Research Article

**PPARγ Regulates Genes Involved in Triacylglycerol Synthesis and Secretion in Mammary Gland Epithelial Cells of Dairy Goats**

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To explore the function of PPARγ in the goat mammary gland, we cloned the whole cDNA of the PPARγ gene. Homology alignments revealed that the goat PPARγ gene is conserved among goat, bovine, mouse, and human. Luciferase assays revealed that rosiglitazone enhanced the activity of the PPARγ response element (PPRE) in goat mammary epithelial cells (GMECs). After rosiglitazone (ROSI) treatment of GMECs, there was a significant ($P < 0.05$) increase in the expression of genes related to triacylglycerol synthesis and secretion: LPL, FASN, ACACA, PLIN3, FABP3, PLIN2, PNPLA2, NR1H3, SREBF1, and SCD. The decreases in expression observed after knockdown of PPARγ relative to the control group (Ad-NC) averaged 65%, 52%, 67%, 55%, 65%, 58%, 85%, 43%, 50%, and 24% for SCD, DGAT1, AGPAT6, SREBF1, ACACA, FASN, FABP3, SCAP, ATGL, and PLIN3, respectively. These results provide direct evidence that PPARγ plays a crucial role in regulating the triacylglycerol synthesis and secretion in goat mammary cells and underscore the functional importance of PPARγ in mammary gland tissue during lactation.

1. Introduction

Lactation is a process highly demanding of lipid synthesis and transport. Although peroxisome proliferator-activated receptor γ (PPARγ) is known to promote lipogenesis and adipogenesis in adipose tissue [1], its role in the lactating mammary gland is less clear. Many candidate genes that regulate lipid synthesis have been identified during the lactation cycle [2]. Researchers have evaluated the expression profiles of 54 genes associated with bovine milk fat synthesis through various periods during lactation and built a regulatory network [3]. Their data showed that PPARγ might be the main factor that regulates the nuclear transcription factor, sterol regulatory element-binding transcription factor 1 (SREBF1), which also affects the expression of some fatty acid metabolism genes during lactation [3, 4].

Much data have been published regarding PPARγ’s role in milk fat synthesis in bovine [5–7], while there is a lack of data on its role in the dairy goat. Whether PPARγ also plays the same critical role in regulation of milk fatty acid synthesis during the lactation process in dairy goat remains to be determined. In the present study, we first identified the sequence of PPARγ in dairy goat mammary tissue and evaluated the activity of the PPRE via luciferase assays. Its function in dairy goat mammary epithelial cells (GMECs) was also investigated through the use of the pharmaceutical ligand rosiglitazone (ROSI) and adenovirus-mediated RNA interference.

2. Materials and Methods

2.1. cDNA Cloning. The primers used in the amplification of the goat PPARγ transcript sequence (PPARG) used for cDNA cloning are reported in Table 1. Primers were designed based on the consensus conserved sequences between humans (AB472042) and bovines (BC116098). The PCR reaction was performed with goat mammary epithelial cell cDNA as a template. The cDNA cloning of the 5’ and 3’ UTR was implemented according to the manufacturer's protocols of
the 5′ RACE system Ver.2.0 (Invitrogen, USA) and 3′-full RACE core set Ver.2.0 (Takara, Japan). The nested gene-specific primers for PPARγ, designed based on its open read fragment (ORF), were used for 3′ RACE. Similarly, the nested gene-specific primers (Table 1) were also designed for 5′ RACE. All the PCR fragments were cloned into pMD-19T plasmid vectors (Takara, Japan) and then sequenced at a commercial facility (Invitrogen, Shanghai, China). The PPARγ protein structure was predicted using PHYRE2 (http://www.sbg.bio.ic.ac.uk/phyre2/html/page.cgi?id=index).

2.2. Vector Construction and shRNA. The luciferase vector (pGL3-basic) containing three copies of PPRE was designed using the WI siRNA Selection Program (http://sirna.wi.mit.edu/home.php) and BLOCK-iT RNAi Designer (http://rniaidesigner.invitrogen.com/rnaiexpress/) using the goat PPARγ gene sequence (HQ589347.1). We selected the highest-ranked shRNA sequences. Additionally, a BLAST search against all EST sequences in GenBank was performed to ensure that the selected sequences were specific for goat PPARγ. Meanwhile, those sequences were selected and synthesized at a commercial facility (Invitrogen, Shanghai, China) with BamHI and Xho I restriction sites suitable for the cloning process (see Table 2). Lastly, three shRNA were generated by heat treatment annealing and constructed into pENTR/CMV-GFP/U6-shRNA. The CDS of PPARγ was subcloned into the pDsRed1-C1 plasmid vector between the Xho I and EcoR I restriction sites to generate pDsRed1-C1-PPARγ.

