Discovery of Novel and Differentially Expressed MicroRNAs between Fetal and Adult Backfat in Cattle

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Abstract

The posttranscriptional gene regulation mediated by microRNAs (miRNAs) plays an important role in various species. Recently, a large number of miRNAs and their expression patterns have been identified. However, to date, limited miRNAs have been reported to modulate adipogenesis and lipid deposition in beef cattle. Total RNAs from Chinese QinChuan bovine backfat at fetal and adult stages were used to construct small RNA libraries for Illumina next-generation sequencing. A total of 13,915,411 clean reads were obtained from a fetal library and 14,244,946 clean reads from an adult library. In total, 475 known and 36 novel miRNA candidates from backfat were identified. The nucleotide bias, base editing, and family of the known miRNAs were also analyzed. Based on stem-loop qPCR, 15 specific miRNAs were detected, and the results showed that bta-miRan25 and miRan26 were highly expressed in backfat tissue, suggesting these small RNAs play a role in the development and maintenance of bovine subcutaneous fat tissue. Putative targets for miRan25 and miRan26 were predicted, and the 61 most significant target transcripts were related to lipid and fatty acid metabolism. Of interest, the canonical pathway and gene networks analyses revealed that PPARα/RXRα activation and LXR/RXR activation were important components of the gene interaction hierarchy results. In the present study, we explored the backfat miRNAome differences between cattle of different developmental stages, expanding the expression repertoire of bovine miRNAs that could contribute to further studies on the fat development of cattle. Prediction of target genes analysis of miRan25 and miRan26 also showed potential gene networks that affect lipid and fatty acid metabolism. These results may help in the design of new intervention strategies to improve beef quality.

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Introduction

MicroRNAs (miRNAs) are a class of small regulatory noncoding RNAs with an average length of 22 nucleotides, which were first identified in C. elegans [1] and were reported to act as endogenous regulators of protein-coding genes at the posttranscriptional level in a sequence-specific manner [2]. MiRNAs have been implicated in fundamental biological processes such as differentiation, proliferation, apoptosis, and homeostasis [3]. In addition, recent reports have also suggested that miRNAs play crucial roles in the regulation of adipogenesis. In 2003, one report suggesting that miR-14 was involved in fat metabolism. The deletion of mir-14 results in animals with increased levels of triacylglycerol and diacylglycerol, whereas increases in the number of mir-14 copies have the converse effect [4]. Another study demonstrated that mir-143 is involved in adipocyte differentiation and may act by targeting the expression of gene extracellular-regulated protein kinase 5 (ERK5) [5]. Furthermore, the mir-8/mir-200 transcripts promote adipogenesis by inhibiting Wnt signaling, which is a negative regulator of adipogenesis [6,7]. In addition to mir-R-122, mir-R-370, mir-R-37b/378*, mir-R-27, and mir-R-355 have also been recently associated with lipid metabolism and adipocyte differentiation [8–12]. Mir-122 inhibition in high-fat fed mice was found to result in reduced plasma cholesterol levels, which was linked to a reduction in cholesterol synthesis and the stimulation of fatty-acid oxidation [8]. Mir-370 has similar effects as mir-122 on lipid metabolism by targeting carnitine palmitoyl transferase (Cpt1a) to reduce fatty acid oxidation [9]. The overexpression of mir-R-37b/378* during adipogenesis increases the accumulation of triacylglycerol [10]. By contrast, mir-R-27 inhibits the expression of peroxisome proliferator-activated receptor (PPAR) γ and C/EBPα, the two master regulators of adipogenesis [11]. In contrast, increased mir-R-355 expression has been associated with an elevated body, liver and white adipose tissue weight, and hepatic triglyceride and cholesterol [12]. Later studies have also demonstrated mir-R-33’s role in the regulation of lipid metabolism, while inhibition of mir-R-33 resulted in increased fatty acid oxidation and the reduced accumulation of fat stores [13,14]. However, the major role of miRNAs in regulating lipid metabolism and adipogenesis remains unknown.

Release 19 of mirBASE (http://www.mirbase.org) contains 21,264 entries representing hairpin precursor miRNAs, expressing...
25,141 mature miRNA products in 193 species. However, the number of known bovine miRNAs is limited, with only 755 reported compared with 2,042 from humans and 1,281 from mice. Cattle are an important species in the animal production industry. They have tremendous importance not only for food production, but also as a mammalian model organism for comparative genomics and biological studies [15]. Additionally, lipid deposition, especially in subcutaneous adipose depots, is directly associated with yield grade and meat quality [16]. Previous studies have provided limited insight into the miRNA population present in bovine species by only investigating general characteristics, expression patterns, and features of their target genes [17–21]. In 2010, Jin et al. reported that approximately 20% of the miRNAs involved in adipogenesis and lipid deposition were identified as being correlated with backfat thickness. Their results suggest that miRNAs play a regulatory role in white adipose tissue development in beef [22].

