Vitamin A Improves Hyperglycemia and Glucose-Intolerance through Regulation of Intracellular Signaling Pathways and Glycogen Synthesis in WNIN/GR-Ob Obese Rat Model.

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ABSTRACT: Vitamin A and its metabolites modulate insulin resistance and regulate stearoyl-CoA desaturase 1 (SCD1), which are also known to affect insulin resistance. Here, we tested, whether vitamin A-mediated changes in insulin resistance markers are associated with SCD1 regulation or not. For this purpose, 30-week old male lean and glucose-intolerant obese rats of WNIN/GR-Ob strain were given either a stock or vitamin A-enriched diet, i.e. 2.6 mg or 129 mg vitamin A/kg diet, for 14 weeks. Compared to the stock diet, vitamin A-enriched diet feeding improved hyperglycemia and glucose-clearance rate in obese rats and no such changes were seen in lean rats receiving identical diets. These changes were corroborated with concomitant increase in circulatory insulin and glycogen levels of liver and muscle (whose insulin signaling pathway genes were up-regulated) in obese rats. Further, the observed increase in muscle glycogen content in these obese rats could be explained by increased levels of the active form of glycogen synthase, the key regulator of glycogen synthesis pathway, possibly inactivated through increased phosphorylation of its upstream inhibitor, glycogen synthase kinase. However, the unaltered hepatic SCD1 protein expression (despite decreased mRNA level) and increased muscle-SCD1 expression (both at gene and protein levels) suggest that vitamin A-mediated changes on glucose metabolism are not associated with SCD1 regulation. Chronic consumption of vitamin A-enriched diet improved hyperglycemia and glucose-intolerance, possibly, through the regulation of intracellular signaling and glycogen synthesis pathways of muscle and liver, but not associated with SCD1.

Keywords: retinoids, adipose tissue, insulin, glucose homeostasis, gene expression

INTRODUCTION

Stearoyl-CoA desaturase (SCD1), a microsomal membrane-associated enzyme, introduces double bonds at the delta-9 position of saturated fatty acids (SFAs), palmitic acid (C16:0) and stearic acid (C18:0), and thereby converting them into monounsaturated fatty acids (MUFA), palmitoleic acid (C16:1) and oleic acid (C18:1), respectively (1). SCD1 is one of the key lipogenic enzymes that controls triglyceride (TG) biosynthesis, through its substrate availability, i.e. MUFA (2,3). Studies from SCD1 knock-out mice models have demonstrated that the absence of SCD1 protects mice from obesity and increases insulin sensitivity due to improved insulin signaling (4, 5). In vivo SCD1 inhibition by pharmacological agents improved insulin sensitivity in an insulin-resistant rat model, while, SCD1 deficiency, under leptin-resistant conditions failed to show improved insulin sensitivity though ameliorated obesity (6,7). In addition, a study on morbid obese human subjects has shown an association between SCD1 expression and insulin resistance markers (8).

Recent understanding of vitamin A metabolism has shed light on the various metabolic conditions including obesity, metabolic syndrome, and insulin resistance, wherein vitamin A and its metabolites play a key role in the control and/or development of diseases and associated complications, in addition to their wide range of biological functions (9). Further, vitamin A and its metabolites are known to regulate SCD1 at the transcriptional level and suppress its activity (10,11). Previously, we have reported in an euglycemic obese rat model (WNIN/Ob) that the feeding of a vitamin A-enriched diet impaired the development of obesity through other mechanisms, independent of SCD1 (12). Further, when young obese rats are challenged with an identical diet, it has not only ameliorated obesity, but also improved insulin
sensitivity through muscle protein tyrosine phosphatase 1B (PTP-1B) regulation accompanied by decreased SCD1 protein expression (13).

With this background, here we tested the hypothesis that chronic consumption of vitamin A-enriched diet (129 mg vitamin A/kg diet) ameliorates hyperglycemia and glucose-intolerance, which are associated with SCD1 regulation of muscle and liver, by employing a glucose-intolerant obese rat model of WNIN/GR-Ob strain. Although candidate gene(s) of obesity in this strain of rats are yet to be identified, obese rats display impaired glucose tolerance along with other features of metabolic syndrome, such as increased body weight, adiposity, dyslipidemia, hyperinsulinemia, hyperleptinemia, and high plasma high-density lipoprotein-cholesterol levels, as compared with their age-and sex-matched lean counterparts (14).

