Sida rhomboidea. Roxb Leaf Extract Down-Regulates Expression of PPARγ2 and Leptin Genes in High Fat Diet Fed C57BL/6J Mice and Retards in Vitro 3T3L1 Pre-Adipocyte Differentiation

Menaka C. Thounaojam¹, Ravirajsinh N. Jadeja¹, Umed V. Ramani², Ranjitsinh V. Devkar¹,* and A. V. Ramachandran¹

¹ Division of Phytothrapeutics and Metabolic Endocrinology, Department of Zoology, The M. S. University of Baroda, Gujarat 390002, India; E-Mails: menakachanu@yahoo.com (M.C.T.); rv_jadeja@yahoo.co.in (R.N.J.); av_rama@rediffmail.com (A.V.R.)
² Department of Animal Biotechnology, College of Veterinary Science and Animal Husbandry, Anand Agriculture University, Anand, Gujarat 388001, India; E-Mail: umedramani@yahoo.co.in

* Author to whom correspondence should be addressed; E-Mail: phyto_met@yahoo.com; Tel.: +91-9825935445; Fax: +91-0265-2226425.

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Abstract: Sida rhomboidea. Roxb leaf extract (SRLE) is being used by the populace of North-East India to alleviate symptoms of diabetes and obesity. We have previously reported its hypolipidemic and anti-diabetic properties. In this study, we report the effect of SRLE on (i) in vivo modulation of genes controlling high fat diet (HFD) induced obesity and (ii) in vitro 3T3L1 pre-adipocyte differentiation and leptin release. Supplementation with SRLE significantly prevented HFD induced increment in bodyweight, plasma lipids and leptin, visceral adiposity and adipocyte hypertrophy. Also, SRLE supplementation reduced food intake, down regulated PPARγ2, SREBP1c, FAS and LEP expressions and up-regulated CPT-1 in epididymal adipose tissue compared to obese mice. In vitro adipogenesis of 3T3L1 pre-adipocytes was significantly retarded in the presence of SRLE extract. Also decreased triglyceride accumulation, leptin release and glyceraldehyde-3-Phosphate dehydrogenase activity along with higher glycerol release without significant alteration of viability of 3T3L1 pre-adipocytes, was recorded. Our findings suggest that prevention of HFD induced visceral adiposity is primarily by down regulation of PPARγ2 and leptin gene expression coupled with attenuation of food intake in C57BL/6J mice. SRLE induced prevention of pre-adipocytes differentiation, and leptin release further
substantiated these findings and scientifically validates the potential application of SRLE as a therapeutic agent against obesity.

**Keywords:** *Sida rhomboidea*. Roxb; obesity; 3T3L1 cells; PPARγ2; leptin

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1. Introduction

Obesity, a fast spreading epidemic, is a major contributor to the global burden of chronic disease and disability. Currently, more than one billion adults worldwide are overweight and at least, 300 million of them are clinically obese [1]. Such individuals are maximally prone to type-2 diabetes, cardiovascular disease and hypertension in the long run [2,3]. Induction of obesity in humans is either ‘genetic’ or ‘lifestyle’ related. The latter is however a complex intermix of sedentary lifestyle and a high calorie diet amounting to nutritional overload [4,5]. Synthetic anti-obesity drugs have often been reported to be very costly with some of them also beset with undesirable side effects, thereby necessitating a need to screen natural/herbal products for treating obesity [6]. In recent times, many traditional herbal preparations have been put through a detailed scrutiny to explore their anti-obesity potential and the underlying mechanism of action [7].