2.3. Cell Culture and Treatments. Goat mammary epithelial cells isolated from a Xinong Saanen goat at peak lactation [9] were allowed to grow in 60 mm culture dishes (NUNC, Denmark) in DMEM/F12 medium (HyClone, China). Routine cultures were incubated at 37°C in 5% CO2 and air. Culture medium was changed every 24 h. Medium was composed of DMEM/F12 with insulin (5 mg/L, Sigma, USA), hydrocortisone (5 mg/L, Sigma, USA), penicillin/streptomycin (10 kU/L, Harbin Pharmaceutical Group, China), epidermal growth factor (1 mg/L, Sigma, USA), and fetal bovine serum (10%, Gibco, USA). ROSI (BioVision, USA) was resuspended in DMSO (Sigma, USA) at a concentration of 50 mmol/L. Cells cultured in 60 mm culture dishes and subcultured at 90% confluence were treated with 50 μmol/L ROSI and harvested at 0, 12, and 24 h after treatment to extract total RNA. The 293A cells for preliminary testing of shRNA and generating recombinant adenovirus were cultured in the basal medium containing 10% fetal bovine serum and 90% DMEM (Gibco, USA).

2.4. Preliminary Screening of shRNA Sequences. In order to get the most effective shRNAs for targeting PPARγ gene, an experiment was done as follows. 293A cells at 80% confluence in 12 plates were transiently transfected with 1.0 μg of three pENTR/CMV-GFP/U6-shRNAs with pDsRed1-C1-PPARγ at a ratio of 3:2 using FuGENE HD Transfection Reagent (Roche, Switzerland). The pDsRed1-C1-PPARγ vector also was transfected alone as a control in the same amount as above. All the steps were performed in accordance with the manufacturer's protocol. The GFP fluorescence was monitored by using a Leica fluorescent microscope (DMI14000B, Germany).

2.5. Adenovirus Generation. shRNA expression cassettes with an EGF reporter gene in the pENTR vector were switched into an adenoviral vector (pAd/PL-DEST) using the Gateway technique (Invitrogen, USA) to generate pAd-shRNA vectors. Pac I linearized adenoviral plasmids were transfected into 293A cells to generate the adenovirus. About

### Table 1: Primer pairs used in PCR for amplification of goat PPARγ from mammary cDNA.

| Name of fragment | Sequence | Product length |
|------------------|----------|----------------|
| PPARγ CDS        | Forward: 5' -ATGGTTGACACAGAGATGCGG-3' 1413 bp  |
|                  | Reversal: 5' -GTAGATTCTCTGTAGAAGTGGTG-3' |
| PPARγ 3′ RACE    | Outer: 5′-AAGTAACTCTCTCTAAAAATACGGCG-3′  516 bp |
|                  | Inner: 5’-CCAGAAAAATGACGGACCTACCGAGA-3′ 160 bp |
| PPARγ 5′ RACE    | GSP1: 5′-CGGTGTATTGTCGTGCTTTTC-3′  750 bp |
|                  | GSP2: 5′-GATACAGGGCTCCAATTTGATTGC-3′ 260 bp |

### Table 2: Characteristics of shRNA used in the experiment.

| Name of shRNA | Sequence |
|---------------|----------|
| sh500-sense   | 5′-GATCCGGAGGAGCATGATGAAAGATCGTAGTCCTC-3′ |
| sh500-antisense| 5′-TCGAGAAAAAGAGAGGAGCATGATGAAAGATCGTAGTCCTGCC-3′ |
| Sh614-sense   | 5′-GATCGGGATGTCTCATAACGCCATGACTGAGTCTGGCTCC-3′ |
| sh614-antisense| 5′-TCGAGAAAAAGAGAGGAGCATGATGAAAGATCGTAGTCCTGCC-3′ |
| Sh1006-sense  | 5′-GATCGGGTTTGAGACTTGACTGACTGAGTCTGGCTCC-3′ |
| sh1006-antisense| 5′-TCGAGAAAAAGAGAGGAGCATGATGAAAGATCGTAGTCCTGCC-3′ |