**MATERIALS AND METHODS**

**Chemicals and materials used**

All the chemicals used were of analytical grade. TG and glucose assay kits (BioSystems, Barcelona, Spain) were procured. Radioimmunoassay-based insulin assay kit from Board of Radiation and Isotope Technology, Mumbai, India and PTP-1B assay kit (Biovision Inc., Milpitas, CA, USA) were purchased. Primary antibodies such as rabbit anti-vacun carboxykinase (PEPCK), were purchased from fatty acid binding protein (FABP), and phosphoenolpyruvate carboxykinase (PEPCK), were purchased from Cell Signaling Technology, Inc. (Danvers, MA, USA). pIRβ (Ytr1162/1163), Glucose transporter type 4 (GLUT4), PTP-1B, AMP-activated protein kinase-α (AMPKα), pAMPKα (Thr172), fatty acid binding protein (FABP), and phosphoenolpyruvate carboxykinase (PEPCK), were purchased from Santa Cruz Biotechnology, Inc. (Dallas, TX, USA). β-Acid monoclonal antibody, secondary antibodies, and protease inhibitor cocktail were purchased from Sigma-Aldrich Co. (St. Louis, MO, USA). Phosphatase inhibitor cocktail, PhosSTOP, was obtained from Roche diagnostics GmbH, Mannheim, Germany. WNIN/GR-Ob strain rats were obtained from the National Centre for Laboratory Animal Sciences, National Institute of Nutrition, Hyderabad, India.

**Animals and experimental design**

Adult (30-week old), male lean and obese rats of WNIN/GR-Ob strain were broadly divided into two groups, A and B, each consisting of 12 lean and 12 obese rats (with impaired glucose tolerance traits), respectively and further divided into two subgroups (A-I, A-II, B-I, and B-II) consisting of 6 rats each. Subgroups A-I and B-I received the stock diet having 2.6 mg of vitamin A/kg diet, while subgroups A-II and B-II received the vitamin A-enriched diet (129 mg of vitamin A/kg diet as retinyl palmitate) for a period of 14 weeks. The composition of diets was identical in all ingredients, except for the vitamin A content. Diet and fatty acid compositions are given in Table 1. Animals were housed individually at ambient temperature, 22.0±10°C, relative humidity of 50–60%, 12-h/12-h light-dark cycle, and cared for in accordance with the principles of the guide to the care and use of experimental animals. The study was approved by the Institutional Animal Ethics Committee of the National Institute of Nutrition, Hyderabad, India (Registered number: 154/1999/CPCSEA & Study number: IAEC/Proj/05).

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Food and water were provided ad libitum. At the end of the experimental period, blood was drawn from the retro-orbital sinus in ethylenediaminetetraacetic acid (EDTA)-coated tubes, after a 12-h fast and the rats were killed by cervical dislocation. Various tissues were excised, immersed in ice-cold saline, weighed, and rapidly frozen in liquid nitrogen and stored at −80°C, until analysis.

**Oral glucose tolerance test and biochemical analyses**

At the end of 12th week, oral glucose tolerance test (OGTT) was performed by administering 3 g glucose per kg body weight to overnight-fasted rats. Blood was drawn at different time intervals and analyzed for glucose and insulin.

Plasma TG, glucose, and insulin levels were quantified, using commercially available kits. PTP-1B activity was measured according to the manufacturer’s instructions. Tissue glycogen content was determined by the

| Table 1. Diet and ground-nut oil fatty acid composition |
|---------------------------------------------------------|
| **Diet composition (g/kg)** | **Major fatty acid composition of ground-nut oil (%)** |
| Wheat flour | C16:0 | 11.0 |
| Roasted Bengal gram flour | C18:0 | 4.1 |
| Skim milk powder | C18:1 | 42.9 |
| Casein | C18:2 | 37.8 |
| Refined groundnut oil | C18:3 | 0.3 |
| Mineral mixture<sup>1</sup> | C20:0 | 1.8 |
| Vitamin mixture<sup>2</sup> | 5 | 1.95 |

Vitamin A of 2.6 mg and vitamin D of 10 μg in the form of vitamin A-enriched diet (129 mg of vitamin A/kg diet as retinyl palmitate) was added per kg diet, Additionally, for vitamin A-enriched diet, 126.4 mg of vitamin A was added per kg diet as retinyl palmitate.

<sup>1</sup>Composition of mineral mixture (g/kg): (d)-α-tocopherol acetate 24.0 g, menadione 0.3 g, thiamine 2.4 g, riboflavin 1.0 g, pyridoxine 1.2 g, niacin 2.0 g, pantothenic acid 2.4 g, cyanocobalamine 1.0 μg, folic acid 0.2 g para-aminobenzoic acid 20 g, biotin 0.08 g, inositol 20 g, and choline chloride 200 g.

<sup>2</sup>Composition of vitamin mixture (g/kg): vitamin A of 2.6 mg and vitamin D of 10 μg in the form of vitamin A-enriched diet (129 mg of vitamin A/kg diet as retinyl palmitate) was added per kg diet, Additionally, for vitamin A-enriched diet, 126.4 mg of vitamin A was added per kg diet as retinyl palmitate.
method of Passonneau and Lauderdale (15).