*Sida rhomboidea*. Roxb (SR; Fam, Malvaceae) is a weed found in marshy places across India. In Ayurveda, it is known as “Mahabala” and is used for treating fever, heart diseases, burning sensations, urinary disorders, piles and all kinds of inflammation [8]. Phytochemical analysis has shown the presence of β-phenethylamine, N-methyl-β phenethylamine, S-(+)-N b-methyltryptophan methylester, vascine, choline, betaine,ephedrine, ψ-ephedrine [9] as the active principals of its aerial parts. Also, presence of sigmasterol, sigmastenol, campesterol, 22-dehydrocampesterol, sitosterol, cholesterol, spinasterol, 24-methylenecholesterol, 22-dihydrosipasterol and some n-alkanes and n-alcohols have been reported [10,11]. In parts of North East India, a decoction prepared from leaves of SR is consumed by rural and urban populace to alleviate symptoms of obesity and diabetes [12]. Recent studies from our laboratory have reported hypolipidemic [13], anti-diabetic [14] and cardioprotective [15] properties of *S. rhomboidea*. Roxb leaf extract (SRLE) in experimental models. SRLE has also been shown to ameliorate high fat diet (HFD) induced steatohepatitis in C57BL/6J mice [16] and prevents in vitro low density lipoprotein oxidation and macrophage apoptosis [17]. Acute and chronic toxicity tests have revealed that SRLE is non-toxic up to a dose of 3000 mg/kg body weight in mice [18].

These findings encouraged us to investigate the anti-obesity potential of SRLE and the possible mechanism thereat. Present study evaluates, the effects of SRLE on the expression of genes associated with adipogenesis, lipolysis and lipogenesis in HFD fed C57BL/6J mice. The focus of this study is on the mRNA expression peroxisome proliferator-activated receptor γ2 (PPARγ2), sterol regulatory element-binding factor 1c (SREBP1c), carnitine palmitol transferase-1(CPT-1), fatty acid synthase (FAS) and leptin in the epididymal adipose tissue of a high fat diet fed C57BL/6J mice. The efficacy of SRLE in controlling in vitro adipocyte differentiation and leptin release is also assessed.
2. Materials and Methods

2.1. Plant Material

Leaves of SR were collected from Imphal district, India in the month of June and shade dried. The plant was identified by Dr. Hemchand Singh, Taxonomist, Department of Botany, D.M. College of Science, Manipur, Imphal, and a sample (voucher specimen No. 216) was deposited at the herbarium of the Department of Botany.

2.2. Preparation of Extract

For preparation of extract, SR leaves were shade dried, manually crushed and grinded in an electric grinder to obtain fine powder. Hundred gm of powdered leaves were boiled in 1000 mL of distilled water at 100 °C for 3 h and filtered using a sterilized muslin cloth. Resulting filtrate was collected in petri plates and concentrated by heating at 100 °C to form a semisolid paste. This paste was kept overnight at 0 °C and a freeze-dried extract was obtained. The net yield obtained at the final step of extract preparation was 24% w/w.

2.3. Experimental Animals

Male C57BL/6J mice (6–8 weeks of age) were purchased from National Centre for Laboratory Animal Sciences (NCLAS), National Institute of Nutrition (NIN), Hyderabad, India. They were housed and maintained in clean polypropylene cages and fed with either low fat diet or high fat diet and provided with water ad libitum. The experimental protocol was carried out according to the guidelines of the Committee for the Purpose of Control and Supervision of Experiments on Animals (CPCSEA), India and approved by the animal ethical committee of Department of Zoology, The M.S University of Baroda, Vadodara (Approval No. 827/ac/04/CPCSEA).

2.3.1. High Fat Diet Induced Obesity in C57BL/6J Mice

A total of 18 mice were randomly allocated into 3 groups of 6 animals each. Group I (LEAN) consisted of mice fed with low fat diet (regular fat diet). Group II (Obese; OB) was fed with high fat diet [16]. Group III (OB + SRLE) consisted of mice fed with high fat diet containing 1% SRLE [14,16]. All animals were fed with their respective diets for 20 weeks.

At the end of the experimental period, overnight fasted animals were given mild ether anesthesia and blood was collected by retro orbital sinus puncture in EDTA coated vials. Plasma was obtained by cold centrifugation (4 °C) of the vials for 10 min at 2000× g. Later, animals were sacrificed by cervical dislocation under mild ether anesthesia and various fat pads were excised and, the epididymal fat pad was stored in RNAlater solution (Ambion) until analysis at −80 °C (Cryo Scientific Ltd., India).

2.3.2. Plasma Lipids and Leptin Assay

Plasma triglycerides (TG) and free fatty acids (FFA) were assayed using commercially available kits (Reckon Diagnostics, Ltd., Baroda, India) whereas; plasma leptin was assayed using anti-mouse
monoclonal antibody coated 96 microtiter plate as per the instructions of the manufacturer (KRISHGEN Biosystems, Ltd.).

