In vitro Silencing of Acetyl-CoA Carboxylase beta (ACACB) Gene Reduces Cholesterol Synthesis in Knockdown Chicken Myoblast Cells

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Abstract

The poultry industry provides cost-effective, healthy, and protein-enriched food for the growing population and achieving the nutritional security to the country. Excessive abdominal and subcutaneous fat deposition is one of the major setbacks to the poultry industry that reduces carcass yield and feed efficiency. In chicken abdominal fat constitutes 20% of total body fat which make up 2–3% of live weight of the bird. In fatty acid metabolism, acetyl-CoA Carboxylase (ACC) is one of the key enzymes with two isoforms i.e. ACACA and ACACB each of which plays a different role. In chicken, ACACB is involved in the β-oxidation of fatty acids and thereby potentially regulating the quality of meat and egg. The RNAi strategy is widely used for silencing the target gene expression. In this study, we designed five shRNA constructs and identified the most efficient shRNA molecule for silencing the ACACB gene under in vitro chicken embryo myoblast (CEM) primary cell culture system. After knocking down the ACACB gene, for understanding how fatty acid metabolism is regulated, we tracked the expression of key fatty acid metabolism genes like ACACA, FASN, SCD, ELOVL2, and CPT1. Also, checked the expression of immune response genes like IFNA, IFNB, and BLB1 in control as well as ACACB knockdown myoblast cells and observed no significant difference. We observed the down-regulation of key fatty acid metabolism genes along with ACACB, which may leads to the less fat accumulation in CEM cells. We also estimated the cholesterol and triglycerides in control and ACACB knockdown myoblast cells and found a significant difference between control and the knockdown cells. In vitro knockdown of the ACACB gene in a cell culture system by a short hairpin RNA (shRNA) expressing construct would help to produce a knockdown chicken with reduced fat deposition.

Introduction

The major objective of poultry production is to provide cheaper, safe, healthy, and protein-enriched food for the growing population thereby achieving the nutritional security to the country. Due to the adoption of improved nutritional strategies and technological innovations creating better rearing conditions, the poultry industry has grown greatly in recent years by making the chicken attaining a finishing body weight of about 2 kg in a very short period. On the contrary, apart from resulting in excessive body fat deposition, this rapid growth rate leads to the high incidence of metabolic diseases and skeletal disorders with increased mortality. In broiler chicken abdominal fat constitutes 20% of total body fat and excessive body fat has been recognized as a major problem in the poultry industry, which make up 2–3% of the total live weight of the broiler chicken (Cahaner et al. 1986; Butterwith 1989; Crespo and Esteve-Garcia 2002; Haro 2005; Leenstra 1986). Fat has limited value both to the poultry producer and consumer. Hence, excessive abdominal and subcutaneous fat deposition is one of the major setbacks to the poultry industry that reduces carcass yield, feed efficiency and ultimately causing consumers non-acceptance of meat (Lippens 2003; Jennen 2004). These negative viewpoints are of significant concern for the rancher and processor as they can bring considerable economic losses (Leenstra 1986; Pym 1987; Griffin 1996; Zubair and Leeson 1996; Buys et al. 1998; Buyse 1999; Havenstein et al. 2003; Nikolova et al. 2007). Moreover, in recent years, preference for leaner meat has been increasing as the consumers are becoming more sensitive about the positive correlation between intake of fatty substances and onset of cardiovascular diseases (Leeson and Summers 1980; Pym 1987; Cable and Waldroup 1990).

In avian species, major site of fat deposition, the abdominal fat pad is the prime indicator of total body fat content (Becker et al. 1979; Thomas et al. 1983). The fat deposition in body tissues is the net result of absorption, de novo synthesis, and β-oxidation of fatty acids (Saadoun and Leclercq 1986). As most of the traditional dietary approaches like restricted feeding techniques to combat excessive fattening in commercial broilers have resulted in decreased genetic potential for weight gain and failed in cost perspective, it would be more appropriate to emphasize the molecular regulation of fatty acid metabolism (Dunnington et al. 1986). In fatty acid metabolism, Acetyl-CoA Carboxylase (ACO) is the key enzyme with two isoforms, ACACA and ACACB each of which plays a different role concerning fat metabolism. The ACACA and ACACB catalyzes the carboxylation of acetyl-CoA to malonyl-CoA and malonyl-CoA generated via ACACA isofom mainly provides carbon units for fatty acid synthesis in lipogenic tissues like liver and adipose tissue whereas malonyl-CoA produced by ACACB isofom regulates fat oxidation by inhibiting carnitine palmitoyl transferase-1 (CPT-1), which controls the entry of long-chain fatty acids into the mitochondrial site of oxidation in nonlipogenic tissues like the heart and skeletal muscles (Abu-Elheiga 2000; Abu-Elheiga 2001). Hence, inhibition of ACACB may lead to reduced fat deposition by increasing fatty acid oxidation. Therefore, we
can reduce the expression of ACACB for improving poultry meat in terms of quality (leaner meat) as well as quantity (weight gain). Feeding experiments on mice revealed that, apart from consuming more food, ACACB knockout mice had lower body fat than their wild-type counterparts and were protected from high fat-induced obesity (Abu-Elheiga 2001; Abu-Elheiga et al. 2005; Oh et al. 2005; Abu-Elheiga et al. 2003). If we knockout the ACACB gene, it may result in deleterious or undesired outcomes, so RNAi-based gene knockdown provides a potential alternative to gene knockout technology. Transgenic RNAi mice showed a gene knockdown phenotype that was functionally similar to gene knockout (Hemmann et al. 2003; Wang et al. 2010). Therefore, down-regulation of ACACB gene expression is a suitable approach to reduce the fat deposition by increased fatty acid oxidation. RNAi has been a standard method in both cultured cells and various model organisms for the controlled down-regulation of gene expression (Tripathi et al. 2012).

