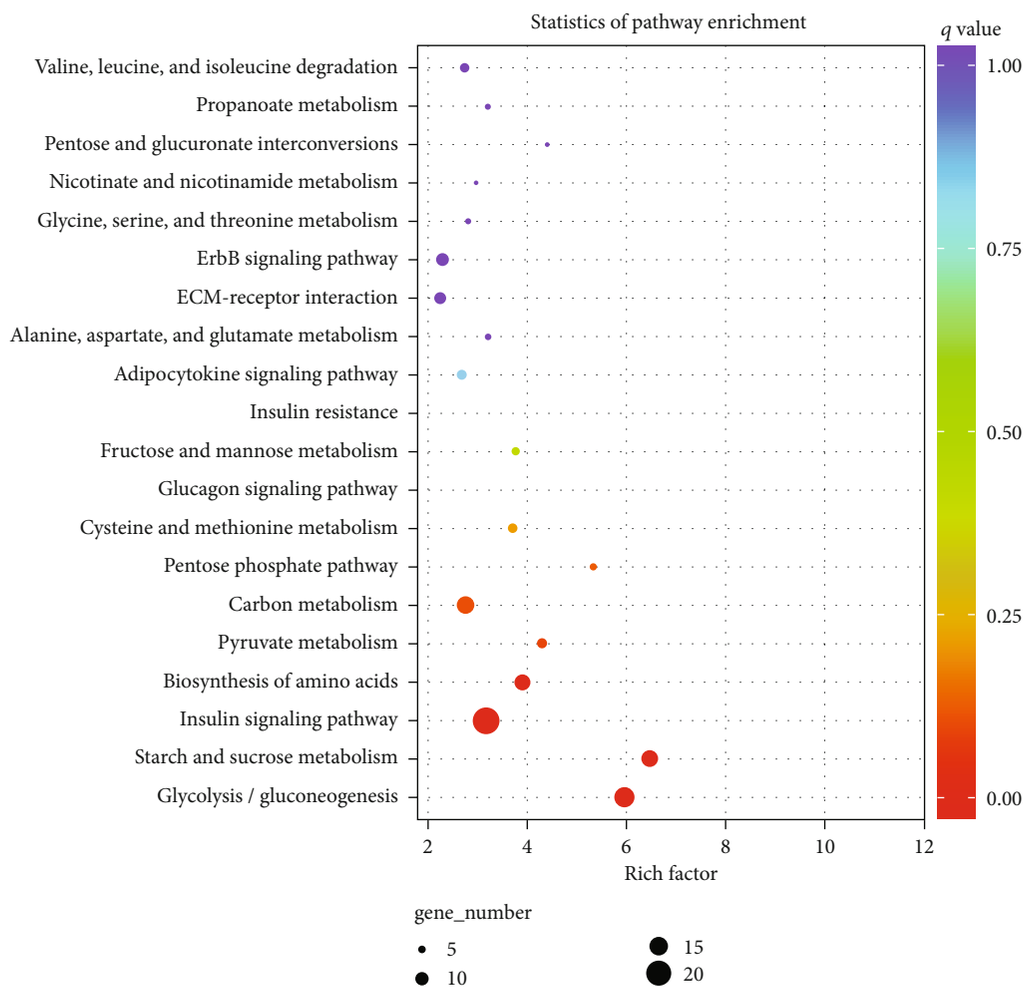


FIGURE 5: Top 20 significantly enriched KEGG pathways.

with the results of the RNA-seq experiment, which proved that the RNA-seq results were reliable (Figure 6(b)).

3.7. Identification of Chicken Skeletal Muscle Satellite Cells. To facilitate the study of muscle fiber growth characteristics in broilers, we isolated skeletal muscle satellite cells from the leg muscle of 15 chicken embryos which were incubated for 10 days *in vitro*. The cells in the process of proliferation and differentiation were identified by immunofluorescence with the skeletal muscle satellite cell-specific markers *Pax7*, *Desmin*, and *MyHC* [39]. The results showed that *Pax7* was positive in the nucleus (Figure 7(a)), and *Desmin* and *MyHC* were positive in the cytoplasm (Figures 7(d) and 7(g)). Therefore, the results showed that the isolated cells are indeed skeletal muscle satellite cells and could be used in the following research.