Given the emerging roles of miRNAs in fat tissue development in multiple species, identifying the differentially expressed miRNAs is an important first step to investigating the function of miRNAs in the course of bovine lipid metabolism and adipogenesis. In the present study, using both next-generation sequencing and reverse transcription, quantitative PCR (RT-qPCR) assays were employed to characterize the genome-wide miRNA expression profile in bovine backfat between fetal and adult periods. We sought to identify a panel of fat depot-specific miRNAs that could serve as targets for further study with a long-term goal of using this information to control backfat deposition in fed cattle.

### Results

#### Deep sequencing of bovine short RNAs

In order to identify novel and differentially expressed miRNAs in the bovine backfat at fetal and adult stages, two small RNA libraries were constructed for Solexa SBS technology sequencing. Solexa sequencing provided a total of 14,071,065 and 14,373,930 reads of 3 nt–30 nt from the fetal and adult backfat tissue libraries, respectively. After removing the low quality reads, adaptors, and insufficient tags and sequences, a total of 13,915,411 and 14,244,946 clean reads of 18–30 nt were obtained (Table 1). The unique and total numbers of the common and tissue-specific small RNA sequences in the two libraries are shown in Figure 1. The fetal-specific unique sequences account for 37.51% of all sequence reads and 50.63% in the adult library, respectively (Figure 1A). The percentages of the fetal-specific and adult-specific sequences were 2.34% and 3.88% of the total number of small RNAs in the two libraries (Figure 1B). Length distribution analysis showed that most reads ranged from 21 to 24 nt, which is typical of the small RNA of Dicer-processed product. The size distribution (18 nt–30 nt) of the small RNA from the fetal and adult stages in Chinese Qinhuang cattle was similar (Figure 2). For example, in the fetal period, the 22 nt and 23 nt sequences were the dominant small RNAs, which accounted for 51.80% and 37.51% of the total reads, respectively.
21.46% of the total sequences as well as 47.42% and 14.68% in the adult bovine library.

Next, all of the reads were aligned against the Bos taurus genome (Btau_6.0) using the SOAP Program [23]. A total of 9,063,361 reads were matched to the bovine genome in the fetal library (127,542 unique sRNAs) and a total of 9,690,034 reads were in the adult library (250,668 unique sRNAs). Subsequently, the genome-matched small RNA sequences were clustered into several RNA classes such as known miRNAs, degraded fragments of mRNA, repeats, rRNA, tRNA, snRNA/snoRNA, and others (Table 2). Known bovine miRNAs accounted for 59.17% of all sequence reads in the fetal library and 47.89% in the adult library, suggesting that mature miRNAs were highly enriched in our small RNA libraries. However, after analyzing the number of unique sequences, the proportion of small RNA sequences derived from known miRNAs represented only a very small fraction of the total sequences, the proportion of small RNA sequences derived from known miRNAs expressed in backfat tissues of some of the cattle. However, 16 and 15 miRNAs from the fetal and adult bovine libraries, respectively, correspond to 0.28 and 0.08 miRNAs per 1 Mbp genomic sequences. Chromosome 21 has the greatest number of known miRNAs in the fetal and/or adult libraries. Most of the conserved miRNAs were found on the autosomes or X chromosome of the cattle. However, 16 and 15 miRNAs from the respective fetal and adult bovine backfat libraries failed to match the genomic sequences, which might be due to an incomplete assembly model of the bovine genome. In detail, the shortest chromosome, 25, and the longest chromosome, 1, encode 12 and 14 miRNAs, respectively, corresponding to 0.28 and 0.08 miRNAs per 1 Mb genomic sequences. Chromosome 21 has the greatest number of known miRNAs distributed in the cattle genome (69 miRNAs). The X chromosome has the second greatest number of the known miRNAs, accounting for 3.08% (38/1,245) of the total miRNAs. It is noteworthy that the expressed miRNAs in the chromosomes X and 21 are dominant in the two libraries (Table S1). To date, 25,141 mature miRNA products have been identified in 193 species. In our study, further analysis identified a total of 432 and 412 conserved miRNAs that belonged to 236
miRNA families in fetal and adult bovine libraries. The identified miRNA families have been shown to be conserved in a variety of species. For example, let-7, miR-1, miR-25, miR-9, and miR-25 families have been found in 68, 69, 68, 72, and 68 species, respectively, while miR-2363, miR-3432, miR-3604, miR-6526, and miR-6536 families have only been detected in *bos taurus* (Table S5). The largest miRNA family size identified was miR-2284, which consisted of 27 members; miR-2285, let-7, miR-30, and miR-376 possessed 22, 8, 6, and 5 members, respectively, whereas other miRNA families such as miR-107, miR-122, miR-140, and miR-1839 had only one member (Table S1). In addition, different family members also displayed drastically different expression levels. For example, the abundance of the miR-2284 family varied from 0 (bta-miR-2284e) to 57,585 reads (bta-miR-2284x) with deep sequencing in the fetal stage. The existence of a dominant member in an miRNA family may suggest that the regulatory role of this family was performed by the dominant member at the developmental time when the samples were collected for RNA extraction. Abundance comparisons of different members in an miRNA family may provide valuable information on miRNAs’ role in that specific stage of bovine development.