Three shRNAs (numbers stand for their position in cDNA) were designed, and each shRNA was added with restriction sites BamH I and Xho I. The loop domain (lower-case nucleotides) contained a Scal I site.
### Table 3: Characteristics of primer pairs used, amplicon length, and efficiency of reaction in the RT-qPCR.

| Accession | Gene | Primer sequence (5′ to 3′) | Product length (bp) | Efficiency |
|-----------|------|-----------------------------|---------------------|------------|
| JN236219.1 | ACACA | Forward: CTCACAACCTCAACCTACGG  
Reversal: GGGAATACACAGAAAGGCAGCC | 171 | 2.09 |
| JI861797.1 | AGPAT6 | Forward: AAGCAAGTTGCCCATCCTCA  
Reversal: AAACGTTGGCTGAATTTCG | 101 | 2.17 |
| X91503 | CD36 | Forward: GTACAGATGCAGCCTCATTTCA  
Reversal: TGGACCTGCAAATATCAGAGGA | 81 | 2.18 |
| DQ380249.1 | DGAT1 | Forward: CCACCTGGGACCTGAGGTGTC  
Reversal: GCATCACCACACACCAATTCA | 101 | 2.11 |
| NM_001009350 | FABP3 | Forward: GATGAGACCACGGCAGATG  
Reversal: GTCAACTATTTCCCGCACAAG | 120 | 2.14 |
| DQ915966.3 | FASN | Forward: GGCTCCACCACCGTGTTCCA  
Reversal: GCTCTGCTGGGCCTGCAGCTG | 226 | 2.13 |
| AJ431207 | GAPDH | Forward: GAAGTTCCACGGCACAG  
Reversal: GGTTCACGCCCATCACAA | 249 | 2.16 |
| DQ997818 | LPL | Forward: AGGACCTGGGACCTGAGGTGTC  
Reversal: GCATCACCACACACCAATTCA | 169 | 2.18 |
| GU332719 | NR1H3 | Forward: CATCAACCCCCATCTCTGGATT  
Reversal: CAGGGGCTCCACATATGTGT | 163 | 2.13 |
| HQ846826 | PLIN2 | Forward: TACGGATGATCAGATGATGAT  
Reversal: CAGGGGCTCCACATATGTGT | 203 | 2.16 |
| HQ846827 | PLIN3 | Forward: GTGGAGGGGTCAGGAGAAA  
Reversal: TCACGGAAATCGGCGAGT | 170 | 1.13 |
| GQ918145 | PNPLA2 | Forward: GGAGCTTATCCAGGCCAATG  
Reversal: TGGCGCCGAGCTGTCACTCT | 226 | 2.24 |
| HQ589347.1 | PPARG | Forward: CCTTCACCACCGTGTTGACTCT  
Reversal: GATACAGGCTCCACTTTGATTGC | 145 | 2.21 |
| DV935188 | SCAP | Forward: CCATGTGCACTTCAAGGAGGA  
Reversal: TGGCGCCGAGCTGTCACTCT | 108 | 2.10 |
| GU947654 | SCD | Forward: CCTTGCCGTGAGGAGGACG  
Reversal: GTCCGGGATACATCAAGGACAT | 257 | 2.10 |
| HM443643.1 | SREBF1 | Forward: CTGTCGACCCGACGAGCAG  
Reversal: GTAGGGCGGGTCAAACAGG | 81 | 2.20 |

Annealing temperature for all primers in this table is 60°C.

ACACA, acetyl-coenzyme A carboxylase alpha; AGPAT6, l-acylglycerol-3-phosphate O-acyltransferase 6; CD36, thrombospondin receptor; DGAT1, diacylglycerol acyl transferase 1; FABP3, fatty acid binding protein 3; FASN, fatty acid synthase; GAPDH, glyceraldehyde-3-phosphate dehydrogenase; LPL, Lipoprotein lipase; NR1H3, liver X receptor α; PLIN2, perilipin2; PLIN3, perilipin3; PNPLA2, patatin-like phospholipase domain containing 2; PPARG, peroxisome proliferator-activated receptor γ; SCAP, cleavage activating protein; SCD, stearoyl-CoA desaturase; SREBF1, Sterol regulatory element-binding transcription factor 1.