**Fatty acid composition by gas-liquid chromatography (GC)**
Gastrocnemus muscle total lipids were extracted, used for TG estimation and fatty acid methyl esters preparation, which were analyzed by GC coupled with a flame ionization detector as described earlier (16). The fatty acid desaturase activity index was calculated as the ratio of product to substrate i.e. C16:1/C16:0, C18:1/C18:0, C20:4/C18:2, and C20:4/C20:3 for delta9-16desaturase (D9-16D), delta9-18desaturase (D9-18D), delta6 desaturase (D6D), and delta5 desaturase (D5D), respectively.

**Protein expression by Western blot**
Briefly, liver or muscle tissues were homogenized in Tris buffer containing 250 mM sucrose, 10 mM Tris (pH 7.4), 1 mM EDTA, and 1 mM dithiothreitol supplemented with protease and phosphatase inhibitor cocktail and subjected to differential centrifugation. From the various cellular fractions, Western blotting was performed, using protein targeted antibodies according to a standard protocol (16). β-Actin was used as the loading control. Images were analyzed by the Image J 1.46r software (National Institutes of Health, Rockville, MD, USA).

**Gene expression by quantitative real time-polymerase chain reaction (qRT-PCR)**
Total RNA from liver and gastrocnemus muscle was isolated, and reverse transcription reaction was performed as described earlier (16). qRT-PCR was performed as suggested by the manufacturer’s instructions, with a LightCycler480 Real Time-PCR system (Roche diagnostics GmbH), using pre-validated probes for rat (UPL probes, Roche diagnostics GmbH) and gene-specific primers (Table 2). Endogenous expression of acidic ribosomal phosphoprotein was used to normalize the expression data and relative expression levels.

**Statistical analysis**
Data are expressed as means±standard error of the mean (SEM). Statistical significance was determined by one-way ANOVA, with post-hoc least significant difference test and Pearson’s correlation analysis was performed. *P*≤0.05 was considered significant. IBM SPSS statistics 19.0 software (IBM Company, Armonk, NY, USA) was used for the statistical analysis.

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### Table 2. Gene-specific primers used for qRT-PCR analysis

| No | Gene | Forward (5'→3') | Reverse (5'→3') |
|----|------|-----------------|-----------------|
| 1  | ACL  | CTTTGGGCGTGAGGCATA | CCACCTTTG6CATCCAGGT |
| 2  | β-Actin | CCCCAGATGCAACCTTCT | CGTCATCCATGGCAGACCT |
| 3  | ADH1 | ATCCATTTCCTGCCCACA | GCCGCTTTG6CACCAGAGG |
| 4  | ADH4 | ATGATATTGGGCCGTTCTGT | ACCAGGTTTGGGACAGAGTC |
| 5  | ADH7 | CAGACACAA6AACCCGAAAA | GGGACAG6C6CTTTG6C6CATTA |
| 6  | Adiponectin | TGGTCACAATGGGATACCG | CCCTTAGG6ACCAAGAACCT |
| 7  | AKT-1 | AACGACGTAGCCATTGTGA | CCACATCTCTG6AGGAGAA |
| 8  | Aldh1a1 | CAAGCTG6CTGACTTTAGGA | CCACCATGTG6T6CTTC |
| 9  | Aldh1a2 | TCTCATG6GTGTCCTG6CA | TCTGAG6CATTG6TG6CGG |
| 10 | Aldh1a3 | GAACG6C6GAG6G6CAAGAA | GGTGAG6C6C6CATC6C66AG |
| 11 | ARPP | GATGCCCAGGG6GAGAGA | CACATTG6AAGT6T6GG |
| 12 | CYP26A1 | GAGAGAGGAGGAGAGGAGGAGGATG6ATA | GGCTGCAGTG6CTG6TAGT |
| 13 | CYP26B1 | AGCG6CAGAGGAGATG6CCA | GCATG6C6G6CG6GAAGATCA |
| 14 | FABP-L | CTTCTCCCGCAAGATC6CA | T6CCCTCTGTGAG6T6G6T6G |
| 15 | GLUT4 | TGCAG6TGCTGTGCTTTT | CCAGTCATC6CTGG6T6G |
| 16 | GSK-3β | ATCA6G6G6CAC6ACATCTGG6GAC | ACGCT6C6ACAG6G6G6GAT |
| 17 | GYS | CTGG6CG6C6AC6AGCTAG | GAG6C6C6T6CTG6CC6T |
| 18 | INSIG1 | ATGTAT6C6GGGTCCTTG6T6T | T6C6GAT6C6A6GTC66CA |
| 19 | IRβ | CAGAGA6A6ACTC6TCTG6G | TTT6AG6G66GAC6CTGG6T6T |
| 20 | RARα | TTGGAT6G6GGCT6CAAGAGC | AGG6C6G6CG6CAT6ACT |
| 21 | RARγ | TCCAG6C6CTTTAC6C66AGGTC | T6C6A6T6G6GTGTC66C6AG |
| 22 | RBP4 | GTAG6G6CG6AG6G6GACG | TTGG66G6GTGTC66T6G |
| 23 | RXRα | ACA6T6G6GAG6CAAGAGC | GGGT6G6GAG66C6C6T6G |
| 24 | RXRβ | GTTC6CTG6CTG6C6CTC | GAG6C6G6GAC6ACTG6G |
| 25 | RXRγ | G6G6CAT6G6AG6AG6G66G6C | T6G6C6CAT6C6TCTG |
| 26 | SCD1 | GAAG6G6G6G6CAACG6GACAG | GGTG66G6CTG6TAG6G6AAG |
RESULTS