In chicken, Acetyl CoA carboxylase beta (ACACB) gene is involved in fatty acid metabolism and thereby, potentially regulating the quality of meat and egg. It is considered that in vitro knockdown of the ACACB gene in cell culture system by developing a short hairpin RNA (shRNA) expressing construct would help, in devising suitable in vivo strategies for knocking down of the gene. This, in turn, might help to produce a knockdown chicken with reduced fat deposition. The present investigation was designed with the objective of silencing ACACB gene with potential shRNA molecule and its effect on other fat synthesis genes under in vitro cell culture system.

Materials

Animals

The experiment was conducted in control broiler chicken line maintained at the Institute farm, ICAR-Directorate of Poultry Research, Hyderabad, India. The fertile eggs were collected from this chicken line and incubated in the incubator for 9 days at 98-100°C with 78–80% relative humidity and turning 6 times a day for embryo development from which embryonic muscle tissues were collected for chicken embryo myoblast cell culture. The entire study was approved by the Institute Animal Ethics committee (IAEC) and Institute Bio-safety Committee (IBSC) of ICAR-Directorate of Poultry Research, Hyderabad, India. All the bio-safety guidelines of IBSC were followed while conducting the experiments.

Designing and cloning shRNA molecules

BLOCK-iT RNAi Designer (https://rnadesigner.thermosher.com/rnaiexpress/) is an online tool that was used to build five separate shRNA sequences that target the ACACB gene's open reading frame (ORF) using Tuschl's motif pattern. The siRNA sequences were then transformed into shRNA with CGAA as the stem-loop sequence and sense loop antisense as the strand orientation. The final shRNA has a 4-nucleotide 5’ overhang (CACC or AAAA) for directional cloning of the ds oligos encoding the shRNA of interest (Table 1). The IDT manufacturers (Integrated DNA Technologies, USA) synthesized the oligos, and scrambled shRNA oligos (lac z) were supplied by the Invitrogen manufacturers (Invitrogen, USA) used in this experiment. The integrity of the ds oligos was checked by loading 5µl annealed ds oligo (500 nM stock) into 4% agarose gel. Both annealed ds oligos (50 bp) and remaining unannealed ss oligos (25 bp) were observed in the gel electrophoresis (Fig. 1). The annealed oligos were cloned into pENTR™/U6 vector (Invitrogen, USA) according to the manual instructions. The transformed recombinant colonies on LB agar plates were screened by using forward (U6: 5’-GGACTATCATATGCTTACCG-3’) and reverse (M13: 5’-CAGGAA ACAGCTATGAC-3’) primers for checking the presence or absence of shRNA inserts (Fig. 1). The plasmid obtained from each pENTR™/U6 entry construct was sequenced to confirm the sequence and orientation of the ds oligos insert.
Table 1
List of top and bottom strand sequences ACACB designed shRNA. Nucleotides in bold letters creates over hangs for cloning in the pENTR™/U6 vector.

| shRNA ID          | Nucleotide sequence (5’ to 3’)                                                                 |
|-------------------|-------------------------------------------------------------------------------------------------|
| Scrambled shRNA top | CACGCCTACACAAATCAGCGATTTCGAAAAATCCTGGATTGTGATG                                              |
| Scrambled shRNA bottom | AAAACTACACAAATCAGCGATTTTCGAAAAATCCTGGATTGTGATG                                                |
| shRNA 1 top       | CACCGGACAACCTCCTCTGATGATGACGAATCATCATCAGAGGATTGTGTC                                          |
| shRNA 1 bottom    | AAAAGGCAACCTCCTCTGATGATGATGGTAGCTATCATCAGAGGATTGTGTC                                          |
| shRNA 2 top       | CACCGGATACCTCCATCTGCTTACGAAATGAGCATGAGGATGATGATGTC                                            |
| shRNA 2 bottom    | AAAAGGCAATGCTTACGAAATGAGCATGAGGATGATGATGTC                                                   |
| shRNA 3 top       | CACCGGCTGGGACCATGTTAATTGAGGAATCGATCACCAGGATGATGATGATGTC                                       |
| shRNA 3 bottom    | AAAAGGCTGTGGGACCATGTTAATTGAGGATGAGGATGATGATGATGTC                                           |
| shRNA 4 top       | CACCGGCTCGGAAACAGCGTTTCTCGAAAGAAACGTGATGATGATGATGATGTC                                       |
| shRNA 4 bottom    | AAAAGGCTCGGAAACAGCGTTTCTCGAAAGAAACGTGATGATGATGATGATGTC                                      |
| shRNA 5 top       | CACCGGACACATCTGGGAAGGATGACGAATGATGCTTTTCCCATGAGGATGATGTC                                    |
| shRNA 5 bottom    | AAAAGGACACATCTGGGAAGGATGACGAATGATGCTTTTCCCATGAGGATGATGTC                                   |

Prediction of secondary structure and thermodynamic properties of ACACB shRNA molecules

The RNA fold program of the Vienna RNA web service version 2.0 (http://rna.tbi.univie.ac.at/cgi-bin/RNAWebSuite/RNAfold.cgi) was used for the prediction of the secondary structures of all five shRNA molecules of chicken ACACB antisense and sense strands (Fig. S1). Further, secondary structures of chicken ACACB mRNA were predicted using a web server Mfold 2.3 version (http://www.unafold.org/mfold/applications/rna-folding-form-v2.php) (Fig. S2).