#The primer sequences are from bovine.

8 to 10 days after transfection, the recombinant virus was collected and subjected to two rounds of amplification in 293A cells. The viral titers were determined in transduced 293A cells through GFP expression as previously described [10–12].

### 2.6. Luciferase Assays.

To assess the degree of PPARγ activation, goat mammary epithelial cells at 80% confluence in 96-well plates were transiently transfected with 0.08 μg of PPRE×3-Luc reporter plasmid along with a Renilla vector (pRL-TK) as a control using the FuGENE HD transfection reagent at a ratio of 25:1. After a 24 h recovery period in medium, cells were treated with 0, 10, 25, 50, and 100 μmol/L ROSI. Forty-eight hours later, cells were harvested and lysates were made using reporter lysis buffer (Promega, USA) according to the manufacturer’s instructions. Luciferase activity in the cell extract was determined using luciferase assay buffer and luciferase assay substrate according to the manufacturer’s protocol (Promega, USA) in a luminometer (BHP9504, China).

### 2.7. Adenovirus Transduction.

Goat mammary epithelial cells at 70–80% confluence were transduced with adenovirus supernatant at a multiplicity of infection (MOI) of 200. The medium was replaced with fresh medium 6 h later. The shRNA negative control adenovirus (Ad-NC) was used as a control. Cells were harvested 48 h after transduction.

### 2.8. RNA Extraction and Real-Time RT-PCR (qPCR).

Total RNA was extracted from cells using RNAprep pure cell kit (Tiangen, China). The first-strand cDNA of different treatments was synthesized from 0.5 μg of purified total RNA using the PrimeScript RT kit (Takara, Japan) according to the manufacturer's instructions. Sufficient cDNA was prepared to
run all the selected genes (Table 3). Primers were designed to span exon-exon boundaries according to BLAST against bovine genome in order to avoid amplification of genomic DNA using Primer 5.0 software. The specificity of the primers was tested using the same protocol as for qPCR in a simple thermocycler (S1000, Bio-rad, USA), and the PCR product was run in a 15μg/L agarose gel. In addition, a dissociation protocol was performed in the RT-qPCR. Only primers with a single band on the agarose gel, a unique peak in the dissociation curve after the RT-qPCR, and devoid of primer-dimers were selected. The efficiency of each primer pair was tested using a standard curve as previously described [3]. All the amplicons were sequenced in order to assess the right amplified genes. Characteristics of all primers used in the RT-qPCR reaction are described in Table 3. RT-qPCR reactions were performed according to the manufacturer’s instructions (SYBR Premix Ex Taq II, Perfect Real Time, Takara, Japan).

Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was selected as an internal control gene [13]. Although we did not verify additional genes as internal controls, GAPDH was used partly because it has been used previously in a goat mammary tissue study [13], and also because it has been widely used as the sole control gene in bovine cell studies [14]. However, we understand the limitation of using a single internal control gene because more reliable data requires the verification and use of at least 3 internal controls [15].

2.9. Western Blot. Whole cell proteins were extracted with RIPA buffer (Solarbio, China) supplemented with PMSF (Pierce, USA). Western blotting was performed using the following primary and secondary antibodies: anti-PPARγ (Abcam, ab19481, Hong Kong, 1:400) and goat anti-rabbit IgG (Tiangen, China, 1:1000). All antibodies were used according to the manufacturer’s recommendations. Signals were detected using the chemiluminescent ECL Western blot detection system (Pierce, USA).

2.10. Statistical Analysis. Each treatment was replicated 3 times, and results are expressed as mean ± SD. Data of RT-qPCR was analyzed relative to the control using the 2−ΔΔCt method. The statistical significance for ROSI treatment was determined by the ANOVA test using SPSS 19.0 software. Treatment means for shRNA interference were separated using Fisher’s least significant difference pair-wise comparisons. Significance was declared at P < 0.05.