Vitamin A improved hyperglycemia and glucose-intolerance

Obese rats of WNIN/GR-Ob strain showed hyperglycemia as compared with their age- and sex-matched lean counterparts. At the end of the experiment, obese rats fed on vitamin A-enriched diet showed significant reductions in fasting plasma glucose, with a concomitant increase in fasting insulin levels as compared with their stock diet-fed control obese rats. On the other hand, no such effects were seen in lean rats (Fig. 1A and 1B). From the OGTT, plasma glucose and insulin levels were measured at various time points, and the calculated area under the curves clearly showed faster glucose clearance upon high vitamin A ingestion in obese rats, which was corroborated with increased insulin levels against obese rats fed on stock diet. However, no such changes were seen in lean counterparts maintained on identical dietary regimen (Fig. 1C and 1D). Further, vitamin A supplementation showed a negative correlation with plasma glucose levels at the $P<0.01$ (Pearson’s correlation).

Fig. 1. Effect of vitamin A on insulin sensitivity-associated parameters. (A, B) Fasting plasma glucose and insulin levels, respectively. (C, D) Oral glucose tolerance test-area under the curve (OGTT-AUC) for glucose and insulin, respectively. (E, F) Glycogen levels of liver and muscle, respectively. Values are means±SEM of 6 rats, except for OGTT-AUC, 4 rats were from each group. Data were analyzed by one-way ANOVA-with post-hoc least significant difference test. Groups bearing different letters (a-c) are statistically different at $P<0.05$. A-I & B-I, stock diet-fed lean and obese rats, respectively; A-II & B-II, vitamin A-enriched diet-fed lean and obese rats, respectively.
**Vitamin A induced intracellular signaling and glycogen accumulation**

The data revealed improvement in the hyperglycemic status of these obese rats. Therefore, we tested whether this resulted in the accumulation of glycogen content in liver and muscle. Obese rats fed on stock diet had higher glycogen content in the liver compared with their age- and sex-matched lean counterparts fed on identical diet.

Further, vitamin A-enriched diet feeding resulted in increased glycogen content of both phenotypes; however, this increase was only significant in obese rats (Fig. 1E). On the contrary, muscle glycogen content was significantly lower in the stock diet-fed obese rats compared to their lean counterparts. However, their levels improved upon vitamin A supplementation in obese rats. Surprisingly, the identical treatment resulted in decreased gly-

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**Fig. 2.** Effect of vitamin A on intracellular signaling pathway proteins and genes of muscle and liver. (A) Western blots of insulin signaling pathway proteins and ratio of phosphorylated to non-phosphorylated protein for glycogen synthase kinase-3β (GSK-3β) and glycogen synthase (GS). (B, C) Relative mRNA levels of insulin signaling pathway genes of muscle and liver, respectively. Values are means±SEM of 3–4 rats from each group. Data were analyzed by one-way ANOVA with post-hoc least significant difference test. Groups bearing different letters (a–c) are statistically different at \( P \leq 0.05 \). A-I & B-I, stock diet-fed lean and obese rats, respectively; A-II & B-II, vitamin A-enriched diet-fed lean and obese rats, respectively.
cogen content in lean rats compared with their stock diet-fed lean rats (Fig. 1F). Notably, Pearson’s correlation analysis showed a positive association between plasma insulin levels and glycogen content of both, liver and muscle at $P<0.003$ and 0.025 levels, respectively.

To understand the vitamin A-induced improvement in hyperglycemia and glucose intolerance, phosphorylation status of intracellular insulin signaling pathways of muscle was studied. In the stock diet-fed obese rats, basal muscle insulin signaling pathway protein phosphorylation (such as GSK-3β and GS) and GLUT4 (membrane to cytosol ratio; data not shown) levels were not significantly different from their identically-treated, age- and sex-matched lean counterparts. However, vitamin A supplementation significantly increased the ratio of pGSK-3β protein to non-phosphorylated GSK-3β protein, and thus, possibly increased the active form of GS (decreased phosphorylated-GS) (Fig. 2A), which was corroborated with concomitant glycogen accumulation in obese rats, without altering the ratio of membrane to cytosol GLUT4 (data not shown), and no such changes were seen in lean rats. Regardless of the treatment, other protein expression pathways including PTP-1B, FABP, PEPCK, and the ratio of pAMPK to AMPK remained unchanged among all groups (data not shown).

