PROTECTIVE EFFECT OF CAMEL MILK AS ANTI-DIABETIC SUPPLEMENT: BIOCHEMICAL, MOLECULAR AND IMMUNOHISTOCHEMICAL STUDY

Ahmed A. Mansour1,2*, Mohammed A. Nassan3, Osama M. Saleh1,4, Mohamed M. Soliman5,6

1Medical Biotechnology Department, Faculty of Applied Medical Sciences (Turbah), Taif Univ., KSA. 2Genetics Department, Faculty of Agriculture, Ain Shams University, Cairo, Egypt. 3Pathology Department, Faculty of Veterinary Medicine, Zagazig University, Zagazig, Egypt. 4National Centre for Radiation Research and Technology (NCRRT), Egyptian Atomic Energy Authority (EAEA), Nasr City, Cairo, Egypt. 5Biochemistry Department, Faculty of Veterinary Medicine, Banha University, Banha, Egypt. 6Medical Laboratories Department, Faculty of Applied Medical Sciences (Turbah), Taif University, KSA.

*Corresponding Author E-mail: Ahmedmansour292@gmail.com

Abstract

Background: Diabetes is a serious disease affects human health. Diabetes in advanced stages is accompanied by general weakness and alteration in fats and carbohydrates metabolism. Recently there are some scientific trends about the usage of camel milk (CM) in the treatment of diabetes and its associated alterations. CM contains vital active particles with insulin like action that cure diabetes and its complications but how these effects occur, still unclear.

Materials and Methods: Seventy-five adult male rats of the albino type divided into five equal groups. Group 1 served as a negative control (C). Group 2 was supplemented with camel milk (CM). Diabetes was induced in the remaining groups (3, 4 and 5). Group 3 served as positive diabetic control (D). Group 4 served as diabetic and administered metformin (D+MET). Group 5 served as diabetes and supplemented with camel milk (D+CM). Camel milk was supplemented for two consecutive months. Serum glucose, leptin, insulin, liver, kidney, antioxidants, MDA and lipid profiles were assayed. Tissues from liver and adipose tissues were examined using RT-PCR analysis for the changes in mRNA expression of genes of carbohydrates and lipid metabolism. Pancreas and liver were used for immunohistochemistry examination using specific antibodies.

Results: Camel milk supplementation ameliorated serum biochemical measurements that altered after diabetes induction. CM supplementation up-regulated mRNA expression of IRS-2, PK, and FASN genes, while down-regulated the expression of PEPCK to control mRNA expression level. CM did not affect the expression of PEPCK gene. On the other hand, metformin failed to reduce the expression of CPT-1 compared to camel milk administered rats. Immunohistochemical findings revealed that CM administration restored the immunostaining reactivity of insulin and GLUT-4 in the pancreas of diabetic rats.

Conclusion: CM administration is of medical importance and helps physicians in the treatment of diabetes mellitus.

Keywords: Camel milk, Diabetes, Gene expression, Immunohistochemistry.

Abbreviations: STZ: Streptozotocin; MET: metformin; CM: Camel milk; GLUT4: glucose transporter4; FASN: fatty acid synthases; CPT-I: carnitine palmitoyl transferase I; PEPCK: phosphoenolpyruvate carboxykinase; PK: pyruvate kinase; IRS-2: Insulin receptor substrate 2; GAPDH: glyceraldehyde-3-phosphate dehydrogenase; TID: type-1-diabetes; T2D: type-2-diabetes; TC: total cholesterol; TG: triglycerides; HDL: high-density lipoproteins; CAT: catalase; GR: glutathione reductase; SOD: superoxide dismutase; MDA: malondialdehyde; GPT: glutamic-pyruvic transaminases; GOT: glutamic-oxaloacetic transaminases; PBS: phosphate-buffer saline.