The Oligowalk software in the RNA structure version was used to determine the thermodynamic properties regulating each shRNA molecule’s binding affinity to its mRNA target region were envisaged (http://rna.urmc.rochester.edu/cgi-bin/server_exe/oligowalk/ oligowalk_form.cgi) (Table 2). Hence, the following parameters were taken into account: $\Delta G_{\text{overall}}$ (Overall Gibbs free energy change): The net energy ($\Delta G$ in kcal/mol) resulting due to binding of oligos to the target site when all energy contributions are considered which includes target structure breaking energy and oligo self-structure energy. The more negative value of $\Delta G$ indicates a more stable duplex. $\Delta G_{\text{duplex}}$: It measures the oligo target binding affinity. A more negative value of $\Delta G_{\text{duplex}}$ indicates more stability of the duplex and vice versa. $\Delta G_{\text{break-target}}$ (disruption energy): The energy cost for disrupting base pairs in mRNA target region so that the binding site becomes single-stranded and completely accessible. A more negative value denotes that the binding site is less accessible. $\Delta G_{\text{intra-oligomer}}$ and $\Delta G_{\text{inter-oligomer}}$: The negative $\Delta G$ of stable structures was greater than that of unstable structures. The free energy changes/differences caused by unimolecular and bimolecular siRNA foldings. $\Delta \Delta G_{\text{ends}}$: It tests the free energy variations in base pairing between the two ends of the antisense strands in the siRNA duplex, i.e. the 3’ and 5’ ends of the antisense strands, also known as differential stability of siRNA duplex ends (DSSE). The main characteristic of an effective siRNA is the unstable 5’end (End-diff is more positive).
| S. No. | Nucleotide sequence of target region | Sequence based features | Thermodynamic features (kcal/mol) |
|-------|--------------------------------------|-------------------------|----------------------------------|
|       |                                      | Position on mRNA | Target Exon | GC% | Overall ΔG | Duplex ΔG | Break-target ΔG | Intra oligo ΔG | Inter oligo ΔG | End diff ΔG |
| 1     | GGACAACCTCCTCTGATGATGA               | 267–287            | 1           | 47.60 | -29.7      | -34.2      | -1.2          | -1.0          | -11.8        | 2.66       |
| 2     | GCGATACTCCCATCTGCTTCA               | 1628–1648          | 11          | 52.30 | -24.6      | -31.1      | -1.9          | -1.9          | -15.2        | 1.14       |
| 3     | GCTGGTGACCATGTTAATTGA               | 3288–3308          | 23          | 42.85 | -26.6      | -32.1      | -1.5          | -1.1          | -12.9        | 2.49       |
| 4     | GCTCCGAAGAATCACGTTTCT               | 4113–4133          | 31          | 47.60 | -26.5      | -31.9      | -1.7          | -1.4          | -14.2        | 2.27       |
| 5     | GGACATCATTGGGAAGGATCA               | 5424–5444          | 40          | 47.60 | -31.3      | -33.9      | -1.0          | -0.1          | -10.9        | 2.76       |

### Chicken embryo myoblasts (CEM) primary cell culture

The 9-day old embryos were used for preparing the chicken embryo myoblast (CEM) primary cell culture (Sato et al. 2006). The collected eggs were cleaned with 70% ethanol and the broad end of the egg was cracked using sterile forceps and peeled off the white shell membrane to reveal the chorioallantoic membrane (CAM) below along with blood vessels. The sterile curved forceps were used for piercing the CAM and gently grasping the embryo under the head and lifted out, and transfer to the sterile 9-cm petri dish containing sterile phosphate buffer saline (PBS) and rinsed thoroughly. The head, limbs, and wings were removed by using scissors, and finally, the ventral side of the embryo was cut open to remove all internal organs and transferred in to a new second petri dish to dissect unwanted tissue like fat or necrotic material, debris and blood tinge was removed by washing with PBS several times. The sterile scissors were used to cut the tissue into fine 3-mm pieces. The minced tissue pieces were transferred to a sterile beaker containing a sterile magnetic bar and 10 ml of 2.5% trypsin, and the beaker was placed on the magnetic stirrer for stirring at about 100 rpm for less than 10 minutes at 37°C and it was further filtered through a sterile double-layered muslin cloth into a fresh beaker and the filtrate was centrifuged for 5 minutes at 1500 rpm. After discarding the supernatant the resulting pellet was resuspended in 5 ml of growth medium (DMEM, HiMedia) with fetal bovine serum (FBS) to stop the trypsin action. The cell suspension was diluted to 1x10⁶ cells/ml in a growth medium with the help of a hemocytometer and seeded approximately 2x10⁵ cells/cm² in each 25-cm² tissue culture ask. The tissue culture asks were incubated at 37°C with 5% CO₂ until a confluent monolayer was obtained.

### Transfection of shRNA construct

The recombinant shRNA clone (pENTR™/U6 Entry vector with respective shRNA insert) against ACACB gene and a control plasmid (plasmid containing scrambled shRNA) was transfected into the chicken embryo myoblast (CEM) primary cell culture using the Gene Pulser Xcell™ Electroporation system (Bio-Rad) to predict the activity of respective shRNAs in myoblasts. Two days before electroporation, the cells were transferred to a new 25-cm² tissue culture flask with fresh growth medium (DMEM) supplemented with 10% FBS and antibiotic antimycotic solution. The cells were grown up to the late-log phase, such that there will be 70–80% confluent on the day of the experiment. Adherent cells were first trypsinized by adding 0.1ml/cm² trypsin, it was finally inactivated with complete medium, and the cells were harvested by centrifuging at 1500 rpm for 5 minutes at room temperature. The cell pellet was re-suspended in growth medium yielding an approximate cell concentration of 2.5 X 10⁶ cells/ml medium. The transfection reactions of all the shRNAs along with scrambled shRNA were carried out in triplicates. Approximately 0.4ml of the cell suspension was transferred into a 0.4 cm electroporation cuvette and the purified plasmid DNA was added to the cell suspension to a final concentration of approximately 50µg/ml. The DNA and cell suspension were mixed
in the cuvette and placed in the shock pod unit holder in the electroporation apparatus and a single square wave pulse was given at 110V with 25 milliseconds pulse length. After the pulse, cell suspension was transferred into a 25-cm² flask containing 5ml of growth medium. The flasks were rocked gently to assure even distribution of the cells over the surface of the flask and incubated at 37°C in a CO₂ incubator. After 48 hours of transfection, the adherent cells were trypsinized and transferred into sterile 15ml conical tubes. The cells were harvested by centrifuging at 1000 rpm for 10 minutes and stored at -80°C until total RNA isolation.