Table 1. Composition of experimental diets.

| Ingredients             | Low Fat Diet (g/kg) | High Fat Diet (g/kg) |
|-------------------------|---------------------|-----------------------|
| Casein                  | 200                 | 200                   |
| L-Cystine               | 3                   | 3                     |
| Corn Starch             | 315                 | 0.0                   |
| Maltodextrin            | 35                  | 125                   |
| Sucrose                 | 100                 | 68.8                  |
| Cellulose               | 50                  | 50                    |
| Soybean Oil             | 25                  | 25                    |
| Lard                    | 20                  | 245                   |
| Mineral Mix ¹           | 10                  | 10                    |
| Di Calcium Phosphate    | 13                  | 13                    |
| Calcium Carbonate       | 5.5                 | 5.5                   |
| Potassium Citrate       | 16.5                | 16.5                  |
| Vitamin Mix ²           | 10                  | 10                    |
| Choline chloride        | 2                   | 2                     |
| Regular chow            | 195                 | 216.25                |
| **SRLE**                | **00**              | **10**                |
| **Energy Content**      | **16.25 kJ/g**      | **25.72 kJ/g**        |
| Protein, % of energy    | 20                  | 20                    |
| Carbohydrate, % of energy | 64                  | 35                    |
| Fat, % of energy        | 6                   | 45                    |

¹ Mineral mix adds the following components (per g mineral mix): sodium chloride, 259 mg; magnesium oxide, 41.9 mg; magnesium sulfate, 257.6 mg; chromium K sulfate, 1.925 mg; cupric carbonate, 1.05 mg; sodium fluoride, 0.2 mg; potassium iodate, 0.035 mg; ferric citrate, 21 mg; manganous carbonate, 12.25 mg; ammonium molybdate, 0.3 mg; sodium selenite, 0.035 mg; zinc carbonate, 5.6 mg; ² Vitamin mix adds the following components (per g vitamin mix): retinyl acetate, 0.8 mg; cholecalciferol, 1.0 mg; dl-a-tocopheryl acetate, 10.0 mg; menadione sodium bisulfite, 0.05 mg; biotin, 0.02 mg; cyanocobalamin, 1 mg; folic acid, 0.2 mg; nicotinic acid 3 mg; calcium pantothenate, 1.6 mg; pyridoxine-HCl, 0.7 mg; riboflavin, 0.6 mg; thiamin HCl, 0.6 mg.

2.3.3. Analysis of Gene Expression by Quantitative RT-PCR (qPCR)

Total RNA was isolated from the epididymal fat pad of control and experimental mice using Tri-reagent (Sigma Aldrich, USA). Quantity and quality of isolated RNA was assessed using nanodrop spectrophotometer (Thermo Scientific, Ltd.) and samples with a ration of $A_{260}/A_{280} > 1.9$ was processed for cDNA synthesis using Omniscript cDNA synthesis kit (Qiagen, USA). A reaction mixture of 20 μL contained, 2 μg total RNA, 10 × RT buffer, dNTP mixture (5mm each), 10 × random hexamer, RNase inhibitor (10 U/μL), Omniscript RT (4 U/μL) and RNase free water. The cDNA synthesis was carried out at 37 °C for 1 h using a Veriti 96 well thermal cycler (Applied Biosystems, USA). Real-time PCR assays were performed in 96-well plates in ABI 7500 Fast real-time PCR machine (Applied Biosystems, USA). Primer sequences for qPCR analysis were as follow: PPARγ2
(sense) TCACAAGAGCTACCAATG, (anti-sense) GCATCCTTCACAAGCATG, SREBP1c
(sense) GATCAAGAGGAGCAGTGC, (anti-sense) TAGATGGTGCTGCTGAGTG, FAS
(sense) GGGTCTATGCCACGATT, (anti-sense) CACAGGGACCGAGTAATG, CPT-1
(sense) CTCAGTGGGAGCGACTCTTCA, (anti-sense) GCCCTCTGTGCTGCTGAGTG, LEP
(sense) GACACCAAAACCCTCAT, (anti-sense) CAGAGTCTGGTCCATCT, GAPDH
(sense) AGGCCGGTGCTGAGTATGT, (anti-sense) GTGGTTGTCACACCCCATCACA.
Syber Green reaction mixture of 20 µL contained 10 µL Quantifast Syber green master mix (Qiagen, USA), 2 µL template, 1 µL of each primer and 6 µL nuclease free water. The following two steps thermal cycling profile was used for qPCR analysis, Step I (cycling step): 95 °C for 10 min, 95 °C for 15 s, 60 °C for 1 min and 95 °C for 15 s for 40 cycles. Step II (Melt Curve step): 60 °C for 15 s, 60 °C 1 min and 95 °C for 30 s.