Further, we tested whether higher intake of vitamin A affects the transcript levels of these proteins in muscle or not. The data suggest that the basal expression of muscle GSK-3β, GS, GLUT4, and insulin-induced gene 1 (INSIG1) genes were significantly lower in obese rats, while AKT mRNA levels remained unaltered compared with their age- and sex-matched lean rats receiving the stock diet. Feeding the vitamin A-enriched diet to obese rats resulted in significant increases in GSK-3β, GS, GLUT4, and INSIG1 transcript levels compared with their stock diet-fed obese counterparts. However, no such effect was observed in identically-treated lean rats (Fig. 2B).

Unlike muscle, the baseline data of liver showed no significant changes in the expression of various insulin signaling pathway genes, such as AKT, GSK-3β, GS, IRβ, and INSIG1 between lean and obese phenotypes. On the other hand, vitamin A supplementation significantly up-regulated their expression levels in both the phenotypes (except INSIG1, whose mRNA levels increased significantly in lean rats) compared to their stock diet-fed controls (Fig. 2C). Further, regardless of treatment, the activity of PTP-1B (both in liver and muscle), one of the important regulators of insulin receptor phosphorylation, was comparable between the groups, which supports the unaltered ratio of muscle phosphorylated insulin receptor to insulin receptor observed in muscles among various groups (data not shown).

**Vitamin A on tissue TG levels and SCD1 regulation**
Feeding the vitamin A-enriched diet to obese rats significantly increased TG accumulation in the liver (17) and muscle, while in the lean rats, hepatic TG levels alone were significantly increased as compared with their stock diet-fed respective controls (Fig. 3A and 3B). To understand the SCD1 regulation (both at mRNA and protein) by high vitamin A-diet feeding, its expression levels were determined in liver and muscle. In the stock diet-fed obese rats, SCD1 mRNA levels were significantly higher in the liver and muscle compared to their age- and sex-matched lean counterparts. Chronic vitamin A supplementation significantly reduced the expression levels of the SCD1 gene in the liver, while augmented its transcript levels in the muscle of the obese phenotype; however, its levels remained unaltered in lean rats compared to their respective stock diet-fed counterparts (Fig. 3C and 3D).

In line with this, protein expression of SCD1 was also significantly higher in both the liver and muscle of the stock diet-fed obese rats compared to that of lean counterparts. Contrary to gene expression, vitamin A-enriched diet-fed obese rats displayed no change in hepatic SCD1 protein expression; however, muscle SCD1 protein levels showed a significant increase compared to their stock diet-fed obese counterparts. No changes in protein expression were observed in the lean phenotype (Fig. 3E and 3F).

**Fatty acid composition reflected SCD1 activity but decreased polyunsaturated fatty acid (PUFA) levels in both liver and muscle**
To identify whether the vitamin A-mediated regulation of SCD1 is reflected in fatty acid composition in these tissues or not, the fatty acid composition of total lipids was analyzed. Muscle fatty acid composition analysis showed a marked increase in MUFA [palmitoleic acid (C16:1) and oleic acid (C18:1)], and a significant reduction in SFA [stearic acid (C18:0)] and long chain PUFA [arachidonic acid (C20:4) and docosahexaenoic acid (C22:6)] with no detectable eicosapentaenoic acid (C20:5) in the obese phenotype compared to their age and sex-matched stock diet-fed lean counterparts. A similar trend was reflected even in total MUFA and PUFAs of muscle fatty acids. Vitamin A supplemented obese rats displayed increased MUFAs, palmitoleic acid (C16:1) and oleic acid (C18:1) although the latter was not statistically significant. Further, this treatment brought down the levels of stearic acid (C18:0) and other long chain PUFAs in obese rats, while in identically-treated lean counterparts, docosahexaenoic acid (C22:6) levels showed a significant reduction, compared to their respective stock diet-fed lean counterparts. Overall, the vitamin A-enriched diet resulted in significant reduction of total SFA, increased
MUFA in obese phenotypes, and no significant change in total PUFA content. However, the fatty acid composition of identically-treated lean rat muscle did not change (Table 3).

On the other hand, compared to lean rats, the liver fatty acid composition of obese rats showed higher SFA [palmitic acid (C16:0)], MUFA [oleic acid (C18:1)], and low levels of stearic acid (C18:0) and long chain PUFA such as eicosatrienoic acid (C20:3) and arachidonic acid (C20:4). Overall, hepatic total SFA and PUFA levels were significantly lower, while MUFA levels were significantly higher in obese rats receiving the stock diet compared to
their lean counterparts. Vitamin A-enriched diet feeding increased the levels of palmitic acid (C16:0), linoleic acid (C18:2), and α-linolenic acid (C18:3) in obese rats (and C16:0 alone in lean rats), but decreased stearic acid (C18:0) and long chain PUFA [eicosatrienoic acid (C20:3) and arachidonic acid (C20:4)] of both phenotypes compared with their respective stock diet-fed controls (Table 3).