**Newly identified miRNAs in backfat**

The miRNA hairpins are mostly located in intergenic regions, introns, or in the reverse repeat sequence of the coding sequence. The characteristic hairpin structure of miRNA precursor can be used to predict novel miRNA. We used the prediction software Mireap to predict novel miRNA by exploring the secondary structure, the Dicer cleavage site, and the minimum free energy of the unannotated small RNA tags that could be mapped to genome. Mireap can be accessed from the following link: http://sourceforge.net/projects/mireap/. Seven of the key conditions for predicting novel miRNA are as follows [25]: (1) the tags that be used to predict novel miRNA are the unannotated tags that can match to reference genome, the tags that can align to the intron region, and the tags that can align to the antisense exon region; (2) hairpin miRNAs can fold secondary structures, and mature miRNAs are present in one arm of the hairpin precursors, which will be considered as candidate miRNA genes; (3) the mature miRNA strand and its complementary strand (miRNA*) present 2 nt 3' overhangs; (4) hairpin precursors lack large internal loops or bulges; (5) hairpins’ secondary structures are steady, with a free energy of hybridization lower than or equal to $-18$ kcal/mol; (6) miRNA precursors with secondary structures have higher minimal

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**Table 2. Distribution of the genome-mapped sequence reads in small RNA libraries.**

| Locus class | Fetal bovine backfat | Adult bovine backfat |
|-------------|----------------------|----------------------|
|             | Unique sRNA | Percent (%) | Total sRNA | Percent (%) | Unique sRNA | Percent (%) | Total sRNA | Percent (%) |
| Total       | 687753 | 100% | 13915411 | 100% | 870506 | 100% | 14244946 | 100% |
| miRNA       | 357 | 0.05% | 496 | 0% | 385 | 0.04% | 683 | 0% |
| Exon_antisense | 23294 | 3.39% | 29572 | 0.21% | 67318 | 7.73% | 94168 | 0.66% |
| Exon_sense  | 6521 | 0.95% | 8108 | 0.06% | 12650 | 1.45% | 14366 | 0.10% |
| Intron_antisense | 12719 | 1.85% | 19684 | 0.14% | 28458 | 3.27% | 37886 | 0.27% |
| Intron_sense | 3827 | 0.56% | 8234182 | 59.17% | 3857 | 0.44% | 6821393 | 47.89% |
| rRNA        | 116439 | 16.93% | 768966 | 5.53% | 136492 | 15.68% | 2630720 | 18.47% |
| repeat      | 12587 | 1.83% | 26041 | 0.19% | 28962 | 3.33% | 78032 | 0.55% |
| stRNA       | 325 | 0.05% | 10067 | 0.07% | 517 | 0.06% | 17019 | 0.12% |
| snRNA       | 3838 | 0.56% | 9407 | 0.07% | 6382 | 0.73% | 24007 | 0.17% |
| snoRNA      | 1932 | 0.28% | 10112 | 0.07% | 3009 | 0.35% | 13710 | 0.10% |
| trpRNA      | 964 | 0.14% | 5291 | 0.04% | 1666 | 0.19% | 21352 | 0.15% |
| tRNA        | 17413 | 2.53% | 105611 | 0.76% | 19699 | 2.26% | 175326 | 1.23% |
| Unknown     | 487537 | 70.89% | 4687874 | 33.69% | 561111 | 64.46% | 4316284 | 30.30% |

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**Table 3. Summary of known miRNA in each sample.**

| miR | miR-5p | miR-3p | pre-miRs | Unique matched to pre-miRs | Read matched to pre-miRs |
|-----|--------|--------|---------|---------------------------|--------------------------|
| Known miRs | 598 | 79 | 78 | 766 |
| Fetal library | 336 | 46 | 50 | 457 | 3890 | 8234368 |
| Adult library | 328 | 40 | 44 | 442 | 3926 | 6821671 |

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free energy indexes (MFEIs) than other different types of RNAs. Based on Solexa sequencing, we identified 36 novel bovine miRNAs that corresponded to 63 genomic loci. Ten novel miRNAs were in the fetal bovine library and nine were in the adult bovine library; 17 of these overlapped in both libraries (Table S7). An examination of pre-miRNAs and other RNAs (tRNA, rRNA, and mRNA) revealed that miRNAs were significantly different from other RNAs [26]. Specifically, most miRNA precursors had an MFEI greater than 0.85, significantly higher than tRNAs (0.64), rRNAs (0.59), or mRNAs (0.65). The results suggested that the MFEI can easily be used to distinguish miRNA from other non-coding and coding RNAs. This provides a more precise criterion to predict miRNAs using computational approaches, and in our database approximately 41 pre-miRNAs had an MFEI greater than 0.85. In addition, the expression profiles of the novel miRNA genes we obtained by measuring the sequencing frequencies are shown in Figure 3B, indicating that 24 miRNAs consisted of 10 up-expressed and 14 down-expressed miRNAs.