The fold change in the expression of genes mentioned herein was done by normalizing the values of threshold cycle (CT) of target gene with the CT value of Housekeeping gene (GAPDH) so as to obtain the ΔCT value. These values were further normalized with ΔCT values of CON (ΔCTOB or OB+SR − ΔCTLEAN) so as to obtain ΔΔCT values. The fold change in expression was then obtained log 2 − ΔΔCT [19].

2.3.4. Microscopic and Morphometric Examination of Epididymal Fat Pad

Epididymal fat pad was fixed in 4% buffered paraformaldehyde, dehydrated in graded alcohol series and embedded in paraffin wax using automated tissue processor. Five µm sections were cut and stained with hematoxyline and eosin and examined under Leica microscope. Photographs of adipocyte were taken with Canon power shot S70 digital Camera at 400× magnification and adipocyte number per 1000 mm were calculated using image analysis software.

2.4. Maintenance of 3T3L1 Cells

3T3-L1 mouse preadipocytes (Obtained from National Centre for Cell Sciences, Pune, India) were maintained in Dulbaco’s modified eagle medium (DMEM) containing 10% fetal bovine serum (FBS) (Himedia Pvt Ltd., Mumbai, India) and 1% antibiotic-antimycotic solution (10×; (Himedia Pvt Ltd., Mumbai, India) and sub cultured every 3rd day using 0.25% trypsin-EDTA solution (Himedia Pvt Ltd., Mumbai, India). 3T3-L1 mouse preadipocytes (Obtained from National Centre for Cell Sciences, Pune, India) were

2.4.1. In Vitro Cytotoxicity Assay

Pre-confluent pre-adipocytes (5.0 × 10³ cells/well) were maintained in 96 well plates (Tarson India Pvt Ltd.) for 72 h in presence of SRLE (10–1000 µg/mL) or vehicle (0.9% NaCl). At the end of incubation period, 10 µL of 3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT; 5 mg/mL) was added to wells and the plates were incubated at 37 °C for 4 h. At the end of incubation, culture media was discarded and the wells were washed with phosphate buffer saline (Himedia Pvt Ltd., Mumbai, India). 150 µL of dimethyl sulphoxide was added to all the wells and were incubated for 30 min. Absorbance was read at 540 nm in ELX800 Universal Microplate Reader (Bio-Tek instruments, Inc., Winooski, VT) and % cell viability was calculated.
2.4.2. *In Vitro* Adipocyte Differentiation Protocol

Freshly sub cultured cells were seeded on 12 well culture plates at the density of $1.0 \times 10^5$ cell/well in DMEM containing 10% FBS and allowed to become confluent. Cells were maintained for 2 days in confluent stage (To arrest cell division). Later (at day 0), culture media was replaced with DMEM containing 0.5 mM 3-isobutyl-1-methylxanthine (Sigma Aldrich, USA), 0.25 $\mu$M dexamethosone (Sigma Aldrich, USA), and 10 $\mu$g/mL insulin(Sigma Aldrich, USA) and cells were maintained for 4 days. At the end of 4 days, culture media was replaced with maturation media containing complete DMEM with 10 $\mu$g/mL insulin and cells were maintained for another 8 days, subsequently media was replaced every 48 hr till the end of the experiment [20].