3. Results and Discussion

3.1. Molecular Cloning and Sequence Analysis of Dairy Goat PPARγ. PPARγ is a member of the nuclear hormone receptor superfamily of transcription factors. It has been fully confirmed in humans and mice that PPARγ directly regulates adipose cell proliferation, maturation, and differentiation [16, 17]. A potential role of PPARγ in controlling milk fat synthesis also has been reported in bovine due to the increase of its expression between pregnancy and lactation [2] and the increase in expression of genes involved in milk fat synthesis after activation with ROSI [5]. However, its role, if any, on milk fat synthesis in the mammary gland of the goat remains relatively unknown. In this study, we cloned the dairy goat PPARγ CDS and then used 5’RACE and 3’RACE procedures to obtain the full-length cDNA. The whole goat PPARγ gene contains a 5’UTR of 114bp, an ORF of 1428bp, and a 3’UTR 215bp. Homology alignment (BLASTN) revealed that the dairy goat PPARγ gene (HQ589347.1) shares 90%, 89%, 98% and 98% identity with human (AB472042), mouse (NM_0011273301.1), sheep (NM_001100921), and bovine (BC116098), respectively. Figure I(a) shows their genetic relationship. The structure prediction using online software revealed that there are two zinc finger structures and a ligand binding domain in the dairy goat PPARγ protein (Figure I(b)). It was also predicted (PredictNLS online software) that the nuclear localization signal sequence (-KKSRNKC-) of the dairy goat PPARγ gene does not exist in either ends of the peptide chain, but it is present in the protein internal compartment.

3.2. A PPARγ Ligand Enhanced Activity of PPARγ Response Element in GMECs. PPARγ is a ligand-dependent nuclear transcription factor, and several unsaturated fatty acids in mammalian tissue are its natural ligands [18]. Binding of ligands to the PPARγ ligand binding domain causes conformational changes in the receptor [16, 19]. Once activated, PPARγ forms a heterodimeric complex with retinoid X receptor (RXR) and binds to thePPRE upstream of target genes [8]. In the present study, dairy goat mammary epithelial cells were incubated with rosiglitazone, a chemosynthetic ligand, which has a high affinity for PPARγ and enhanced its activity. As shown in Figure 2, treatment with ROSI caused an activation of PPARγ in GMECs. The luciferase levels between the treatment group and the control group (treatment with 0μmol/L ROSI) were statistically significant (P < 0.05). Data also indicated that the activation of the PPARγ by ROSI reached a peak at 50μmol/L dose.

3.3. Activation of PPARγ by ROSI Affects Expression of Genes Related to Triacylglycerol Synthesis and Lipid Droplets in GMECs. Genes related to de novo fatty acid synthesis (acyl-coenzyme A carboxylase alpha (ACACA), fatty acid synthase (FASN)), desaturation (Stearoyl-CoA desaturase (SCD)), TAG synthesis (Diacylglycerol acyl transferase 1, (DGAT1)) and other genes including fatty acid binding protein 3 (FABP3) and Perilipin2 (PLIN2) were upregulated in adipose tissue of rats [20], humans [21], and bovine mammary epithelial cells [5] treated with ROSI. As summarized in Figure 3, treatment with ROSI increased the expression of ACACA, FASN, SCD, FABP, LPL, and also those associated with lipid droplet formation and hydrolysis (PLIN2 and patatin-like phospholipase domain containing 2, PNPLA2), and transcription regulators (SREBF1; liver X receptor α, NRIH3) (Figure 3). The significant (P < 0.05) increase in gene expression suggests that these genes are putative PPARγ target genes in goat mammary gland. In a previous study, the expression of genes associated with long-chain fatty acid uptake or intracellular activation and transport, including LPL, was not affected by ROSI treatment of bovine mammary...
cells for 12 h [5]; however, our results revealed that LPL was upregulated significantly with ROSI treatment but only after 24 h. These contrasting responses may be related at least in part with inherent species differences in the regulatory mechanism via PPARγ [5, 22].