miRNAs differentially expressed between adult bovine muscle and backfat

In an earlier miRNA discovery study, we observed 521 miRNAs including 104 novel miRNA candidates from the Chinese Qinchuan bovine longissimus thoracis [21]. In order to identify the fat-specific miRNAs involved in bovine lipid metabolism and adipogenesis, we compared the backfat library with the muscle library and found that miRNA expression was different in the two tissues. In the muscle, bta-let-7a/b/f, bta-miR-1, and bta-miR-206 were the dominant expressed miRNAs, with more than 100,000 reads. They constituted 96.83% of the total known miRNA sequencing reads, suggesting that they have abundant in muscle tissue. The sequencing frequencies of the 58 miRNAs (e.g., bta-miR-122, bta-miR-1185, bta-miR-1224) were much lower (1 ≤ sequencing reads ≤ 10). However, in fat tissue, seven miRNAs (bta-let-7a/b/c/e/f, bta-miR-154c, and bta-miR-199a-3p), each with more than 100,000 reads, were the most abundant. In the two libraries, a total of 233 known miRNAs were co-expressed. Among the known miRNAs, only one fat-specific and 17 muscle-specific miRNAs were found (Table S8A), and the expression levels of the tissue-specific expressed miRNAs were low. In comparison with miRNA expression in muscle tissue, 195 miRNAs in backfat tissue were significantly up-regulated, while 28 were significantly down-regulated (Figure S2A). In addition, five novel miRNAs were identified, and the results are shown in Table S8B. The expression levels of novel miRNAs in the muscle and backfat tissue were also shown in Table S8C and Figure S2B. The read number for each novel miRNA was much lower than that of the majority of conserved miRNAs. For instance, bta-miRn26 and bta-miRn11 were both novel miRNAs, and the total reads were only 286 and 15 in the backfat library, respectively. However, the total reads of two conserved miRNAs (bta-let-7a and bta-miR-154c) were striking, at 1,636,370 and 120,128, respectively. This might suggest that these novel miRNAs play a specific role during the developmental stages. Since this study investigated Chinese Qinchuan bovine muscle and backfat tissue miRNAs, whether these low-abundant miRNAs are expressed at higher levels in other tissues and organs remains to be investigated.

Validation of miRNA expression

To validate our sequencing data, stem-loop qPCR [26] analysis of miRNA expression was performed in the heart, liver, lung, kidney, intestines, stomach, muscle, and backfat of female animals. At least five animals were used for each sample. The expression...
levels of the 15 miRNAs were determined, and the selected miRNAs included five reported tissue-specific miRNAs (bta-miR-9, miR-124, miR-122, miR-27, and miR-103), seven backfat-predominant miRNAs in fetal and/or adult bovine backfat libraries (let-7a, miR-140, miR-199a, miR-320, miR-2234a, miRn9, and miRn25) as well as three high-read miRNAs (miR-154, miR-1839, and miRn26) in backfat that were compared with the muscle library. A comparison of miRNA expression profiles among tissues revealed that very few miRNAs had tissue-specific expression (e.g., miR-122 and -2234a in the liver). Several miRNAs were found to be highly expressed in particular tissues: miR-1839 in the heart, miR-103 in the liver, miR-9 and -154 in the kidneys, miRn25 in fat, and miRn26 in muscle and fat tissue, respectively. In contrast, another seven miRNAs were found in six or more tissues, and the two most abundant miRNAs across the eight tissues were miRNA-199a and miRNA-320 (Figure 4A and Figure S3). To further explore the highly expressed miRNAs between cattle of different genders, we also performed a quantitative analysis of the miRNAs in adult cow, bull, and steer backfat. miRNAn25 and miRNAn26 were expressed at significantly higher levels in cow than bull and steer backfat (Figure 4B).

**Prediction of target genes**

The miRNA expression patterns in different tissues have been investigated. In cattle, miR-9 and miR-124 in the brain, miR-122 in the liver, and miR-1, miR-133a, and miR-206 in muscle are all tissue-specific [27]. In the present study, miRnAn25 and n26 were also specifically identified in backfat. The similar expression pattern indicates that tissue-specific miRNAs may have certain roles in the respective tissues. We therefore could postulate that miRn25 and n26 may influence lipid metabolism and adipogenesis in backfat tissue. To identify the potential function of the two novel miRNAs, target prediction was performed using RNAhybrid software. A total of 2,416 putative target sites for miRNA25 (Table S9A) and 672 target sites for miRNA26 (Table S10A) were identified in the cattle’s backfat, respectively. Next, all the target gene candidates were submitted for homology and annotation analyses and Gene Ontology (GO) annotation using an online version of the Blast2GO program (www.Blast2GO.com). In total, 2,416 target sites of miRNA25 and miRNA26 were annotated with 1,818 GO terms (Table S9B), and 841 GO terms (Table S10B) were associated to 672 target sites of miRNA26 in the Non-Redundant database. The distributions of GO term categories for the target genes of miRNA25 and miRNA26 were highly similar (Figure S4). The top five biological functions identified by the Blast2GO program included categories related to a wide variety of physiological and biological events, such as cellular processes, metabolic processes, biological regulation, responses to stimulus, and multicellular organismal processes. Further analyses revealed that 126 miRNA25 target genes relating to lipid metabolism were annotated with 32 GO terms (Table S9C), and 44 target sites for miRNA26 were annotated within 10 GO terms (Table S10C), respectively.