2.4.3. Qualitative and Quantitative Analysis of Adipocyte Differentiation

3T3-L1 pre-adipocytes were differentiated as described above in presence of absence of SRLE (10–200 $\mu$g/mL). At the end of 12th day, Oil Red O staining for adipocyte lipid accumulation was performed. At the end of incubation, cells were washed twice with PBS and fixed in a 4% buffered paraformaldehyde for 10 min, washed twice with Milli Q water (Millipore India Pvt Ltd.) and then stained using 0.5% Oil Red O (ORO) for 15 min at room temperature. Excess ORO dye was washed with Milli Q water and photographs were taken in Leica DMIL inverted microscope using Canon power shot S70 digital camera [16].

In another set of experiment the stained adipocytes were treated with 100% isopropanol (to extract intracellular Oil Red O stain) and then the absorbance (Optical density; OD) of the extracts was measured at 490 nm. Reagent blank and cell blank assay was also performed simultaneously to minimize non-specific staining. The difference in absorbance between cells with and without ORO dye was calculated. Percentage adipogenesis was calculated as: (OD of treated cells)/(OD of untreated cells) × 100.

2.4.4. Leptin Release and Triglyceride Accumulation Assays

3T3-L1 pre-adipocytes were differentiated as described above in presence of SRLE (10–200 $\mu$g/mL) or vehicle (0.9% NaCl) and leptin and TG contents were assayed in supernatant and cells respectively. On day 12, supernatants from each well were collected and leptin content was analyzed using mouse specific leptin ELISA kit (Krishgen, Biosystems) as per the instructions of the manufacturer. After removal of supernatants, cells were washed twice with Phosphate buffer saline (PBS) and solublized in 100 $\mu$L of 1% Triton X 100 (in PBS) and, assayed for total TG using commercially available enzymatic kit (Reckon Diagnostics, Baroda, India) using Merck Microlab L300 Semi-autoanalyzer. Results were expressed as % TG [20].

2.4.5. Glycerol Release Assay

3T3-L1 pre-adipocytes were differentiated as described above for 12 days. For glycerol release assay, differentiated adipocytes were incubated with SRLE (10–200 $\mu$g/mL) or vehicle (0.9% NaCl) for 48 h. At the end of incubation, supernatant was collected from each well and glycerol content was determined by spectrophotometric assay [21].
2.4.6. G3PDH Activity Assay

3T3-L1 pre-adipocytes were differentiated as described above in presence of SRLE (10–200 µg/mL) or vehicle (0.9% NaCl). On day 12 after removal of supernatants, cells were washed twice with ice-cold PBS on and lysed in Tris-EDTA buffer (25 mM Tris/1 mM EDTA, pH 7.5) and glyceraldehydes-3-phosphate dehydrogenase (G3PDH) activity was determined [22]. Protein content in the cell lysate was determined using bovine serum albumin as a standard [23].

2.5. Statistical Analysis

Statistical evaluation of the data was done by one way ANOVA followed by Bonferroni’s multiple comparison test. The results were expressed as mean ± S.E.M using Graph Pad Prism version 3.0 for Windows, Graph Pad Software, San Diego, CA, USA.

3. Results

3.1. Body Weight Gain, Food Intake and Feed Efficiency

As shown in Figure 1A, HFD fed OB mice recorded significant weight gain after 20 weeks compared to the lean control. However, HFD induced weight gain was significantly controlled by SRLE supplementation of OB mice. There was also time dependent decrement in food intake in SRLE supplemented OB mice (Figure 1B). There was a significant increment in the feed efficiency ratio (body weight gain/food intake) in OB mice compared to LEAN mice (0.071 ± 0.004 vs. 0.030 ± 0.002). However, FER was successfully controlled in the OB+SRLE mice compared to OB mice (0.058 ± 0.003 vs. 0.071 ± 0.004).

Figure 1. Effect of S. rhomboidea. Roxb leaf extract feeding on (A) body weight gain and (B) food intake. Results are expressed as mean ± S.E.M., n = 6. Where, time points not sharing common letter indicate significant differences (p < 0.05).
3.2. Plasma Lipids and Leptin

OB mice recorded significant increase in plasma TG, FFA and LEP compared to LEAN mice while, SRLE supplementation to HFD fed mice resulted in significant decrement in plasma TG, FFA and LEP compared to OB mice (Table 2).