3.8. Expression Profile of *MyHC* Genes. In order to further explore the growth of muscle fibers in broilers, the expression patterns of *MyHC* subunits during cell differentiation were analyzed by qRT-PCR. *MyHC* IIA is a genetic marker of fast oxidative muscle fibers [40]. *MyHC* IIB is a genetic marker of fast glycolytic muscle fibers [41]. *MyHC* I is a genetic marker

of slow oxidative muscle fibers [42]. The results suggested that the expression of *MyHC* IIA was significantly upregulated during cell differentiation (Figure 8(a)). However, the expression of *MyHC* IIB was significantly downregulated on the second day of differentiation and then gradually upregulated from day 4 to day 6 (Figure 8(b)). Additionally, the expression of *MyHC* I was significantly upregulated at the beginning and decreased after the 4th day of differentiation (Figure 8(c)). The results indicated different development of muscle fiber types in broilers.

4. Discussion

The skeletal muscle of vertebrates is composed of muscle fibers with different morphological, contractile, and metabolic characteristics. Meanwhile, the properties of muscle fibers have an important effect on the meat quality of livestock [8]. In our study, we found that slow/fast muscle fiber marker genes were differentially expressed between the breast and leg muscles of broilers. The results were consistent with several previous studies demonstrating that the breast muscle of the chicken was a fast muscle, while the leg muscle was a slow muscle [43]. Therefore, RNA-seq was performed

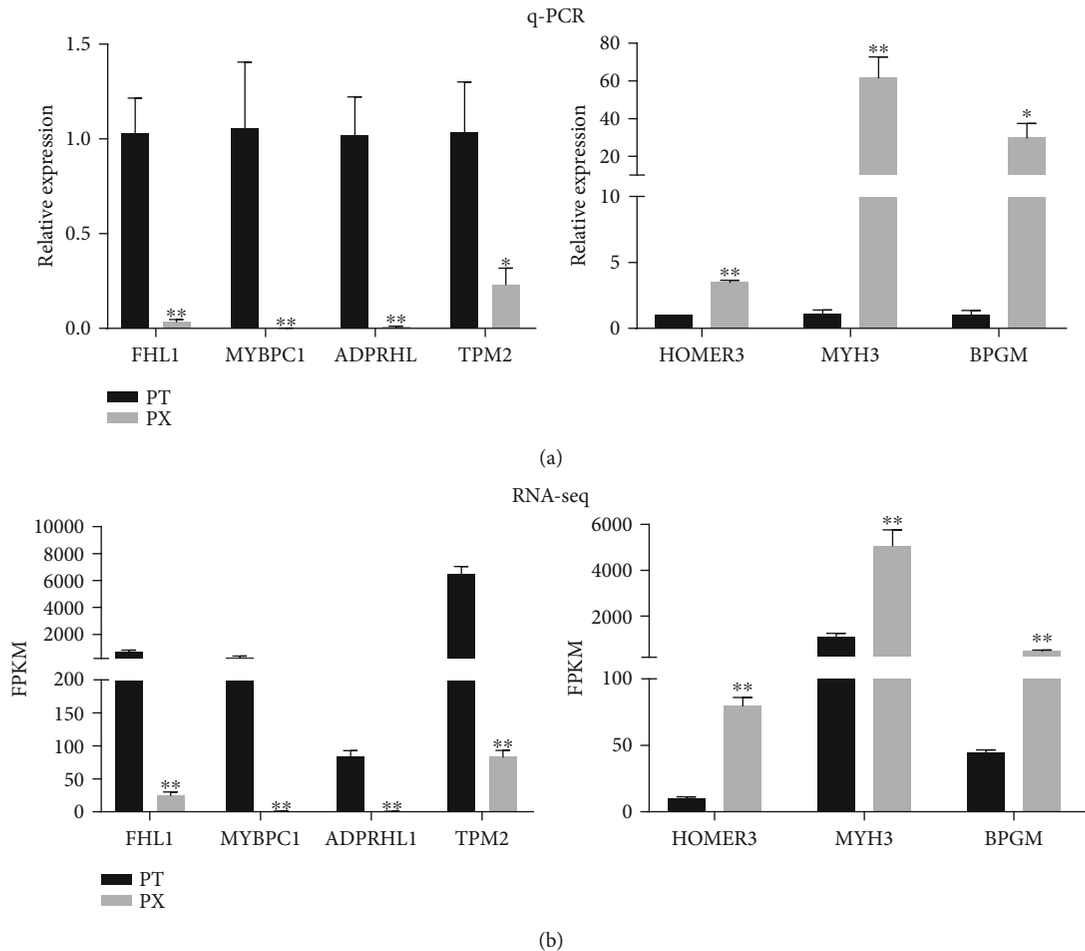


FIGURE 6: Validation of DEGs by qRT-PCR: (a) q-PCR results and (b) RNA-seq results (** $P < 0.01$, * $P < 0.05$; PT represents broiler leg muscle; PX represents breast muscle, $n = 5$).