Previous reports have documented the functional connection between RNA editing and miRNA-mediated post-transcriptional gene silencing [28]. Therefore, to characterize the potential effects of RNA editing on miRNAs, we first identified differentially expressed transcripts between fetal and adult bovine backfat based on RNA-Seq technology. In total, 7,495,616 reads could be uniquely aligned in the fetal library and 6,088,485 reads were in the adult library, representing ~18.04 Gbp and ~14.89 Gbp of expressed sequences in the respective libraries. Of the 3,879 known genes that were expressed in bovine backfat, 62 and 95 were differentially expressed at fetal and adult stages, respectively. Of those, 26% up-regulated and 74% down-regulated transcripts were identified by comparing the adult stage to the fetal stage (Table S11). Additionally, 40.47% miRNA25 target genes related to lipid metabolism were expressed in the fetal and/or adult stages (Table S9D), as well as 36.36% target genes for miRNA26 (Table S10D), respectively. An MA-plot indicated that most of the target genes related to lipid and fatty acid metabolism were down-regulated, accounting for about 91.80% of the total transcripts (Figure 5).

The 61 most significantly different miRNA25 and n26 target transcripts related to lipid and fatty acid metabolism were investigated using Ingenuity Pathways Analysis software (IPA, www.ingenuity.com), and the results are shown in Figure 6. In comparison with the adult stage, five of these transcripts had increased gene expression in the fetal stage and 56 had decreased expression. Briefly, genes that play a crucial role in lipid synthesis (ABCA1, ACACA, ACAD1, BMP2, ESR1, FGFI, and SDR5D1), lipid accumulation (CD36, LDLR, and SREBF1), lipid homeostasis (EHHADH and HNF4A), fatty acid metabolism (PPARG, ACSL1, SLC16A1, FOX, and LDLR), fatty acid oxidation (LIPE and FTO), and fatty acid efflux (SREBF2) were down-regulated in the fetal stage rather than in the adult stage. On the contrary, the gene involved in fatty acid concentration, lipid uptake, and steroid synthesis (HSD3B7) was up-regulated. When comparing the fetal stage with the adult stage, the most representative canonical pathways significantly modulated in bovine backfat were involved in peroxisome proliferator-activated receptor α/retinoid X receptor alpha (PPARα/RXRα) and farnesoid X receptor/retinoid X receptor (FXR/RXR) activation, etc. (Table 4).

**Discussion**

Adipose tissue is a highly specialized structure in energy storage. In mammals, the adipose tissue pool is composed of at least two functionally different types of fat: white and brown fat. White adipose tissue is the primary site of energy storage and of the release of hormones and cytokines [29]. Excess accumulation of white adipose tissue causes obesity. Brown adipose tissue, on the other hand, affects whole-body metabolism and may alter insulin sensitivity [30] and modify susceptibility to weight gain [31]. However, brown adipose tissue is primarily only found in fetuses and young individuals and is believed to have no physiologic relevance in adults, resulting in significant differences between fetal and adult adipose tissue development. In addition, potential regulators of adipogenesis include miRNAs, which encode an abundant class of 22 nucleotide evolutionarily conserved RNAs that control gene expression at the posttranscriptional level by targeting miRNAs for degradation and/or translational repression [32]. Computational and experimental analyses suggest that miRNAs may regulate the expression of about 30% of human and mouse genes [33]. Recently, miRNA expression profiles and functions have been extensively investigated, but little is still known about the role of miRNAs in metabolic tissues, particularly adipose tissue. Of particular relevance, miR-14 [34] and miR-278 [4] in the fat of flies regulate lipid metabolism, and miR-122 in the liver of mice controls triglyceride metabolism and cholesterol biosynthesis [8]. In the current investigation, we therefore hypothesized that miRNA expression would precede de novo lipogenesis in adipose tissue. To address our hypothesis, fetal and adult Chinese Qinhuian bovine backfat samples were collected, and two miRNA libraries were constructed for Solexa SBS technology sequencing.

Of the mappable sequences, the majority of the small RNAs were 21–24 nt in size, which coincided with the known specificity for Dicer processing and the features of mature miRNAs [35].
In the present study, the 22 nt sequences in bovine backfat (including the fetal and adult samples) were the dominant small RNAs. This was in agreement with Chen et al., who reported that 21–23 nt sequences was significantly greater than others, and almost half of the sequences in the backfat of Large White and Meishan pigs were canonical 22 nt miRNA [36]. Recently, similar results in Qinchuan bovine [21], Texel [37], and carp [38] muscle and goat mammary glands [39] have also been reported. However, our findings are not consistent with the previous studies on bovine testis and ovaries [40,41] and even of those on maize [42]. The differences observed between our study and previous studies may have been due to different experimental approaches (Solexa Sequencing vs random cloning), different species (cattle vs maize), and different tissues (adipose tissue vs testis and ovary tissues). In addition, the variations in sequence length may be also attributed to differences in functional roles, such as RNA editing in miRNA-mediated gene silencing [43], 3’-editing [44], and degradation of microRNAs by a family of exoribonucleases [45].