Table 2. Effect of *S. rhomboidea*. Roxb leaf extract feeding on plasma triglycerides, free fatty acids and leptin titer.

|                      | LEAN | OB     | OB+SRLE |
|----------------------|------|--------|---------|
| **Plasma**           |      |        |         |
| Triglycerides (mmol/L) | 0.53 ± 0.02 a | 2.16 ± 0.05 b | 0.61 ± 0.04 a |
| Free fatty acids (mmol/L) | 1.80 ± 0.07 a | 4.68 ± 0.09 b | 2.33 ± 0.08 a |
| Leptin (ng/L)        | 12.00 ± 1.98 a | 46.24 ± 2.13 b | 25.01 ± 1.89 c |

Results are expressed as mean ± S.E.M., *n* = 6. Where, timepoints not sharing common letter indicate significant differences (*p* < 0.05).

3.3. Visceral Adiposity

Visceral adiposity was evident in the form of increase in various fat pad weights (Perirenal, abdominal and epididymal) in OB mice after 20 weeks of HFD feeding (Figure 2). OB mice supplemented with SRLE extract showed significant attenuation in HFD induced visceral adiposity and increment in fat pad weights (Figure 2).

Figure 2. Effect of *S. rhomboidea*. Roxb leaf extract feeding on morphological and anatomic evaluation of visceral adiposity in Lean (A and D), OB (B and E) and OB + SRLE (C and F) groups and abdominal, epididymal and perirenal fat pad weights. Results are expressed as mean ± S.E.M., *n* = 6. Where, timepoints not sharing common letter indicate significant differences (*p* < 0.05).
3.4. Microscopic and Morphometric Evaluation of Epididymal Fat Pad

Microscopic evaluation of epididymal fat pad of OB mice recorded adipocyte hypertrophy that was marked by significant reduction in the adipocyte number (120.0 ± 12.0 vs. 270.0 ± 19.00) per 1000 mm area compared to LEAN. However, OB+SR mice recorded significantly higher number of adipocytes per (170.0 ± 15.0 vs.120.0 ± 12.0) 1000 mm area compared to OB.

3.5. Quantitative RT PCR Analysis

HFD fed OB mice recorded significantly up-regulated mRNA expression of PPARγ2, SREBP1c, FAS and LEP, while CPT-1 expression was significantly down-regulated in epididymal fat pad compared to lean mice. However, these set of changes were significantly prevented in SRLE supplemented OB mice (Figure 3).

Figure 3. Effect of S.rhomboidea.Roxb leaf extract feeding on quantitative RT-PCR analysis of PPARγ2, SREBP1c, CPT-1 FAS and LEP mRNA expression. Results are expressed as mean ± S.E.M., n = 6. Where, timepoints not sharing common letter indicate significant differences (p < 0.05).

3.6. Cytotoxicity Assay

Cytotoxicity analysis of SRLE in pre-adipocyte cells registered non significant alteration in the cell viability in the dose range of 10–1000 µg/mL (Figure 4).
**Figure 4.** Effect of *S. rhomboidea*. Roxb leaf extract on cell viability. Results are expressed as mean ± S.E.M., *n* = 3. Where, *P* < 0.05, **P** < 0.01 and ***P*** < 0.001 and ns non significant compared to 0 µg/mL SRLE.

3.7. Qualitative and Quantitative Analysis of Adipocyte Differentiation

ORO staining of differentiated adipocytes at the end of 12 days revealed significant cytoplasmic lipid accumulation in the untreated differentiated adipocytes (Figure 5A) whereas, SRLE extract supplementation to differentiating 3T3L1 pre-adipocytes significantly reduced adipocyte differentiation, characterized by lesser cytoplasmic lipid accumulation (Figure 5B–F). Quantitative analysis of ORO staining recorded 30% to 75% inhibition in adipogenesis in the SRLE supplemented 3T3L1 cells.