3.4. Preliminary shRNA Screening and Adenovirus Generation. The GFP protein on the pENTR/CMV-GFP/U6-shRNA vector was used to assess the efficacy of transduction via intensity of green fluorescence inside the cells. The 293A cells were either transfected with only the pDsRed1-C1-PPARγ construct (red fluorescent cells) or cotransfected with both constructs. Once the shRNA enters the cell, if specific for PPARγ, it would enhance pDsRed1-C1-PPARγ construct with a concomitant reduction of red fluorescence. In this way, the shRNA efficacy in knocking down PPARγ was assessed by the disappearance of red fluorescence in the cells. As shown in Figure 4, sh1006 and sh614 were more efficient than sh500 to silence PPARγ (Figures 4(b2) and 4(c2)). There was more dsRED fusion protein being coded and detected in the sh500 group (Figure 4(a2)), indicating that sh500 had weaker silencing effect on goat PPARG. This was probably also due to the lower transfection observed for the sh500 construct (Figure 4(a3)).

Although the approach depicted in Figure 4 is not quantitative, it represents a relatively easy way to screen efficient shRNAs.
According to the results of the preliminary screening, sh1006 and sh614 were selected to generate adenovirus Ad-sh614 and Ad-sh1006. Judging by the RT-qPCR and western blot analysis (Figure 5), compared with Ad-sh614 (about 20%), the Ad-sh1006 (about 60%) was more efficient in knocking down goat PPARγ.

3.5. Knockdown of Goat PPARγ in GMECs Affects Expression of Genes Involved in Triacylglycerol Synthesis and Lipid Droplet Formation in GMECs. Based on the above results, the Ad-sh1006 was selected to block expression of PPARγ in GMECs, and expression analysis of genes known to be involved in milk fat synthesis and lipid droplet formation was evaluated (Figure 6). Results demonstrated that FASN (−58%), ACACA (−65%), and SCD (−65%) decreased significantly after PPARγ knockdown (Figure 6(a)). With the exception
Figure 4: Efficacy screening of the three designed shRNA via images analysis. pDsRed1-C1-PPARγ vector was transfected as a control ((a1), (b1), and (c1)). The three tested shRNA (sh500, sh614, and sh1006) as pENTR/CMV-GFP/U6-shRNA construct were cotransfected with pDsRed1-C1-PPARγ vector. The transduction efficiency was estimated by the level of green fluorescent protein (GFP) expression ((a3), (b3) and (c3)). Shown are representative images of the PPARγ expression (in red) after a 48 h cotransfection. (a1), (b1), and (c1) show high transfection and expression of PPARγ construct vector. (a2), (b2), and (c2) show reduction of PPARγ expression after addition of shRNA construct, while (a3), (b3), and (c3) show efficacy of shRNA transfection as shown by the green color (i.e., GFP). Images were obtained by a fluorescence microscope (Leica, DMI4000B, Germany) at 100x magnification. The images clearly show that the sh1006 had the highest effect on PPARγ vector expression (c2).

Figure 5: Efficacy screening of the two designed shRNA via RT-qPCR and western blot. The efficiency of Ad-sh614 and Ad-sh1006 (transduced with two adenoviruses at 200 multiplicity of infection for 48 h) in decreasing PPARG expression in dairy goat mammary epithelia cells was assessed by RT-qPCR (a) and western blot (b). The data revealed that Ad-sh1006 had the highest knockdown of PPARγ transcript and protein; thus, it was used in the subsequent experiments.
Figure 6: Effect of PPARγ knockdown on genes coding for proteins involved in milk fat synthesis in GMECs. The expression of genes related to fatty acid synthesis (a), cellular fatty acid uptake (b), triacylglycerol synthesis (c), lipid droplet formation and triacylglycerol hydrolysis (d), and transcriptional regulation (e) was assessed in goat epithelial cells (GMECs) after transduction with Ad-sh1006 at 200 MOI for 48 h. The data represent the mean ± SD of cells transfected with control (Ad-NC) or Ad-sh1006 vector in triplicate per experiment. \(^{b}P < 0.05\) versus the control group.
of SCD, those data are in agreement with observations in bovine [5] and suggest that PPARγ regulates de novo fatty acid synthesis and desaturation in goat mammary cells.