Total RNA isolation and cDNA synthesis

The total RNA was isolated from the CEM cell pellet using 1ml of Trizin (GCC Biotech, India), according to the manufacturer's protocol. After homogenization, the sample was incubated for 5 minutes at room temperature and then chloroform (200µl/sample) was added to the sample, shaken vigorously for 15 seconds, and incubated for 5 minutes at room temperature followed by centrifugation at 12000 rpm for 15 minutes at 4°C. The upper aqueous phase was transferred to a new 1.5ml RNase free sterile microcentrifuge tube and 500µl of isopropanol was added for precipitation. The tubes were incubated for 30 minutes at -20°C and then pelleted by centrifugation at 12000 rpm for 10 minutes at 4°C. The pellet was washed with 750µl of 75% ethanol, air-dried the pellet for 5 to 10 minutes, and then dissolved in 20µl of RNase-free water. The total RNA was treated with DNase I (HiMedia, India) to remove any trace amount of genomic DNA. The RNA quality and quantity were checked by 1.2% denatured agarose gel and Jenway™ Genova Nano Micro-volume Spectrophotometer (Fisher Scientific, USA) respectively. For each sample, 2µg of total RNA was reverse transcribed using the Verso cDNA Synthesis Kit (Thermo Scientific, USA) in a 20µl reaction using Oligo dT and random primers. The cDNAs were diluted at 1:3 with nuclease-free water before the qPCR analysis.

Primer designing and qRT-PCR

A total of 9 genes were selected on the basis of their role in de novo fat synthesis and immune response viz. ACACA (Acetyl-CoA Carboxylase Alpha), ACACB (Acetyl-CoA Carboxylase Beta), FASN (Fatty acid synthase), SCD (Stearoyl-CoA Desaturase), ELOVL2 (ELOVL Fatty Acid Elongase 2), CPT1 (Carnitine palmitoyltransferase 1), IFNA (Interferon alpha), IFNB (Interferon Beta), and BLB1 (Major histocompatibility complex class II beta chain BLB1). The sequences were downloaded from NCBI and CDS region was identified by using the ExPASy translation tool (https://web.expasy.org/translate/) and the primers were designed based on CDS using IDT Primer Quest software (https://www.idtdna.com/Primerquest/Home/Index) (Table 3). The expression levels of target genes (ACACA, ACACB, FASN, SCD, ELOVL2, CPT1, IFNA, IFNB, and BLB1) and reference gene Albumin (ALB) were quantified using thermal cycler Himedia Insta Q96™ with Bright Green 2X qPCR Master mix ROX (abm, Canada). The qPCR experiments were performed in a 10µl reaction volume [containing 1µl of diluted cDNA, 5µl of BrightGreen 2x qPCR MasterMix, 0.3µl of each primer] under the following program: an initial denaturation for 5 minutes at 95°C, followed by 40 cycles of amplification with the 30s of denaturation at 95°C and 60s of annealing/extension at 60°C. The dissociation curve was obtained by heating the amplicon from 55 to 95°C. All qPCR reactions were carried out in three biological replicates. Non-template controls (NTC) were also included in each run for each gene.
Table 3
The List of primers used for quantitative real-time PCR (qRT-PCR).

| S. No. | Gene Symbol | Primer Sequences (5'-3')                                      | Amplicon Size (bp) | Annealing temp (°C) | Accession No. |
|--------|-------------|----------------------------------------------------------------|-------------------|---------------------|---------------|
| 1      | ACACA       | F: CAGATTTGTTGTCATGGTGAC                                       | 162               | 60                  | NM_205505.1   |
|        |             | R: ACAGCCTGCACTGGAATGC                                           |                   |                     |               |
| 2      | ACACB       | F: GCTCCTGCTGCCCATATATTA                                       | 94                | 60                  | NC_006102     |
|        |             | R: GTCCGTGATGACACCTTTCT                                          |                   |                     |               |
| 3      | FASN        | F: GTTCTCTGTACAGAGAAATGTG                                       | 168               | 60                  | NM_205155.3   |
|        |             | R: CCATGGTTGACTTGTGGATC                                          |                   |                     |               |
| 4      | SCD         | F: TGACGCTGATCCCTTCTGC                                          | 152               | 60                  | NM_204890.1   |
|        |             | R: AATAGTCAAGAGATCCGGCAG                                         |                   |                     |               |
| 5      | ELOVL2      | F: ATGTTTTGGACCTCGAGATGC                                         | 221               | 60                  | NM_001197308.1|
|        |             | R: CACGTGGCAAGATCCGGC                                            |                   |                     |               |
| 6      | CPT1        | F: GCCTTCGTGCGCAGAT                                              | 146               | 60                  | DQ314726.1    |
|        |             | R: ACGTAGGGCAGAAGAGG                                             |                   |                     |               |
| 7      | ALB         | F: TCCTGATCGCTACACTAAG                                           | 98                | 60                  | NM_205261.2   |
|        |             | R: CTGGCAGCAGTTAGTACCAATA                                         |                   |                     |               |
| 8      | IFNA        | F: CTGTCACGCTCCTTCTG                                            | 170               | 60                  | NM_205427     |
|        |             | R: GTGTGTTGAGGAGCAG                                              |                   |                     |               |
| 9      | IFNB        | F: CTCTTCAGAATACGGGCT                                           | 164               | 60                  | NM_001024836  |
|        |             | R: GTGTGTGGCTGCTAAG                                              |                   |                     |               |
| 10     | BLB1        | F: TGAGTGCCACTACCTGAAC                                           | 200               | 60                  | NM_001044679  |
|        |             | R: GGCAGTACGTGTCCCAG                                             |                   |                     |               |