**Figure 5.** Photomicrograph of Oil Red O stained differentiating 3T3L1 cells (A) untreated, (B) treated with 10 µg/mL SRLE, (C) treated with 20 µg/mL SRLE, (D) treated with 50 µg/mL SRLE, (E) treated with 100 µg/mL SRLE and (F) treated with 200 µg/mL SRLE and qualitative (whole well image of Oil Red O stained adipocytes) and quantitative (% adipogenesis) evaluation of adipocyte differentiation (G and H). Results are expressed as mean ± S.E.M., *n* = 3. Where, *P* < 0.05, **P** < 0.01 and ***P*** < 0.001 and ns non significant compared to 0 µg/mL SRLE.
3.8. Triglyceride Accumulation and Leptin Release in Differentiated Adipocytes

Figure 6A,B shows LEP release and TG accumulation in the untreated and SRLE supplemented differentiated adipocytes at the end of 12 days. Whereas, untreated adipocytes showed higher levels of LEP release and TG accumulation compared to pre-adipocytes (Figure 6A,B). SRLE supplementation to differentiating adipocytes recorded significant decrement in the LEP release and TG accumulation compared to differentiated untreated adipocytes (Figure 6A,B).

**Figure 6.** Effect of *S. rhomboidea*. Roxb leaf extract on *in vitro* (A) leptin release, (B) triglyceride accumulation, (C) glycerol release, and (D) G3PDH activity during 3T3L1 pre-adipocyte differentiation were evaluated as described in materials and methods. Results are expressed as mean ± S.E.M., *n* = 3. Where, *P* < 0.05, **P* < 0.01 and ***P* < 0.001 and ns non significant compared to 0 µg/mL SRLE.

3.9. Glycerol Release and G3PDH Activity Assay

SRLE supplementation to differentiating pre-adipocytes resulted in higher indices of glycerol release and lowered cellular G3PDH activity compared to untreated differentiated adipocytes (Figure 6C,D).

4. Discussion

The onset and progression of HFD induced experimental obesity in C57BL/6J mice has a physiological resemblance to high calorie diet induced obesity in humans [24]. The high fat diet induced obesity in C57BL/6J mice characterized by selective deposition of fat in the mesentery, is comparable to visceral adiposity in humans [25,26]. Hence, this experimental model is widely used to investigate anti-obesity potential of herbal formulations [27,28]. Significant prevention of body weight
gain and reduction in food intake caused due to SRLE supplementation of HFD fed mice recorded in the present study are in agreement with our previous observations [13]. SRLE induced decrement in food intake could be attributed to presence of ephedrine and pseudo-ephedrine in various species of Sida [29], including Sida rhomboidea. Roxb [9,30]. Hence, it is assumable that, ephedrine induced appetite suppression is the key aspect that inhibits body weight gain in SRLE supplemented OB mice. Previous studies have reported that, ephedrine promotes the release of noradrenaline and inhibits the uptake of noradrenalin that eventually; results in decreased food intake and satiety [31,32]. However, mechanism responsible for SRLE induced decreased food intake needs further investigation. Ephedrine content of SRLE is <0.01% and, in the present study, it amounts to 50 µg/kg body weight, a dose that is almost 400 times lesser than the upper limit toxic dose (24 mg/kg body weight/day) set by USFDA [33]. Further, the safety evaluation of SRLE reported from our laboratory had shown that dose of 3000 mg/kg body weight is non-toxic in Swiss albino mice [18].

Adipose tissue represents a dynamic endocrine organ that not only maintains energy balance but also controls lipid homeostasis mainly by two reciprocal processes; lipogenesis and lipolysis [34]. Moreover, adipogenesis is a complex process in which both adipocyte growth and differentiation contribute to overall adipose mass [35]. Therefore, adipocyte differentiation is crucial in the maintenance of adipose tissue integrity. The net balance between the three processes determines the onset of visceral obesity. In the present study, OB mice recorded significantly increased mass of various fat pads. These results are further substantiated by concomitant decrement in adipocyte number, due to augmented TG deposition and adipocyte hypertrophy. The observed changes in epididymal fat pad of OB mice are in agreement with the recorded elevation in plasma leptin titre and the up-regulated expression of leptin mRNA in adipose tissue. However, SRLE supplementation of HFD fed mice show higher number of non-atrophied adipocytes, decrement in fat pad weights and plasma TG levels coupled with lower plasma leptin titer and down-regulation of epididymal leptin mRNA expression compared to OB mice. Since, plasma leptin and its mRNA expression are synonymous with obesity [36] and TG induced adipocyte hypertrophy [37], the results recorded herein are indicative of the mitigative nature of SRLE against HFD induced visceral adiposity, essentially by altered leptin gene expression in adipocytes.