Introduction

Diabetes mellitus type 1 (T1D) is a complex disease resulting from the interplay of genetic, endocrine, epigenetic, and environmental factors (Hakonarson and Grant 2011). Worldwide, the T1D represents an increasing global public health burden, and the incidence of T1D among humans has been rising (Forlenza and Rewers 2011). Over the last few decades, there has been an overall increase in the incidence of T1D of ~3% to 5% per year, and it is estimated that there are ~65 000 new cases per year in children, 15 years old (Borchers et al 2010). Moreover, the number of peoples with diabetes will rise to 500 million within the next 20 years (Malik et al 2012). Africa will experience a largest increase in the next generation.
for diabetes incidence. But till now Middle East is characterized with high incidence of diabetes but Africa still the most affected (Al-Baghihi et al 2011).

Failure to control insulin level in our bodies is the leading cause of diabetes. Only 5%-10% of diabetic patients do not produce insulin because of complete destruction in pancreatic β-cells that results in hyperglycemia and ketoacidosis in type-1 diabetic patients (Gaba et al 2015). Type 2 diabetes (T2D), a non-insulin-dependent diabetes mellitus, is associated with insulin resistance as cells of the body can’t respond to insulin. Accidental drug treatments such as glucocorticoids, β-adrenergic agonists, statins, Streptozotocin (STZ), and alloxan induce diabetes. STZ and alloxan are used to induce diabetes in rodent lab animal models (Eleazu et al 2013). The pancreatic β-cells take up the STZ via the plasma membrane GLUT2 transporter2, resulting in inflates, and ultimately irreversible destroys the pancreatic β-cells (Eleazu et al 2013; Ikebukuro et al 2002; Akbarzadeh et al 2007; Muhammad and Syed 2010; Sboui et al 2010; Sharma et al 2012). STZ treatment induces T1D (Muhammad and Syed 2010), and may induce T2D (Arulmozhhi et al 2004; Martha et al 2009).

Camel milk has valuable therapeutic effects on treatment of wounds as it enhance wound healing and increase immune cell proliferation and chemotaxis state of experimental animals (Badr 2012 and 2013; Badr et al 2011 and 2012). Presence of peptides and proteins in camel’s milk are the cause of its biological in digestion, absorption, growth and immunity (Korhonen and Pihlanto 2001). The most important uses of camel’s milk are as drug against autoimmune diseases, dropsy, jaundice, splenomegaly, asthma, anemia, piles and diabetes with antimicrobial and antitoxic effects (Soliman et al 2015). The improvement in immune function using dietary antioxidants as camel whey can play an important role in the prevention of many human diseases and diabetic complications. Dietary whey suplementations may improve wound healing by increasing GSH synthesis and cellular antioxidant defense (Velioglu Ogünç et al 2008). As known, camel milk contains insulin-like peptides and other substances that modulate glucose level (Mohamad et al 2009; Zagorski et al 1998; Agrawal et al 2003).

There is a growing worldwide trend that camel milk consumption helps in prevention and control of diabetes (Malik et al 2012). The presence of peptides in camel’s milk exhibits its biological activities that have a valuable effect on digestion, absorption, growth and immunity (Korhonen and Pihlanto 2001) and antitoxic effect against some heavy metals and drugs (Al-Hashem 2009; Afifi 2010; Khan and Alzohairy 2011). CM is beneficial in reducing the dose of insulin needed to induce glycemic control and improves fasting blood glucose, glycosylated hemoglobin, serum anti-insulin antibodies, urinary albumin excretion, and mass body index (Agrawal et al 2009; Mohamad et al 2009). Regular consumption of CM for few months improved the condition of diabetic patients and experimental animal models (Sboui et al 2010; Mohamad et al 2009; Agrawal et al 2005 and 2011). Most published data about the way by which any treatment may affect the response of diabetic patients and experimental animal models is not completely explained (Mima et al 2008; WHO 2011; Hesham et al 2012). Therefore, this study focused on examining the changes in biochemical alterations during diabetes, changes in gene expression of lipids and carbohydrate metabolism and how to prevent or modulate diabetes incidence by camel milk supplementation. Moreover, the histopathological changes using immunohistochemical stains for pancreas was used to clarify the medical importance of CM supplementation in treatment of diabetes.

**Materials and Methods**

**Camel's milk preparation**

Camel’s milk samples were collected daily in the morning from camel farm in Turabah governorate, Mecca principedom, KSA. Milk was collected from healthy camels (4 years old) by hand milking and kept in sterile screw bottles and kept under cooling conditions until transported to the laboratory. Rats supplemented with unpasteurized camel milk.