In bovine mammary cells, SREBF1 has attracted much attention because of its regulation of FASN and SCD expression and the major role played in milk fat synthesis [6, 23]. PPARγ indirectly regulates SREBP1 protein activity through regulation of the expression of insulin-induced gene 1 (INSIG1) and directly regulates SREBF1 expression in adipose cells of mice [4]. We observed that the expression of SREBF1 and SCAP decreased by 50% and 43% after knockdown of PPARγ (Figure 6(e)). The mRNA of NRIH3 gene also was reduced by 75% when PPARγ was knocked down (Figure 6(e)). Our data agree to a large extent with a previous bovine study, where an increase of SREBF1 expression after ROSI treatment was observed [5]. We speculate that there might be two different signaling networks regulating de novo fatty acid synthesis in ruminant mammary cells. One pathway is under direct regulation of PPARγ and encompasses genes such as LPL, NRIH3, and FABP3 (Figures 6(b) and 6(e)); another is under indirect regulation of PPARγ through SREBF1 and NRIH3 (Figure 6(e)) which would, in turn, participate in upregulation of the transcription of FASN and ACACA [23–25]. Regardless of the specific mechanism, our data support the previous hypothetical milk fat synthesis transcriptional networks proposed for bovine mammary [3].

In agreement with that previous proposal, our data support a complex regulatory network that controls mammary triacylglycerol synthesis in goat mammary cells such that several protein factors serve as putative checkpoints to regulate milk fat synthesis. PPARγ appears to be one of those factors in dairy goats.

PPARγ plays multifaceted roles in the regulation of triacylglycerol synthesis and secretion besides the de novo synthesis of fatty acids. As an adiposity factor, PPARγ is able to regulate triacylglycerol synthesis and deposition and then dominate the process of differentiation of fat cells [26]. In the present study, the mRNA expression of genes related to triacylglycerol synthesis DGAT1 (~52%) and AGPAT6 (~67%) decreased greatly after infection with Ad-sh1006 (Figure 6(b)), which suggests that PPARγ regulates triacylglycerol synthesis in mammary cells as in fat cells.

Triacylglycerols are deposited in fat cells, while in the mammary cells they are secreted in the form of lipid droplets in milk. To investigate the role of PPARγ in transcription of milk fat globule protein genes, we measured the mRNA expression of PLIN2, PLIN3, and PNPLA2 after PPARγ knockdown (Figure 6(d)). The expression of PLIN2 was largely induced while the expression of PLIN3 and PNPLA2 decreased approximately 24% and 50%, respectively, in cells transfected with Ad-sh1006, while it is not extremely for PLIN3. Previous data from humans [27] indicated that there is a PPRE on the promoter of the PLIN2 gene; thus, it is considered as a downstream target and would be decreased after PPARγ knockdown. However, our data showed that the expression of PLIN2 had an unexpected increase. Such response might have been caused by compensatory effects of other unidentified transcription factors.

Other data also support the evidence [28] that PPARγ could affect not only the genes related to fatty acid transport, but also genes that control triacylglycerol hydrolysis in goat mammary cells (Figure 6(e)). For instance, expression of PNPLA2 is significantly increased during lactation in bovine mammary tissue [29]. However, judging by differences in milk fatty acid profiles between goat milk and bovine milk [30], goat mammary lipid synthesis differs in some respects from bovine. From a mechanistic standpoint the upregulation of PNPLA2 after PPARγ activation may be functionally related with the unique characteristics of goat milk.

Our data showed that even if there is great similarity between two ruminant dairy species such as goat and cow [5], there are still some inherent differences between them. Such differences may at least in part be caused by different target genes of PPARγ in each species. Attempts to compare in vitro data among studies performed in different laboratories are obviously challenging because of differences in cell culture conditions (e.g., culture medium, absence of prolactin in our study and not in bovine [31]) and also different protocols. The comparisons of data from the present study with data generated in bovine mammary [31] are likely also slanted because of the use in the present study of GAPDH as the only internal control for RT-qPCR normalization versus multiple genes used in the bovine study.

### 4. Conclusions

In the present studies, we cloned the PPARγ gene in dairy goat mammary gland and explored its function in vitro. As proposed in bovine mammary gland, PPARγ plays a multifaceted role in regulating the overall process of fatty acid and triacylglycerol synthesis and secretion. Our overall data indicate that PPARγ in goat mammary plays a role in controlling milk fat synthesis directly or via the activation of the transcription regulators SREBF1 and NRIH3. Together, our data provide strong evidence that PPARγ is the key regulator of milk fat synthesis in ruminants. Hence, controlling PPARγ activation may prove useful in regulating milk fat production in the lactating dairy goat.

### Conflict of Interests

The authors declare that they have no conflict of interests.

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