on these two skeletal muscles to screen candidate genes related to muscle fiber-type conversion in our research. A total of 767 DEGs were discovered between both investigated groups, of which 7 DEGs were randomly selected and verified by qRT-PCR analysis. Although the expression fold changes of the selected genes between the two methods were not the same, the expression trend was highly consistent, proving that the results obtained by chip analysis were credible.

The basal processes of muscle cell growth and muscle fiber formation in broilers include cellular processes, single biological processes, and biological regulation. GO functional analysis showed that 11 DEGs were enriched in GO term mitotic cell cycle processes (Table S3). Among these DEGs, the *FHL1* gene was found to be highly abundant in oxidative muscle fibers [44]. Overexpression of *FHL1* promoted the transformation of muscle fibers to the oxidative type, while knockdown of *FHL1* could cause muscle fiber hypertrophy [45]. Slow muscle fibers are usually small in diameter, whereas the fast muscle fibers are usually large in diameter [46]. In this study, the *PDS5A* gene was also screened as an upregulated DEG in the breast muscle. *PDS5A* was found to be highly expressed in the skeletal muscle but low in other tissues and to have an effect on the proliferation of skeletal muscle cells [47].

These genes might play a key role in regulating muscle cell proliferation and muscle fiber transformation. In addition, intracellular calcium signalling is also an important factor that influences muscle fiber types [48]. Our study showed that 15 DEGs were enriched in the GO term calcium ion binding (Table S3). Among these genes, *NKDI* was upregulated in leg muscles. Some studies have shown that Wnt/ β -catenin signalling could transform slow muscle fibers into fast muscle fibers and that *NKDI* has a strong antagonistic effect on Wnt/ β -catenin signalling [49, 50]. Another gene, *RYR3*, controls calcium ion release channels in muscle cells [51]. *RYR3* has a high calcium sensitivity and regulates rapid muscle contraction together with the *RYR1* gene [52]. Some studies have shown that calcium sensitivity is related to the generation of fast myofibril subtypes, which was consistent with the results of our study [53]. KEGG pathway analysis also showed that the expression of calcium signalling pathway-related genes was altered significantly (Table S4). The function of these aforementioned DEGs in skeletal muscle fiber transformation needs further investigation.

One of the methods to classify fast or slow muscle is by their energy metabolism [54]. GO functional annotation analysis showed that a total of 198 DEGs were enriched in