The XX/XY chromosomes are a biological system that determines the development of sexual characteristics in most mammals. We calculated the densities of the miRNAs on the chromosomes using the miRNA data from the miRBase and found that there were no available data on the Y chromosome in cattle. This was supported by a previous deep-sequencing study in cattle [40] and sheep [37]. The bovine X chromosome is 149 Mbp in size, ranking the 2nd largest among all of the chromosomes. In this study, 38 unique miRNAs were located on the X chromosome, accounting for 8% (38/475) of genome-mapped miRNAs. The density distribution of miRNA loci on the X chromosome reached 0.25 miRNA loci per Mbp, which ranked 3rd in the density distribution of miRNA loci among all of the chromosomes. Higher densities of miRNAs on X chromosomes were also identified in eight other mammalian species [46], consistent with prior studies.
that demonstrated their resistance to meiotic sex chromosome inactivation [47].

MiRNAs from adipose tissue of distinct breeds of cattle had different levels of expression and backfat thickness [22]. Further, a recent study showed variations in miRNA profiles among different locations of subcutaneous backfat, and in beef cattle, miR-9 and miR-124 in the brain and miR-122 in the liver were all tissue-specific [27]. In the present study, these miRNAs were also identified, and the results suggest that some of the previously reported tissue-specific miRNAs maybe have certain limits in terms of their distribution in tissues. Liver-specific miR-122 has a diversity of roles in liver, e.g., metabolism, hepatocarcinogenesis.

Figure 6. Functional and pathways analysis using miRNASeq target genes related to lipid and fatty acid metabolism as input data. The node colour indicated the expression of genes: (red) up-regulated and (green) down-regulated in adult stage relative to fetal stage. The shapes of nodes indicated the functional classes of the gene products. Relevant canonical pathways that feature modulated genes were indicted as well (e.g. LXR/RXR Activation and PPARα/RXRα Activation).

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| Table 4. Description of the top ten canonical pathways significantly modulated in bovine backfat tissue when comparing fetal stage with adult stage. |
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| **Ingenuity canonical pathways** | **Genes** |
| PPARα/RXRα Activation | ACADL, CD36, ABCA1, ERK1/2, HDL-cholesterol, LDL, N-cor, Pka, Proinsulin, Rxr |
| FXR/RXR Activation | HDL-cholesterol, PPARG, SLC10A1, SREBF1, HNF4A, LDL, Nh1h, PEPCK, Proinsulin, Rxr |
| LXR/RXR Activation | LDLR, SREBF1, CD36, ACACA, ABCA1, HDL-cholesterol, LDL, N-cor, Nh1h, Rxr |
| Hepatic Cholestasis | HDL-cholesterol, SLC10A1, SREBF1, HNF4A, ESR1, HSD3B7, N-cor, Pka, Proinsulin, Rxr |
| LPS/IL-1 Mediated Inhibition of RXR Function | SLC10A1, FABP5, SREBF1, ALAS1, ACSL1, ABCA1, HDL-cholesterol, LDL, Nh1h, Rxr |
| RAR Activation | RDH10, PIK3R1, BMP2, ERK1/2, HDL-cholesterol, LDL, N-cor, Rxr, SRC |
| TR/RXR Activation | LDLR, HDL-cholesterol, SREBF1, N-cor, PEPCK, ACACA, Rxr, PCK1 |
| AMPK Signaling | PIK3R1, LIPE, ACACA, PPP2R5A, ERK1/2, Pka, SRC, Proinsulin |
| ERK/MAPK Signaling | ERK1/2, ESR1, Pka, PPARG, PPP2R5A, SRC |
| PPAR Signaling | ERK1/2, N-cor, Nh1h, PPARG, Proinsulin, Rxr |

Note: Statistical significance of pathway modulation was calculated via a right-tailed Fisher’s Exact test in Ingenuity Pathway analysis and represented as –log (P value): -log values exceeding 1.30 were significant FDR < 0.05.
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bovine tissues suggest that these miRNA may be more relevant to expressed at significantly higher levels in cow tissue (Figure 4B).

To better understand the functions of the identified miRNAs in lipid metabolism and adipogenesis, putative targets of the highly expressed miRn25 and n26 in bovine backfat were predicted (Table S9A and S10A). According to the analysis and annotation, targets of miRn25 and n26 had diverse functions, ranging from genes encoding transcription factors involved in signal transduction, enzymes involved in metabolism, various kinases, and isomerase and helicase to genes regulating oxidative reduction and transport (Table S9B and S10B). Further analysis identified that 61 most significantly different miRNAn25 and n26 target transcripts related to lipid and fatty acid metabolism overlapped with the bovine backfat transcriptome profile obtained by RNA-Seq (Table S9D and S10D). More specifically, an important target was the PPARγ gene (Figure 6), which was a dominant regulator of adipogenesis and fat cell gene expression [61]. Other important genes targeted by the highly expressed miRNAs included acyl-CoA synthetase long-chain family member 1 (ACSL1), lipase, hormone-sensitive (LIPE), CD36 molecule (thrombospondin receptor) (CD36), and acetyl-CoA carboxylase alpha (ACACA), which are all known to be involved in lipid metabolism and adipogenesis. We therefore hypothesized that miRNAn25 and n26 can target sequences in these genes to regulate the development of bovine fat tissue.