**Experimental Design**

Seventy-five adult male rats of the albino type, 8 weeks old, 200 gram body weight were purchased from King Fahd center for Scientific Research, King Abdel-Aziz University, Jeddah, KSA. Rats were kept for one week for acclimatization. All animal procedures were approved by the Ethical Committee Office of the dean of scientific affairs of Taif University, Saudi Arabia. Rats were divided into five groups (each group were handled in triplicate; 5 rats per one cage in triplicate; 15 rat per group). Group 1 as used as a control and received free access to food and water. Group 2 supplemented with camel milk (CM). Diabetes was induced in the remaining forty-five rats by intraperitoneal injection of STZ (60 mg/kg bw, Sigma-Aldrich St. Louis, MO, USA) based on protocol stated by Saleh et al (2016). To avoid the sudden death of the diabetic rats due to hypoglycemia, rats were supplemented with 5% glucose solution for the next 12 h following STZ injection. Diabetes was confirmed after 72 hrs of STZ injection by measuring blood glucose levels using commercially available glucose detector. Rats with glucose levels over 220 mg/dL considered diabetic and used for

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subsequent groups. The positive diabetic rats divided into groups 3, 4 and 5. Group 3 served as positive diabetic control (D). Group 4 diabetic and administered metformin (D+MET). Group 5 served as diabetes and supplemented with CM (D+CM). CM was given orally to group 2 and 4 at a dose of 250 ml/24 hours/15 rats based on study of Althnaian (Althnaian et al 2013). CM dose can be equated as 83.33 ml/kg bw daily for 2 consecutive months. The experiment was repeated three times each with five animals for reproducitively. Twenty-four hours after last administrations, all rats were sacrificed after anesthetization by diethyl ether inhalation. Blood, liver, and adipose tissues were collected. Liver and adipose tissues were kept in TRizol® reagent (Life Technologies, USA) at -80°C in a deep freezer for ribonucleic acid (RNA) extraction and maintained in 10% neutral buffered formalin (NBF) at room temperature for 24 hours for immunohistochemical study.

**Serum biochemical measurements**

Triglycerides (TG), total cholesterol (TC), and high-density lipoproteins (HDL) were measured using commercial spectrophotometric analysis kits (Bio-Diagnostic Company, Giza, Egypt). Insulin and leptin levels were measured using spectrophotometric commercial kits bought from Chrystal Chem Co, Downers Grove, IL 6051 USA. Glucose was measured using commercial glucose monitoring set (ACCU-Chek). Liver, renal biomarkers, antioxidants and MDA were measured spectrophotometrically as indicated in instruction manuals of each kit.

**RNA Extraction**

Total RNA was extracted from adipose and liver tissues samples as stated previously (Ahmed and Ahmed 2016). RNA concentration and purity determined by BIO-RAD SmartSpecPlus UV/Visible Spectrophotometer (USA). RNA integrity was examined by running through 1.5% denaturated agarose gel stained with ethidium bromide. Semi-quantitative RT-PCR analysis

Three micrograms of total RNA and 0.5 ng oligo dT primer (Qiagen Valencia, CA, USA) were used for cDNA synthesis. The total volume was adjusted up to 11 µl by sterilized DEPC water and incubated in the Bio-Rad T100TM Thermal cycler at 65°C for 10 min for denaturation. Then, 2 µl of 10X RT-buffer, 2 µl of 10 mM dNTPs and 100U Moloney Murine Leukemia Virus (M-MuLV) Reverse Transcriptase (Sib Enzyme. Ak, Novosibirsk, Russia) were added. The total volume was adjusted up to 20 µl by adding DEPC water. The mixture was then kept in BIO-RAD thermal cycler at 37°C for one hour, re-heated for 10 min at 90°C for enzyme deactivation. Specific primers for examined genes (Table 1) were designed using Primer3Plus online software (http://primer3plus.com/) and synthesized by Macrogen (Macrogen Company, Korea). 1 µl cDNA was mixed with 1 µl of 10 pm of each primer, and 12.5 µl PCR master mix (Promega USA).