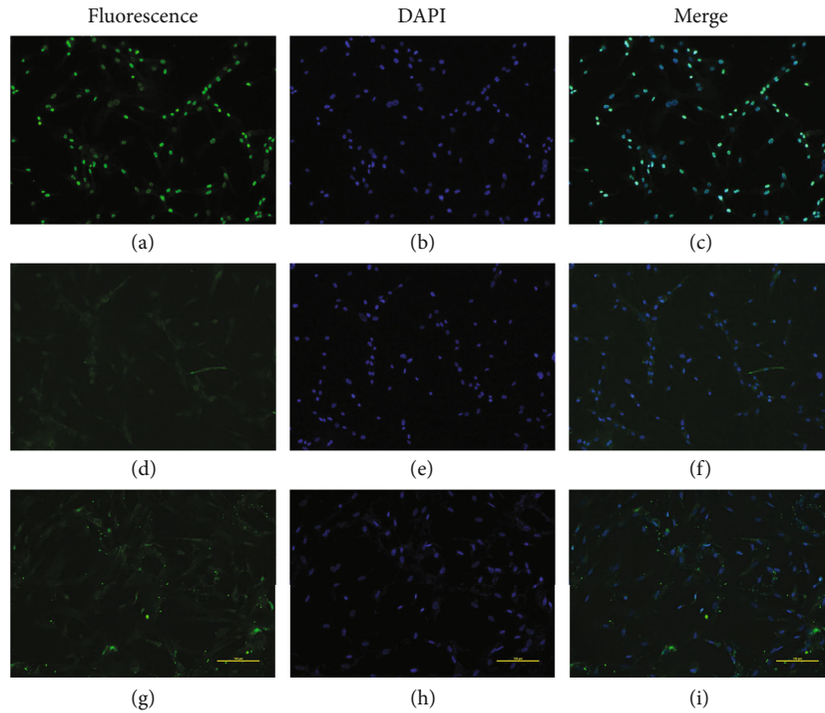


FIGURE 7: Identification of skeletal muscle satellite cells: (a) *Pax7* was detected with immunofluorescence; (b) DAPI stain of the nuclei of skeletal muscle satellite cells; (c) merged image of (a) and (b); (d) *Desmin* was detected with immunofluorescence; (e) DAPI stain of the nuclei of skeletal muscle satellite cells; (f) merged image of (d) and (e); (g) *MyHC* was detected with immunofluorescence; (h) DAPI stain of the nuclei of skeletal muscle satellite cells; (i) merged image of (g) and (h).

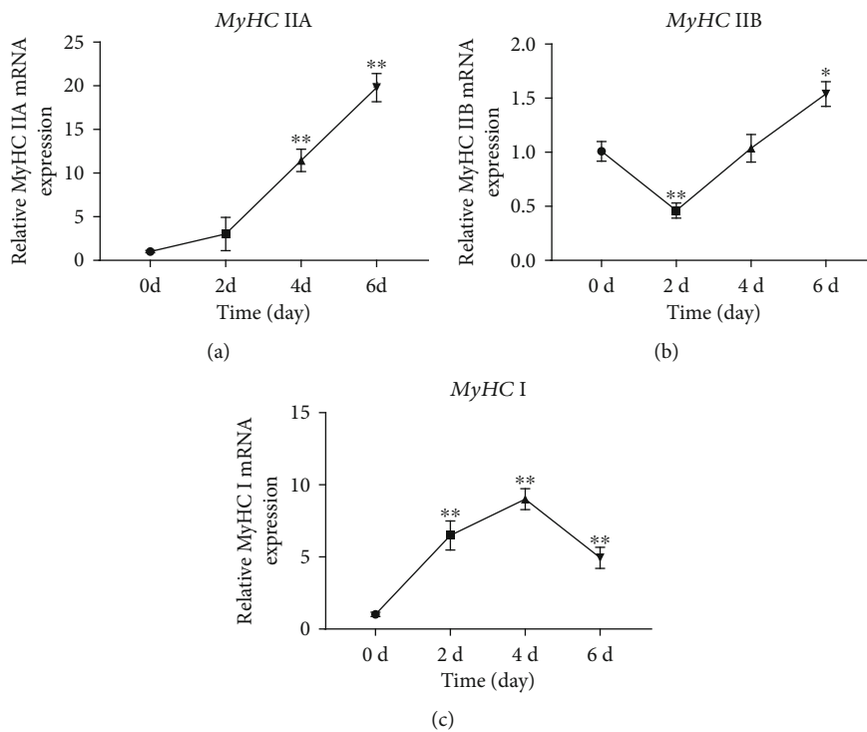


FIGURE 8: Expression of *MyHC* genes in the differentiation of chicken skeletal muscle satellite cells (** $P < 0.01$, * $P < 0.05$; $n = 3$).

metabolic process-related GO terms (Table S3). Previous studies showed that slow-twitch muscles have lower glycolytic enzyme activity, while fast-twitch muscles act mainly through the glycolytic pathway to meet the energy requirements for rapid muscle contraction [55, 56]. In our study, the KEGG pathway enrichment analysis results indicated that the four most significant enrichment pathways were related to the energy metabolism of muscle cells. Most of the DEGs involved in the glycolysis/gluconeogenesis pathway were upregulated in the breast muscle (15 of 17) (Table S4). Protein synthesis during skeletal muscle development is accompanied by energy metabolism, and the glycolysis/gluconeogenesis pathway plays a significant role [57]. *PKM* is a gene that catalyses the last step of glycolysis [58] and was upregulated in the breast muscle. The muscle type of pyruvate kinase (*PKM*) is one of the key mediators of the Warburg effect and tumor metabolism [59] and was closely related to the development of PSE meat [60]. Another gene, *LDHA*, the predominant *LDH* isoform in the skeletal muscle, is also an important enzyme involved in glycolysis [61]. The mutation at the C.423A>G site in exon 6 of porcine *LDHA* had a significant effect on the muscle fiber type II ratio [62]. In addition, 13 DEGs were enriched in the biosynthesis of amino acid pathway, of which 12 DEGs were upregulated in the breast muscle (Table S4). Another study found that genes related to glycolysis/gluconeogenesis and the biosynthesis of amino acid pathway were gradually upregulated during the growth and development of chicken breast muscle [63]. Therefore, we inferred that these DEGs were activated during fast-twitch muscle fiber formation.