Estimation of cholesterol and triglycerides

After forty-eight hours of transfection, control as well as ACACB knockdown adherent cells were trypsinized and transferred into 15ml sterile conical tubes and pelleted by centrifugation for 10 minutes at 10000 rpm. The pellet was washed with ice-cold PBS by centrifugation at 10000 rpm for 10 minutes at 4°C. After harvesting of cells we counted the cell numbers from each culture flask in neubauer chamber and thus normalized the cell numbers before proceeding to estimate cholesterol and triglycerides. For each category (knockdown group and control group) equal number of cells (1x10^6 cells/ml) were collected for estimation of cholesterol and triglycerides. The pellet was lysed by using 1ml complete cell extraction buffer (150 mM NaCl, 1% NP-40, 0.1% Sodium dodecyl sulphate, and 50 mM Tris, pH8) and the lysate was transferred to the 1.5ml sterile microcentrifuge tubes, vortexed the tubes and incubated on ice for 30 minutes with occasional vortexing. The lysate was centrifuged at 4°C for 20 minutes at 14000 rpm and aqueous phase was collected into 1.5ml sterile microcentrifuge tubes. This cell extract was used for cholesterol and triglycerides estimation with the help of the Turbo Chem 100 automatic blood analyzer. The commercially available cholesterol and triglyceride reagent kits (Identi cholesterol test kit, Identi Triglyceride test kit CPC Diagnostics, Chennai, India) were used for cholesterol and triglycerides estimation.

Statistical analysis
The experiments were repeated twice, the relative expression of each gene was calculated by using $2^{-\Delta \Delta Ct}$, and statistical analysis was carried out using the trial version of SPSS 25. A univariate general linear model with Tukey’s HSD and DMRT as post hoc test was used to study the significant difference between different shRNA groups due to the knockdown effect of target genes. Data from representative experiments were presented as Mean ± SE for different samples with differences determined by least significant differences at 5% level ($p \leq 0.05$).

**Results**

Secondary structures and thermodynamic properties for shRNAs

The evaluation of the shRNA constructs 1 and 5 were devoid of any secondary structures in their antisense strands while shRNA constructs 2, 3, and 4 formed secondary structures (Fig. S1). Hence, the shRNAs 1 and 5 showed minimum free energy (MFE) of 0.00 kcal/mol whereas, the shRNA 2, 3, and 4 showed MFE of -1.40, 0.30, and 0.10 kcal/mol respectively, due to the secondary structure formation (Fig. S1). The stem loop structures were formed for all the ACACB mRNA target regions of anti-ACACB shRNA molecules (Fig. S2). The GC percent of shRNAs ranged from 43–52% where shRNA 3 had the lowest (43%) of all the shRNAs studied, while shRNA 2 had a higher percentage (52%). The predicted values (negative) about overall/net $\Delta G$ value, $\Delta G$ duplex, and $\Delta G$ break-target were found to be highest in shRNA 5, shRNA 1, and shRNA 2 respectively, while the lowest values were observed in shRNA 2, shRNA 2, and shRNA 5, respectively. The DSSE was found to be highest for shRNA 5 and lowest for shRNA 2 (Table 2).

Cloning and confirmation of ACACB anti shRNA in pENTR™/U6 Vector

The RNAi-Ready pENTR™/U6 vector was used to ligate all the five shRNA oligos of the ACACB gene and transformed into One Shot TOP10 chemically competent E. coli cells. The recombinant clones of all the shRNAs were confirmed by colony PCR (Fig. 1). Further, the plasmid DNA was isolated from the recombinant clones and confirmed by plasmid PCR (Fig. 1). Each recombinant pENTR™/U6 construct was sequenced to confirm the sequence of the shRNA, which revealed the absence of mutations. The DNA was isolated from the transfected cell culture of each construct and used as a template in PCR to check for the presence of pENTR™/U6 Entry vector contains shRNA and found a product of 293 bp length signifying the successful transfection (Fig. 1).

Silencing efficiency of ACACB shRNAs

The qRT-PCR was performed with ACACB and ALB gene specific primers with all five shRNA treated CEM samples and scrambled shRNA treated samples was used as a control. The initiation phase, exponential phase, and plateau phase of amplification curves were all optimal, indicating that the product was amplified exponentially, i.e. fluorescence emission was corresponding to the amplified template. Melting curve analysis was performed at the end of the qRT-PCR cycle to verify the specificity of amplification, revealed a single peak for all genes, suggesting that the PCR products were homogeneous. The knockdown performance of shRNA 1 and 5 against scrambled shRNA was found to be significantly different ($P \leq 0.05$) in the qRT-PCR study. The fold change of ACACB gene in different shRNA constructs was 0.33, 0.75, 0.61, 0.62 and 0.31 in the cells with shRNA 1, shRNA 2, shRNA 3, shRNA 4 and shRNA 5 constructs respectively. In contrast to the scrambled shRNA, the knockdown (KD) percent of ACACB mRNA caused by different shRNA ranged from 69% (shRNA 5) to 25% (shRNA 2), respectively (Fig. 2).

Effect of shRNAs on immune response genes

In the knockdown cells, the relative expression of immune response genes such as IFNA, IFNB, and BLB1 was also monitored by qRT-PCR in target as well as control samples. However, when compared with scrambled shRNA, no significant difference was observed in all five shRNA groups (Fig. 3).

Relative quantification of de novo fat synthetic genes
Once the knockdown of Acacb gene was found, the relative expression of de novo fat synthetic genes such as, Acaca, Fasn, Scd, Elov12, Cpt1 have been studied. The Acaca, Fasn, Scd and Cpt1 genes were down-regulated and Elov12 gene was significantly up-regulated. The fold change of Acaca, Fasn, Scd, Cpt1 and Elov12 was 0.55, 0.08, 0.01, 0.05 and 2.87 respectively (Fig. 4).