**Table 3. Effect of vitamin A on the fatty acid composition of liver and muscle**

| Major fatty acids (%) | Liver |  |  |  |  |  |  |  |
|-----------------------|-------|-------|-------|-------|-------|-------|-------|-------|
|                       | Lean A-I | Lean A-II | Obese B-I | Obese B-II | Lean A-I | Lean A-II | Obese B-I | Obese B-II |
| C16:0                 | 19.6±0.5 a | 32.6±1.4 b | 26.4±2.0 c | 29.2±0.6 d | 23.5±1.1 a | 21.9±1.5 a | 22.8±0.8 a | 20.3±1.0 a |
| C16:1                 | ND     | 0.4±0.1 a | 5.7±0.8 b | 4.4±1.0 c | 2.1±0.4 a | 3.5±1.2 a | 7.3±1.3 b | 11.6±0.7 c |
| C18:0                 | 22.3±0.4 a | 17.8±0.7 a | 9.2±2.5 b | 6.2±0.8 c | 13.2±1.0 c | 11.1±1.6 c | 8.2±1.7 b | 3.1±0.3 a |
| C18:1                 | 9.6±1.1 a | 9.5±0.9 a | 26.3±2.5 b | 24.2±1.1 b | 20.3±1.9 a | 25.7±3.8 b | 29.2±3.3 b | 35.9±1.3 c |
| C18:2                 | 19.9±0.7 a | 20.6±2.1 a | 18.9±3.0 a | 28.5±1.0 a | 24.6±1.3 b | 25.6±3.2 b | 23.1±1.5 a | 25.4±0.5 |
| C18:3(n-3)            | ND     | ND     | 0.4±0.1 b | 0.8±0.1 b | ND     | ND     | ND     | ND |
| C20:3                 | 1.5±0.2 b | 1.0±0.1 b | 0.7±0.2 ab | 0.3±0.1 a | 0.9±0.1 a | 0.9±0.2 a | 1.0±0.1 b | 0.7±0.1 |
| C22:0                 | ND     | ND     | ND     | ND     | ND     | ND     | ND     | ND |
| C20:4(n-6)            | 27.1±0.6 a | 18.2±0.8 b | 12.4±3.0 b | 6.4±1.0 a | 10.1±0.7 b | 7.5±1.1 b | 5.3±1.0 b | 1.8±0.3 a |
| C20:5(n-6)            | ND     | ND     | ND     | ND     | ND     | ND     | ND     | ND |
| C24:0                 | ND     | ND     | ND     | ND     | 0.2±0.0 a | 0.2±0.0 b | 0.3±0.1 b | 0.2±0.1 |
| C22:5                 | ND     | ND     | ND     | ND     | 1.2±0.2 ab | 0.9±0.2 b | 1.0±0.2 b | 0.4±0.1 a |
| C22:6                 | ND     | ND     | ND     | ND     | 2.9±0.2 a | 1.9±0.3 b | 1.4±0.3 b | 0.4±0.1 a |
| Total SFA             | 41.9±0.7 a | 50.3±2.0 b | 35.5±2.4 a | 35.4±1.4 a | 37.1±2.0 a | 33.4±3.1 a | 31.7±2.4 a | 23.9±1.4 a |
| Total MUFA            | 9.6±1.1 a | 9.9±0.9 a | 32.1±3.5 b | 28.6±1.7 b | 22.5±2.0 a | 29.2±4.6 ab | 36.5±4.4 b | 47.5±0.8 b |
| Total PUFA            | 48.5±1.1 c | 39.7±1.3 c | 31.9±2.0 c | 35.3±0.5 a | 40.5±1.1 c | 37.6±2.4 bc | 32.0±2.5 ab | 28.8±0.3 a |

Values are means±SEM of 4 rats from each group. Data were analyzed by one-way ANOVA with post-hoc least significant different test. Groups bearing different letters (a–d) are statistically different at P<0.05 compared within the same tissue. ND: Non-detectable.

**Vitamin A on fatty acid desaturase activity indices**

Increased hepatic D9-18D and decreased D6D were observed in obese rats compared to their stock diet lean counterparts. Further, the desaturase activity index of the latter was significantly reduced in both lean and obese rats by vitamin A-enriched diet consumption, while other desaturase activities were not affected compared to their stock diet-fed respective controls (Fig. 4A).

Muscle fatty acid desaturase activity indices for D9-16D and D9-18D were significantly higher, whereas D6D and D5D activity indices were significantly lower in obese rats receiving the stock diet compared to their age- and sex-matched lean counterparts. Vitamin A supplementation increased the activity indices of D9-16D and D9-18D, and decreased D6D and D5D indices significantly in the obese phenotype. On the other hand, in lean rats, the D5D activity-index showed a significant reduction compared to stock diet obese and lean rats respectively (Fig. 4B).