Finally, the identified target genes were mapped to the genetic networks, and an IPA network score of 49 and 25 genes of interest (GOI) presented functions related to lipid metabolism, small molecule biochemistry and molecular transport (Figure 6). Many of these GOI have been previously associated with lipid metabolism and adipogenesis (e.g., fatty acid oxidation and lipid accumulation and homeostasis). These results support the utility of our methods for identifying the significant genes related to this biological function. Additionally, several GOI related to lipid and fatty acid metabolism that appear to be novel in the context of the IPA network were also identified, providing new targets for future study.

This study expands the repertoire of bovine miRNAs and could initiate further study in the fat development of cattle. In addition, the miRNA expression patterns among eight tissues in beef cattle showed that most miRNAs are ubiquitously expressed, suggesting that these miRNAs may play a role in a broad range of biological processes in various tissues. However, miRn25 and n26 were significantly expressed in bovine backfat, suggesting that they are likely to play a role in the development of bovine fat tissue and could be potential molecular targets for future study.

Conclusions

We have identified 475 known miRNAs and 36 novel miRNAs in the backfat of fetal and adult Qinhuang bovines using deep sequencing technologies. This study expands the repertoire of bovine miRNAs and could initiate further study in the fat development of cattle. In addition, the miRNA expression patterns among eight tissues in beef cattle showed that most miRNAs are ubiquitously expressed, suggesting that these miRNAs may play a role in a broad range of biological processes in various tissues. However, miRn25 and n26 were significantly expressed in bovine backfat, suggesting that they are likely to play a role in the development of bovine fat tissue and could be potential molecular targets for future study.

Bulls have less fat than cows and steers, resulting in less tasty meat, particularly because bulls are slaughtered at a younger physiological age than other meat-producing cattle. In addition, they have higher levels of energy expenditure and protein retention than cows and steers, contributing to a lower availability of nutrients for fat deposition. Nowadays, molecular biology techniques, especially genomics, continue to further expand our knowledge of the adipocyte. In addition, new genetic markers, transcription factors, and genes are continuously being discovered [59]. In order to further expand the key mechanisms that can act to improve fat level between cattle of different sexes, highly expressed miRn25 and n26 in bovine backfat were analyzed in adult cow, bull, and steer backfat tissues, and that they were expressed at significantly higher levels in cow tissue (Figure 4B). The expression patterns and level of miRn25 and n26 in the tested bovine tissues suggest that these miRNA may be more relevant to the highly conserved biological process in different sexes. Further research is needed to uncover their regulatory functions. Interestingly, most of the potential novel miRNAs discovered in this study were expressed at low levels (Table S8C), explaining why they were not discovered in previous efforts and showing the advantage of next-generation sequencing in miRome analysis. This pattern of expression is consistent with the previous results of human studies [60].

In conclusion, we have completed a comprehensive miRNA study of bovine backfat using deep sequencing technology, revealing novel miRNAs and potentially providing new targets for the development of bovine fat tissue.
markers for genetics and breeding. Putative targets for miRn25 and n26 were predicted, and the 61 most significantly different target transcripts related to lipid and fatty acid metabolism were identified using the Blast2GO program. Canonical pathways and gene network analysis revealed that PPARz/RXRz activation and LXR/RXR activation stood out in our gene interaction hierarchy. The results obtained may help in the design of new selection strategies to improve beef quality.

Materials and Methods

Ethics statement

All animal protocols were approved by Institutional Animal Care and Use Committee (IACUC) of Northwest A&F University and Qinbao Animal Husbandry Co., Ltd, respectively. All surgery was performed under sodium pentobarbital anesthesia, and all efforts were made to minimize suffering. Bovine embryos of slaughtered cows were collected from a local slaughterhouse of Xi’an, P.R. China. The adult Qinchuan bovine tissues were obtained from Qinbao Animal Husbandry Co., Ltd, which is a cattle breeding and slaughter corporation in Xi’An, P.R. China.

Tissue collection and high-throughput sequencing

On day 200 (d200), bovine embryos (gestation period 280 days) were collected into sterile physiological saline immediately after removal from the reproductive tract of slaughtered cows at a local abattoir. Fetal age was estimated based on crown-rump length [64]. Bovine tissue samples including the heart, liver, lungs, kidneys, stomach, intestine, muscle, and fat were collected from the adult Chinese Qinchuan cattle. These tissues were snap-frozen in liquid nitrogen and stored at −80°C until use. In this study, two miRNA libraries were constructed. Total RNAs were extracted from the backfat of three fetal and three adult Chinese Qinchuan bovines, which were pooled respectively. Subsequently, low molecular weight RNAs were removed. Then, the flanking sequences that have more than 10 hits to the genome or that match to known potential novel miRNA precursor sequences, unique sequences in the miRBase were considered conserved miRNAs. To discover precursor sequences perfectly matched known bovine miRNAs in miRBase, conserved miRNAs. Only those small RNAs whose mature and precursor sequences perfectly matched known bovine miRNAs in miRBase were considered conserved miRNAs. To discover potential novel miRNA precursor sequences, unique sequences that have more than 10 hits to the genome or that match to known non-coding RNAs were removed. Then, the flanking sequences (150 nt upstream and downstream) of each unique sequence were extracted for secondary structure analysis with Mfold [http://www.bioinfo.rpi.edu/applications/mfold] and evaluated by Mireap [http://sourceforge.net/projects/mireap/]. Specifically, the miRNA candidates that passed Mireap were deemed as highly probable if their corresponding miRNA*s were also found in the small RNA libraries. After prediction, the resulting potential miRNA loci were examined carefully based on the distribution and numbers of small RNAs on the entire precursor regions. Those sequences residing in the stem region of the stem-loop structure and ranging between 20–22 nt with free energy hybridization lower than −20 kcal/mol were considered [65].