Our KEGG results further discovered that 22 DEGs were enriched in the insulin signalling pathway (Table S4). Insulin plays a key role in a series of biological reactions, such as glucose uptake, lipid and protein synthesis, cell growth, and glycogen synthesis [64]. In skeletal muscle cells, the insulin signalling pathway regulates glycogen synthesis and decreases the rate of glycogen breakdown [65]. It has been shown that the content of glycogen in type I skeletal muscle fibers is significantly lower than that in type II muscle fibers [66]. We found that 5 DEGs in the insulin signalling pathway were also enriched in the GO term glycogen metabolic process (Table S3), including *GYS1* and *PYGM*; these were all upregulated in the breast muscle. The insulin signalling pathway stimulates the expression of *GYS1* mRNA and promotes the synthesis of glycogen in the skeletal muscle [67]. Skeletal muscle fiber type IIB predominantly uses glycogen and glucose as fuel, and the upregulation of *PYGM* accelerates glycogen phosphorylation [68]. In addition, 3 DEGs in the insulin signalling pathway were upregulated in the leg muscle, including *HRAS* (Table S4). The expression of *HRAS* was negatively correlated with metabolic traits related to the insulin signalling pathway [69]. Thus, *HRAS* may be a candidate gene involved in slow muscle fiber formation.

Skeletal muscle fibers are generally divided into four types according to the myosin heavy chain (*MyHC*) isoforms, namely, types I, IIA, IIB, and IIX [70]. However, there are very few type IIX muscle fibers in poultry skeletal muscle

[71]. A previous study showed that *Pax7*, *Desmin*, and *MyHC* were specific genetic markers of skeletal muscle satellite cells [72]. To further explore the mechanism of skeletal muscle fiber transformation in broilers, an in vitro cell model was successfully constructed in our study. Then, the expression profiles of *MyHC* isoforms (type I, type IIA, and type IIB) during broiler skeletal muscle satellite cell differentiation were detected. *MyHC* IIA mRNA expression increased throughout the time course. Expression of the *MyHC* IIB isoform was low on day 2 but rose until day 6. *MyHC* I expression first grew dramatically from day 0 to day 4 and then declined until day 6. Interestingly, these results coincided with those in a C2C12 mouse myoblast cell differentiation model [73]. Therefore, future studies will use these approximate time points of skeletal muscle fiber transformation in vitro to explore the function of the key DEGs in broilers.

5. Conclusion

In the current research, RNA-seq was carried out to compare the leg and breast muscles of broilers. A total of 767 DEGs were identified. Compared with leg muscle, there were 429 upregulated genes and 338 downregulated genes in the breast muscle. The expression profile of *MyHC* isoforms during broiler skeletal muscle cell differentiation was detected. Therefore, our study provides useful information for understanding the molecular mechanisms of muscle fiber-type transition in broilers. However, the transformation of skeletal muscle fiber in vivo is a dynamic process and these transcriptome data are preliminary, so the function of the DEGs still requires further investigation.

Data Availability

The data used to support the findings of this study are available from the corresponding author upon request.

Conflicts of Interest

The authors declare that there is no conflict of interest regarding the publication of this paper.

Authors' Contributions

Han Wang and Zhonghao Shen analyzed the data and drafted the manuscript. Han Wang, Xiaolong Zhou, Songbai Yang, Feifei Yan, and Ayong Zhao designed the experiments. Zhonghao Shen, Ke He, and Han Wang performed the experiments. All authors provided help for sample collection and reviewed the final version of the manuscript. Han Wang and Zhonghao Shen contributed equally to this work.

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Supplementary Materials

Table S1: primer sequences for the genes selected for qRT-PCR. Table S2: DEGs identified under certain filter criteria. Table S3: GO enrichment analysis of DEGs. Table S4: KEGG pathway analysis of DEGs. (*Supplementary Materials*)

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