**Vitamin A meagerly regulated its metabolic pathway transcripts of liver and muscle**

To understand the role of vitamin A metabolic pathway in improved hyperglycemia and glucose clearance, some of the transcript levels were analyzed in both liver and muscle. Hepatic vitamin A metabolic pathway gene expression analysis revealed that basal expression levels of alcohol dehydrogenase (Adh) namely, Adh1, Adh4, and Adh7, and aldehyde dehydrogenase (Aldh), such as Aldh1a2 and Aldh1a3 remained unaltered, while Aldh1a1 gene was up-regulated in obese rats compared to their age- and sex-matched lean counterparts. Although none of these gene expression levels were altered by vitamin A-enriched diet feeding, Aldh1a2, cytochrome p26 (CYP26)A1, and CYP26B1 mRNA expressions were up-regulated in both lean and obese rats compared to their respective control groups receiving the stock diet.

Further, the isoforms of retinoic acid receptor (RAR) and retinoid X receptor (RXR) mRNA levels were comparable among all the groups, except, RXRα mRNA expression was down-regulated in lean rats receiving the vitamin A-enriched diet compared with their stock diet-fed lean counterparts (Fig. 4C).
In muscle, among the various isoforms, Adh1 and Aldh1a1 expression levels were detectable and no significant changes were observed between phenotypes or treatments, except the basal expression of retinoic acid catabolizing enzyme CYP26A1, which was under-expressed. On the other hand, CYP26B1 was over-expressed in the stock diet-fed obese rats compared to their age- and sex-matched lean counterparts. Higher basal expression levels for RARα and lower expression levels of RXRγ were observed in the stock diet-fed obese rats compared with their lean counterparts. Vitamin A supplementation augmented RXRβ expression only in the obese phenotype, without affecting other genes in both phenotypes (Fig. 4D).

**DISCUSSION**

SCD1 is the key lipogenic enzyme responsible for the biosynthesis of MUFA and thereby regulates TG and energy homeostasis. Its role in insulin signaling and glucose homeostasis has been well demonstrated through SCD1 deficiency, using both *in vivo* and *in vitro* models (5-7,18). Contrary to many of these studies, the hepatic SCD1 protein expression data did not correlate with either circulatory or hepatic TG levels, which suggests that, only under steady state condition, the activity/level of SCD1 expression may correlate with fatty acid synthesis and/or TG accumulation. In addition, other factors, including substrate availability, synthesis, secretion, clearance, and...
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turnover-rate influence TG levels/accumulation, independent of SCD1. Unaltered MUFA levels and fatty acid desaturase activity-indices, particularly D9-16D and D9-18D of liver are suggestive of unaffected SCD1 protein status (and possibly its activity) due to increased vitamin A consumption. On the other hand, the fatty acid composition showed high levels of MUFA (C16:1 and C18:1), the functional indicator of SCD1 activity, as a result of SCD1 over-expression in the stock diet-fed obese rats, which are in concurrence with our previous findings, with regard to the high MUFA content of obese rat liver.

Unlike liver, SCD1 expression (both protein and gene) of muscle was positively associated with elevated TG levels and corroborated with increased MUFA levels (C16:1) and D9D activity-indices of muscle. Further, the study of Schenk et al. (19) has shown amelioration of fatty acid-induced insulin resistance in obese human subjects through increased TG accumulation in skeletal muscle in response to acute exercise. However, still it is debatable whether increased TG levels are due to SCD1 over-expression or increased uptake of circulatory lipids or both. Previously, Samuel et al. (20) have demonstrated SCD1 induction by RA and the involvement of nuclear transcription factors retinoic acid receptor/retinoid X receptor (RAR/RXR) and DNA response elements 1 (wherein RXR homodimer or RXR/ peroxisome proliferator-activated receptor heterodimer may bind) through gel mobility shift assay, and thus speculated the presence of one or more retinoic acid response elements in the promoter of SCD1. Further, the study of Repa et al. (21) has demonstrated the transcriptional regulation of SCD1 through sterol regulatory element binding protein-1c (SREBP-1c), the master regulator of lipid metabolism. In the current study, vitamin A metabolic pathway transcript levels did not go hand-in-hand with the observed SCD1 expression of liver and muscle. However, feeding of the vitamin A-enriched diet resulted in decreased hepatic SREBP1 mRNA levels in obese rats, while remained unaltered in lean rats compared to their respective controls, which is in agreement with the expression of hepatic SCD1. Thus, we postulate that vitamin A-mediated regulation of SCD1 may involve the SREBP1-dependant pathway, at least in the context of the present animal model, which partly explains the differential expression levels of the SCD1 gene in lean and obese phenotypes.