MicroRNA expression analysis

A comparison of the known miRNA expression in the two samples was conducted to determine the differentially expressed miRNAs. The samples’ miRNA expression was shown by plotting the log2-ratio figure and scatter plots. The procedures are shown below: (1) Normalize the expression of miRNA in two samples (fetal and adult backfat) to get the expression of transcripts per million. Normalized expression (NE) = Actual miRNA count/Total count of clean reads. (2) Calculate the fold-changes and P-values from the normalized expression. Then, generate the log2-ratio and scatter plots. Fold-change formula: Fold_change = log2 (fetal NE/adult NE). P-value formula: \( p(X|Y) = \left( \frac{1}{N} \right)^{X} \cdot \binom{Y}{X} \cdot \left( \frac{1}{N} \right)^{Y} \left( \frac{1}{N} \right)^{1} \), where \( N \) = Actual miRNA count, \( X \) = fetal NE and \( Y \) = adult NE.

The x and y represent normalized expression levels, and N1 and N2 represent the total count of clean reads of a given miRNA in small RNA libraries of the fetal and adult stages, respectively. Stem-loop real-time reverse transcription polymerase chain reaction (RT-PCR) with SYBR Green was used for the analysis of miRNA expression [66]. Total RNA (1 μg from tested tissues) was converted to cDNA with a RT primer mixture (250 nM) using a PrimeScript® RT reagent Kit with gDNA Eraser (TaKaRa, Dalian, China). The cDNA was then used for the real-time PCR quantification of miRNA using the miRNA-specific primer and the universal primer. The bovine ribosomal protein S18 (RPS18) (GenBank NO. NM_001033614.1) gene was used as an endogenous control. The primers for miRNAs and the control gene are listed in Table S12. Real-time quantitative PCR was performed using a Bio-Rad CFX 96™ Real Time Detection System and SYBR Green PCR Master Mix (TaKaRa, Dalian, China) in a 20 μl reaction. All reactions were carried out in triplicate. The threshold cycle (Ct) was defined as the cycle number at which the fluorescence intensity passed a predetermined threshold. The quantification of each miRNA relative to the RPS18 gene was calculated using the equation: \( N_{2} = 2^{-\Delta \Delta Ct} \).

Target gene prediction

A search for miRNA target genes was performed using an approach described earlier [67,68], and the rules used for target prediction include the following six aspects: (1) no more than four mismatches between the sRNA & target (G-U bases count as 0.5 mismatches); (2) no more than two adjacent mismatches in the miRNA/target duplex; (3) no adjacent mismatches in positions 2~12 of the miRNA/target duplex (5’ of miRNA); (4) no mismatches in positions 10~11 of the miRNA/target duplex; (5) no more than 2,5 mismatches in positions 1~12 of the miRNA/target duplex (5’ of miRNA); (6) the minimum free energy (MFE) of the miRNA/target duplex should be greater than or equal to 75% of the MFE of the miRNA bound to its perfect.
Availability of supporting data

The data set supporting the results of this article is available in the NCBI Gene Expression Omnibus (GEO, http://www.ncbi.nlm.nih.gov/geo/) repository, with access number GSE48569.

Supporting Information

Figure S1 Nucleotide bias of sRNA tags. Note: miRNA nucleotide bias of fetal bovine backfat (A) and adult bovine backfat (B), respectively.

Figure S2 The differential expressions of bovine conserved (A) and novel (B) miRNAs between adult bovine muscle and adult bovine backfat tissues were shown. Note: Expression level (AM): Expression level of adult bovine muscle; Expression level (AF): Expression level of adult bovine backfat. Each point in the figure represents a miRNA. Red points represent miRNAs with fold change > 2, blue points represent miRNAs with 1/2 < fold change < 2, green points represent miRNAs with fold change ≤ 1/2.

Figure S3 The expression of miRNAs in bovine tissues and organs were detected by RT-qPCR.

Figure S4 GO term distribution. Standard configuration of the Blast2GO web application (http://www.blast2go.de) was applied to generate level 2 graphs for GO-term distributions to the biological process.

Table S1 Bovine conserved miRNAs.

Table S2 Summary of base edit in fetal bovine library.

Table S3 Summary of base edit in adult bovine library.

Table S4 End variants of the fetal bovine library identified in this study.

Table S5 Family analysis of known miRNA.

Table S6 The known miRNAs expression profiles between two libraries.

Table S7 Novel miRNAs identified in this study.

Table S8 The known miRNAs expression profiles between adult muscle and adult backfat libraries.

Table S9 Predicted targets for novel miRan25.

Table S10 Predicted targets for novel miRan26.

Table S11 The known miRNAs expression profiles between two libraries.

Table S12 Stem-loop RT-PCR Primers.

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