Quantification of cholesterol and triglycerides

The cholesterol and triglycerides were estimated from the control and Acacb knockdown cell lysate and estimated by using Turbo Chem 100 automatic blood analyzer. We found significant (P < 0.01) reduction in cholesterol and triglyceride levels in Acacb knockdown cells at 47.71% and 34.91% respectively as compared to the control (Fig. 5).

Discussion

Synthesis of potential shRNAs

The success of RNAi depends on the designing of the shRNA for specific target recognition and minimization of off-target effects. In the present study, unique/specifc shRNA molecules were designed based on the Reynolds ranking criteria for the Acacb gene and also, the specifcity of the shRNA sequences is important for the formation of RISC (Reynolds et al. 2004; Paddison et al. 2004). In addition to specifcity, the G-C content plays an enormous role in the formation of duplex siRNA molecule. Low G-C content is known to decrease afnity for the target sequence, whereas higher G-C content interferes with RISC formation that eventually cleaves the mRNA molecules. The G-C content of the designed shRNA molecules in the present study was moderate for the duplex siRNA formation. In addition to the G-C content, the lack of internal repeats, an A/U-rich 5' end, Tuschl motifs, and other features were included which improves the silencing afnity of siRNA (Fuchs et al. 2004).

Finally, a simple local alignment search tool (BLAST) was also employed to ensure that shRNA had no signifcant homologies with genes other than the target to avoid possible off-target effects. Further, silencing performance was also positively associated with the siRNA-mRNA duplex (Gduplex) stability (Pascut et al. 2015). Similarly, 1 and 5 shRNAs had higher Gduplex values than shRNA 2, 3, and 4 (which had a low Gduplex value), meaning that 1 and 5 shRNAs could bind the target site more efciently. The knockdown efciency of 1 and 5 shRNAs was high, supporting the above predictions. The duplex asymmetry (DSSE) and target site accessibility could improve knockdown efciency by about 26% and 40%, respectively (Shao et al. 2007). For enhanced RNAi potency, the siRNAs suitable disruption energy and DSSE is < −10 kcal/mol and > 0.0 kcal/mol, respectively (Shao et al. 2007). We noticed that the expected disruption energies and DSSE for all of the modeled shRNAs were less than −10 kcal/mol and greater than 0.0 kcal/mol, respectively, suggesting that all shRNAs have the ability to silence genes. It has been discovered that the development of self-structures in shRNA (both unimolecular and bimolecular shRNA folding) reduces the equilibrium afnity for the target mRNA (Lu and Mathews, 2007). For all of the shRNAs the predicted self-structure energies in this sample were low, indicating their efciency in silencing.

Gene silencing caused by shRNAs is mostly due to sequence-specifc mRNA degradation by antisense/guide strand of shRNA (Martinez et al. 2002). The ‘guide strand’ is inserted into the active RNA-induced silencing complex (RISC) to locate the mRNA, which has a complementary sequence leading to the endonucleolytic cleavage of the target mRNA leading to gene silencing (Hannon, 2002). The degree of secondary structure in the antisense strand was of utmost importance in determining the highly active shRNAs among the several factors controlling the efciency of gene silencing (Patzel et al. 2005). The development of secondary structure in the antisense strand is a signifcant factor in shRNA-induced gene silencing (Wolfman et al. 2003; Qiao et al. 2008). The guide-RNA structures are categorized as those, greatest silencing caused by sequences that do not form secondary structures, second best are stem-loop structures with ≥ 2 free nucleotides at 5’ end and ≥ 4 free 3’ nucleotides, followed by internal-loop, two stem-loop, and short free end stem-loop structures (Patzel et al. 2005). Therefore, it is understood that secondary structure formation correlates negatively with the eficient silencing of the gene. Hairpin-structured shRNAs are unable to fully open during their function, resulting in low gene silencing efciency. As a result, the RISC-siRNA complex formed would not be that much effective while interacting with the complementary mRNA. The mRNA local structure is one of the key factors with a strong effect on silencing of the shRNA molecule (Schubert et al. 2005; Gredell et al. 2008;
The mRNA contains loop structures that provide easy access to the guide strand for binding the target region which enhances the gene silencing efficiency but the presence of paired nucleotides and hairpins reduced the gene silencing, respectively (Schubert et al. 2005; Holen 2005; Li and Cha 2007). The GC content of the mRNA target region plays a crucial role in loading shRNA into RISC complex (Reynolds et al. 2004; Shah et al. 2007; Wang and Mu 2004; Kretschmer-Kazemi Far 2003; Bohula et al. 2003; Stewart et al. 2008; Luo and Chang 2004). In our study, all the ACACB mRNA target regions of shRNAs revealed stem-loop structures and optimum GC% content (42.86 to 47.6). However, there might be some other factors which also contribute to the different silencing efficiency of shRNA constructs.

Following these findings, shRNA constructs 1 and 5 were devoid of any secondary structures in their antisense strands while shRNA constructs 2, 3, and 4 formed secondary structures. Accordingly, shRNA 1 and 5 had higher knockdown efficiency (67 and 69%) respectively, while shRNA 2, 3, and 4 had lower knockdown percentages (25–39%) which confirms the earlier reports (Patzel et al. 2005; Wolfman et al. 2003; Qiao et al. 2008). As compared to the control, all the shRNAs showed lower expression of the ACACB gene and among the five shRNAs, the shRNA 5 showed higher knockdown whereas shRNA 2 showed lower knockdown respectively.

**ACACB gene silencing in chicken myoblast cells**

Transfection of shRNA constructs into the CEM resulted in a notable down-regulation of ACACB mRNA, implying that CEM culture can be used as an *in vitro* model for some functional studies. Owing to the lack of secondary structures, shRNA 1 and 5 had a higher silencing efficiency than the other shRNAs. In the case of shRNA 2, intrinsic factors such as low duplex energy and high disruption energy may have made the mRNA-shRNA hybridization complex less stable, resulting in incomplete accessibility of the target mRNA region. Secondary structure formation in shRNA 2, 3, and 4 might have reduced their efficiency of silencing by influencing the hybridization of the siRNA/RISC to its target site (Schubert et al. 2005).