Vitamin A-supplemented diet feeding to glucose-intolerant obese rats differentially regulated SCD1 (unaltered in liver and elevated in muscle) expression, normalized circulatory glucose levels, and improved glucose clearance rate with a concomitant increase in insulin levels. It is well known that insulin stimulates tissue glycogen synthesis and thereby regulates glucose homeostasis (22). The current data on increased tissue glycogen (both liver and muscle) contents and positive correlation between plasma insulin and tissue glycogen go in favor of our hypothesis that vitamin A enhances glucose clearance and its storage as glycogen, thereby improving hyperglycemia, glucose tolerance and thus, suggesting improved insulin sensitivity. This derives support from the observed over-expression of adiponectin mRNA, while no change in retinol binding protein 4 in white adipose tissue (data not shown) was observed in vitamin A-challenged obese rats. These observations are also in line with earlier reports, which have shown a positive association between glycogen content and insulin sensitivity (23,24). Intracellular insulin signaling pathway of muscle revealed that the increased glycogen accumulation by vitamin A in obese rats was associated with decreased phosphorylation of glycogen synthase and thereby keeping it in its active form through regulation of glycogen synthase kinase, an upstream inhibitor of glycogen synthase, without changing GLUT4 protein expression.

On the other hand, the glycogen synthesis pathway was not affected, but there was a drastic reduction in the muscle glycogen content of lean rats receiving high vitamin A diet. Liver glycogen primarily plays the key role in maintaining glucose levels during starvation, and muscle glycogen is utilized, when liver stores are exhausted (25, 26). Therefore, the muscle glycogen reduction may not be due to increased degradation to maintain glucose homeostasis, especially in the presence of adequate liver glycogen stores, which was comparable to that of the control group. Further, it is possible that increased muscle glucose oxidation (which accounts for nearly 90% of glycolytic flux) through the activation of pyruvate dehydrogenase enzyme complex led to the higher demand for glycogen breakdown, particularly in the high vitamin A-challenged lean rats, resulting in decreased muscle glycogen content. Previously, vitamin A-induced activation of protein kinase Cδ, resulting in the activation of pyruvate dehydrogenase complex and increased oxidation has been demonstrated in vitro cell lines (27). Therefore, the involvement of such pathway may be considered, which partly explains the reduction of muscle glycogen at least in the high vitamin A-fed lean rats. However, further studies are needed to understand the glucose partitioning mechanisms of muscle in both phenotypes to explain the observed differences in the utilization of primary energy source i.e. glucose. In addition, the vitamin A-induced over-expression of insulin signaling pathway/effector genes in both liver and muscle supports the fact that vitamin A regulates insulin signaling not only at phosphorylation, but also at the transcriptional level. Further, the role of other insulin signaling pathway proteins such as PTP-1B, insulin receptor, and AMPK-phosphorylation status seem to be negligible in this glucose-intolerant obese rat model.

The association between TG accumulation and insulin
resistance appears to be complex with the advent of reports stating that hepatic TG accumulation in fact is a protective mechanism to counteract lipotoxic effects of excess non-esterified fatty acid (NEFA) and hence is not associated with or may not aggravate insulin resistance (28). Thus, it appears that vitamin A-induced TG accumulation of liver and muscle alleviates lipotoxicity and insulin resistance due to high levels of circulatory NEFA and/or TG, wherein decreased hepatic lipogenesis with no defective very-low-density lipoprotein secretion and/or fatty acid oxidation is observed (17).

Importantly, long chain PUFA, particularly arachidonic (C20:4) acid levels of liver and muscle were significantly lower in the obese phenotype, which were further reduced by high vitamin A consumption. In muscle, besides arachidonic acid, other long chain PUFA levels were decreased in the high vitamin A-fed obese rats. Similar effects were seen even in the identically-fed lean phenotype. As the source of dietary fat for all groups was the same, we speculate that the conversion of linoleic acid (C18:2) to arachidonic acid (C20:4) is affected due to the inhibitory action of vitamin A on fatty acid desaturases, particularly D6D and D5D (as reflected by their activity indices), which is in line with previous findings on vitamin A and its metabolite retinoic acid-mediated regulation of D5D (29). Arachidonic acid (C20:4) deficiency in streptozotocin-induced diabetic rat model and low levels in muscle of obese zucker rats (fa/fa) have been previously reported by Holman et al. (30) and Voss et al. (31). The current study could not relate the changes in long chain PUFA content, insulin signaling, and glycogen accumulation and their molecular link. Unlike other effects that are very specific to the obese phenotype, decreased arachidonic acid (C20:4) in response to vitamin A supplementation was seen in both lean and obese rats. This observation warrants further investigations to address the regulation of arachidonic acid (C20:4) by the bio-synthetic pathway of vitamin A, its biological significance and relevance to clinical research, especially in various chronic diseases including cancer, wherein vitamin A and its metabolites are of great therapeutic potential. In conclusion, chronic feeding of vitamin A-enriched diet to glucose-intolerant obese rats improved hyperglycemia and glucose intolerance, possibly by enhancing glycogen accumulation through the regulation of intracellular signaling and metabolic pathways of muscle and liver, but not associated with SCD1.

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**AUTHOR DISCLOSURE STATEMENT**

The authors declare no conflict of interest.

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