The sequence of events beginning with adipocyte differentiation and ending with obesity is marked by up-regulation of prominent adipogenic genes [38]. PPARγ2, a nuclear receptor is a transcription factor that plays a crucial role in stimulating differentiation of pre-adipocytes and progenitor cells into adipocytes [39]. Recent studies have shown that some of the lipid lowering plant extracts acts as PPARγ2 antagonists thus preventing fat accumulation and adipocyte differentiation [40,41]. SREBP1c is another adipogenic transcription factor that controls the production of endogenous ligands for PPARγ2 [42] thus regulating the downstream cascade of lipid homeostasis and adipogenesis [43]. SREBP1c mediated lipogenesis in hepatic and adipose tissue is differential [44]. PPARγ2 has been reported to mediate lipogenesis in adipose tissue either directly or via modulation of SREBP1c [45,46]. In the present study, OB mice recorded significant up-regulation of PPARγ2 and SREBP-1c expression in the epididymal adipose tissue. However, SRLE supplementation not only prevented the up-regulated expression of these genes but also attenuated adipogenesis.

FAS gene expression is governed by PPARγ2 and is involved in the facilitated conversion of FFA into TG in adipocytes thus contributing to adipocyte hypertrophy and visceral adiposity in HFD fed
OB mice [47]. This physiological condition is parallel by with elevated plasma levels of TG and FFA. However, the recorded relatively lowered plasma TG and FFA levels in SRLE supplemented HFD fed mice is attributable to the reduced food intake of these animals. Further, the down regulation of FAS in adipocytes can again be correlated with lower expression of PPARγ2 in SRLE supplemented OB mice and the resultant anti-adipocyte hypertrophy. In the present study, HFD fed OB mice recorded significant up-regulation of FAS expression along with down-regulation of CPT-1 expression thus shifting the overall balance towards lipid anabolism. However, up-regulation of CPT-1 expression observed in SRLE supplemented OB mice along with down-regulation of PPARγ2 and FAS provides ample evidence for the anti-lipogenic potential of SRLE. Also, previous studies have documented PPARγ2 induced down regulation of ACC along with up-regulation of CPT-1 expression that eventually controls visceral adiposity [48]. Hence, inhibitory effect of SRLE on expression of ACC can not be ruled out and requires further scrutiny. It is inferable from the results observed herein that, the anti-obesity property of SRLE is a two pronged process that acts via reduction of dietary intake and down regulation of PPARγ2.

Anti-obesity potential of herbal extracts is also assessed in vitro using 3T3L1 pre-adipocytes as; these cells accumulate TG and release LEP when they differentiate into adipocytes [49,50]. In the present study, experimentally induced adipocyte differentiation was characterized by plumping of adipocytes due to accumulation of red colored lipid droplets and higher percentage of adipocyte differentiation. Also, TG accumulation and G3PDH activity level were all elevated in untreated adipocytes. The fully differentiated state of adipocytes was confirmed by the augmented released of leptin from these cells. However, in the presence of SRLE, differentiating adipocytes results in an overall decrement in the qualitative and quantitative indices. Qualitatively, the inhibitory effect of SRLE on adipocyte differentiation was marked by admixture of undifferentiated and differentiated cells and minimal lipid accumulation. These observations were further substantiated by the dose dependent reduction in TG content, G3PDH activity and LEP titer. Further, treatment of SRLE to differentiated adipocytes resulted in dose dependent increment in the glycerol release. These findings provide unequivocal evidence for the potential of SRLE in preventing adipocyte differentiation, lipid accumulation and LEP release and promoting lipolysis.

5. Conclusion

It can be concluded from the present study that the mechanism governing the anti-obesity potential of SRLE is a two pronged process that involves: (i) attenuation of food intake; and (ii) down regulation of PPARγ2 and related lipogenic genes that control visceral adiposity. These contentions are substantiated by the demonstration of SRLE induced prevention of in vitro adipocyte differentiation and leptin release. However, detailed scrutiny of its active phyto-ingredie nts along with clinical investigations is warranted to develop a herbal anti-obesity formulation.

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