In goat fibroblast cells, observed that substantial silencing of ACTRIIB gene as 33–66 % (Patel et al. 2014). Besides, several researchers have carried out knockdown experiments on the MSTN gene in different animals, including chicken. However, the MSTN gene was silenced up to 68 and 75% in chicken embryo fibroblast cells (Sato et al. 2006; Tripathi et al. 2013). Later on, in the same chicken embryo fibroblast cells shRNA was used against MSTN, ACTRIIA, and ACTRIIB genes and observed the knockdown percentage of 68, 82 and 87, respectively (Tripathi et al. 2013; Satheesh et al. 2016; Guru Vishnu et al. 2019). Even in *in vivo* studies, MSTN knockdown chicken showed 28% more body weight during 42 days of age compared to the control broiler chicken (Bhattacharya et al. 2017). In duck embryonic fibroblasts, different lentivirus-mediated shRNA groups were compared and showed decreased the MSTN mRNA expression by 61.6, 76.9, and 79.1%, respectively (Tao et al. 2015). Further, in caprine foetal fibroblasts, transient transfection of anti-myostatin shRNA decreased the mRNA level by 89 and 72%, respectively (Kumar et al. 2014; Jain et al. 2015). The importance of using thermodynamic features in shRNA designing was highlighted by observing the relationship between shRNA thermodynamic parameters and the silencing performance of different shRNAs. Based on the findings, it is hypothesized that every designed shRNA with $G_{overall}$, $G_{duplex}$, $G_{breaktarget}$, and $G_{ends}$ of shRNAs in the range of 25 to 32 kcal/mol, 31 to 35 kcal/mol, 1.0 to 1.9 kcal/mol, and > 0.0 Kcal/mol kcal/mol, respectively, would appease for maximum silencing efficiency.

**Effect on immune response genes**

Although RNA interference has promised to be a powerful experimental tool to manipulate gene function, there has been a growing concern about the use of shRNA due to off-target effects such as activation of immune response. Few studies have stated that both non-immune cells and immune cells can recognize shRNAs independent of the sequence leading to interferon (IFN) induction and inflammatory cytokines both *in vivo* and *in vitro* (Sledz et al. 2003; Judge et al. 2005). The IFN response caused by the activation of dsRNA-dependent protein kinase R (PKR) leads to the global inhibition of protein synthesis (Judge et al. 2005). The dsRNA (> 23-bp) can affect cell viability and induce a powerful interferon response (strong up-regulation of the dsRNA receptor, Toll-like receptor 3) in a cell type-specific manner (Reynolds et al. 2006). It was concluded that the length threshold of siRNA-induced interferon response was not constant, but it differed between various types of cells significantly.
However, shRNAs shorter than 30 bp can evade PKR activation (Robbins et al. 2006) and some experiments exhibited a significant increase in the expression of immune response genes including Scramble shRNA. It was also observed that activation of interferon response in goat myoblast cells due to exogenous administration of shRNA against the ACTRIIB gene (Patel et al. 2014). Interferon modulation in chicken embryonic myoblast cells varied between 46–112 folds of OAS1 and 2-7.2 folds for IFNβ compared to mock-transfected control due to anti myostatin shRNA (Tripathi et al. 2013). It has been reported that the shRNA-mediated myostatin knockdown in transgenic sheep showed increased MHCI expression (Hu et al. 2013).

There are several reports regarding shRNA-induced interferon responses suggested that a high level of shRNA expression might be due to the accumulation of unprocessed or aberrantly processed transcripts triggering interferon response (Stewart et al. 2008; Cao et al. 2005; Watanabe et al. 2006). The shRNA transfected porcine embryo cells, showed induction of the OAS1 and IFNβ genes by 1000 and 50 folds respectively (Stewart et al. 2008). Further, reported that the introduction of H1 and U6 promoter-based shRNA constructs by pronuclear microinjection led to induction of OAS1 gene and early embryo lethality (Bridge et al. 2003). In the present experiment, the expression of interferon genes IFN α, IFN β, and BLB1 were analyzed in control (scrambled shRNA treatment) and knockdown cells possessing different shRNA molecules to explore the impact of shRNA on immune function. It was observed that there was no significant difference of expression of IFN α, IFN β, BLB1 genes between knockdown and scrambled shRNA treated cells. Hence, shRNA molecules used in the present study have not been captured by the interferon mechanism in vitro as any foreign DNA fragment of a specific length is normally detected and destroyed by the interferons. It may therefore be construed that these shRNA molecules have been very much effective to silence ACACB expression without interfering with the body’s immune system. However, the introduction of short (<30 nt) dsRNAs with 2-base 3 overhangs resembling dicer processing does not activate the interferon pathway and also shRNA expression from vectors in the nucleus resemble endogenous miRNA (Elbashir et al. 2001). The administration of naked, synthetic siRNAs in mice showed down-regulation of endogenous or exogenous targets without inducing an interferon response (Heidel et al. 2004). Similarly, the in vitro siRNA study showed the absence of IFN induction in human CD34 + progenitor cells (Robbins et al. 2006). On the contrary, activation of PKR, OAS, RIG-1, TLR 7, and TLR 8 by sequences shorter than 19 bp (Gantier et al. 2007).

**Effect of ACACB gene silencing on de novo fat synthetic genes**

Now-a-days, researchers are more focused on ACACA and ACACB to understand the chemistry and biological activity because they are very important enzymes in fatty acid synthesis and oxidation. The ACACB gene is localized subcellularly on the mitochondrial membrane and is involved in the synthesis of malonyl-CoA, and this inhibits the CPT1, which plays an important role for controlling the two opposing pathways i.e. fatty acid synthesis and oxidation. In this study, we knocked down the ACACB gene in chicken myoblast cells and tracked the expression of key fatty acid metabolism genes such as ACACA, FASN, SCD, ELOVL2, and CPT1. We observed down-regulation of ACACA, FASN, SCD, and CPT1 gene and up-regulation of ELOVL2 in ACACB knockdown myoblast cells. The down-regulation of ACACA, FASN, SCD genes indicates suppression of the fatty acid synthesis, and up-regulation of ELOVL2 indicates the enhancement of long-chain fatty acids formation (Fig. 6). The CPT1 is a rate-limiting enzyme and down-regulation of this enzyme indicates blocking the β-oxidation for balancing the fatty acid synthesis and oxidation because of fewer fats accumulation in the tissues.

**Effect of ACACB gene silencing on cholesterol and triglyceride synthesis**

In chicken egg and meat the high cholesterol and triglycerides are most undesired components and which compels us to eat chicken products in less quantity. The ACVR2B knockdown chicken showed significantly low cholesterol (Bhattacharya et al. 2019). In mice, LNP-formulated siRNAs were used for knockdown of the ApoB gene resulted in significant reduction of total cholesterol and LDL cholesterol, which suggested that targeting ApoB is a therapeutic approach for hyperlipidaemia treatment (Tadin-Strapps et al. 2011). In this study, we observed a significant reduction of cholesterol and triglycerides at 47.71% and 34.91%, respectively in ACACB knockdown cell lysate compared to the control. Based on these results, we suggest that the ACACB knockdown chicken may produce low cholesterol and triglycerides.
In conclusion, from the present study we have identified potential shRNA molecules against the ACACB gene where shRNA1 and shRNA5 showed more than 60% knockdown efficiency on the expression of the ACACB gene under in vitro myoblast cell culture system. The silencing of the ACACB gene showed to have a direct effect on the down-regulation of ACACA, FASN, SCD, and CPT1 genes, and up-regulation of ELOVL2 gene in myoblast cells. We suggest that these shRNA molecules may be used under in vivo system for the development of knockdown chicken having a potential of producing lean meat by silencing of expression of the ACACB gene.

Declarations

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Authors’ Contributions G. S conducted the wet lab experiments and prepared the first draft. P. J. L carried out the statistical analysis. S. T. V. R prepared all the tables. R. M. V. P prepared all the figures. M. K edited the draft. C. S. P and A. R. P helped in cell culture. D. D helped in wet lab experiments. T. K. B conceived the idea, prepared the plan of work and final editing of the draft.

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Compliance with ethical standards

The entire study was approved by the Institute Animal Ethics committee (IAEC) and Institute Bio-safety Committee (IBSC) of ICAR-Directorate of Poultry Research, Hyderabad, India. All the bio-safety guidelines of IBSC were followed while conducting the experiments.

Conflict of interest The authors have no conflict of interest.

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**Figures**
Figure 1

(A) generation of double-stranded oligo (ds oligo). Lane 1 and 8: 20 bp DNA ladder; Lane 2: annealed scrambled oligo; Lane 3-7: annealed shRNA oligos 1-5. (B) and (C) Agarose gel electrophoresis of colony and plasmid PCR amplified fragment (293 bp) of anti ACACB shRNA constructs. Lane 1: negative control; Lane 2-5, 7: anti ACACB shRNA constructs 1-5; Lane 8: Scrambled shRNA construct; Lane 6: 100 bp plus DNA ladder. (D) Agarose gel electrophoresis of DNA from cell lysate amplified fragment (293 bp) of anti ACACB shRNA constructs. Lane 1-5: anti ACACB shRNA constructs; Lane 6: scrambled shRNA construct; Lane 7: 100 bp plus DNA ladder. Lane 8: negative control.
Figure 2

(A) The CEM cells transfected with different shRNA constructs and before harvesting the cells were photographed by using a DeltaPIX microscope. (B) Knockdown efficiency of anti ACACB shRNA constructs in CEM. ShRNA 1-5: Anti ACACB shRNA constructs; shLac Z: negative control (scrambled shRNA).
Figure 3

Induction of immune response genes due to anti ACACB shRNA constructs. shRNA 1-5: anti ACACB shRNA constructs, shLacZ: negative control (scrambled shRNA). (A) IFNA: Interferon A; (B) IFNB: Interferon B; (C) BLB1: Major Histocompatibility complex II beta chain BLB1.
Figure 4

Gene expression associated with fatty acid metabolism related genes in ACACB knockdown chicken myoblast primary cells compared with control cells. Mean values were different at *P ≤ 0.05.
Figure 5

Estimation of cholesterol (A) and triglycerides (B) in ACACB knockdown chicken myoblast primary cells compared with control cells. Mean values were different at *P ≤ 0.01.
Diagrammatic representation of targeted gene expression analysis conducted in the present study. In chicken tissues, Acetyl-coenzyme A carboxylase 1 (ACACA) and acetyl-coenzyme A carboxylase 2 (ACACB) play distinct roles in lipid metabolism. Diet fat, carbohydrate, and protein are digested, and the fatty acids (FA), glucose, and amino acids are transported to various tissues, including liver, adipose, and muscle. In the cytosol acetyl-CoA is carboxylated to malonyl-CoA by ACC1 and utilized through fatty acid synthase (FAS) and Stearoyl CoA Desaturase (SCD) reactions to generate palmitate and palmitoleate, which is utilized in the synthesis of triglycerides (TG) and VLDL. Also, acetyl-CoA is carboxylated by ACC2 at the mitochondrial membrane to form malonyl-CoA, which inhibits the CPT1 and reduces acyl-CoA transfer to mitochondria for β-oxidation. The down and up arrow (↓) indicates the down and up-regulation of genes in ACACB knockdown CEM primary cells compared to the control cells. Mean values were different at *P ≤ 0.05.

**Supplementary Files**

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