The final volume was brought up to 25 ml using sterilized, deionized water. PCR was carried out using Bio-Rad T100TM Thermal Cycler with a cycle sequence at 94°C for 5 minutes one cycle, followed by 31 cycles (Table 1) each of which consists of denaturation at 94 °C for one minute, annealing at the specific temperature corresponding to each primer (Table 1) and extension at 72 °C for one minute with an additional final extension at 72 °C for 7 minutes. Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) mRNA was used as a reference. The PCR products were electrophoresed on 1.5% agarose gel, stained with 0.1μg/ml of ethidium bromide. PCR products were photographed and subjected to band intensity analysis using Image J software v. 1.47 (http://imagej.en.softonic.com/).

| Table (1): PCR conditions and primers sequence of examined genes. |
|---|---|---|
| Gene | Primer sequence (5' - 3') | Annealing Temp. | Band size |
| GAPDH | AGATCCACAACCGGATACATT | (F) | 52 | 309 bp |
| | TCCCTCAAGATTGTCAGCAA | (R) | | |
| PK | ATTGCTGTGACTGGATCTGC | (F) | 52 | 229 bp |
| | CCCGCACTGATGTTGATGATAG | (R) | | |
| PEPCK | TTTACTGGGAAGGCATCGAT | (F) | 52 | 236 bp |
| | TCCTGAGAACAGGGGCCAC | (R) | | |
| IRS-2 | CTACCCACTGACGCCCAAGAG | (F) | 55 | 151 bp |
| | CCAGGGATGAAAGCAGGACTA | (R) | | |
| FASN | CCAGAAGCCCAGACAGAGAAG | (F) | 61 | 345bp |
| | GACGCCAGTGTCTCCTCC | (R) | | |
| CPT-1 | TATGTGAGATGCTGTCTCC | (F) | 52 | 628 bp |
| | CTCGGAGAGCTAAGCTTGC | (R) | | |
Immunohistochemical analysis of pancreatic insulin

Rat Pancreas was fixed in 10% buffered neutral formalin then washed, dehydrated in ascending grades of ethyl alcohol then cleared in xylene, embedded in paraffin, sectioned at 5 micron thickness, deparaffinized, and then treated with 3% H2O2 for 10 min to block the peroxidases activity. Subsequently, the tissue samples were heated at 121°C in 10 mM citrate buffer for 30 min, blocked for 20 min in 5% normal serum. The pancreatic tissue was incubated overnight with rabbit polyclonal anti-insulin primary antibody (1:100; sc-9168; Santa Cruz Biotechnology, Inc., Dallas, TX, USA) or with GLUT4 Antibody (1:100; sc-53566; Santa Cruz Biotechnology, Inc., Dallas, TX, USA) in phosphate-buffered saline (PBS) at 4°C. After three extensive washes with PBS, the sections were incubated with a goat anti-rabbit IgG biotin conjugated secondary antibody (1:2,000; sc-2040; Santa Cruz Biotechnology, Inc.) for 20 min at 32°C. After further incubation with horseradish peroxidase-labeled streptavidin, antibody binding was visualized using diaminobenzidine, and the sections were counterstained with hematoxylin.

Statistical analysis

Data are presented as means ± standard error of means (SEM) for 3 independent experiments per each treatment. Results were analyzed using analysis of variance (ANOVA) and post hoc descriptive tests by SPSS software v 11.5 (SPSS, IBM, Chicago, IL, USA) with \( p<0.05 \) regarded as statistically significant.

Results

Analysis of serum chemistry in diabetic rats:

Effect of camel milk supplementation on glucose, insulin and leptin levels in diabetic rats

Diabetic rats supplemented with camel milk showed a decrease in glucose levels compared to diabetic rats (D). This is coincided with the decrease reported in metformin administered diabetic rats (D+MET) as a positive therapeutic group. The changes in glucose levels are mainly due to the increase in insulin secretion reported in camel milk administered diabetic rats (D+CM) as seen in the Table (2). Leptin levels were decreased in diabetic group and was increased in metformin diabetic rats (D + MET) while CM administration normalized and increased it in a way to increase peripheral glucose utilization and homeostasis (D+CM) as seen in table 2.

Table (2): serum changes in glucose levels, insulin and leptin in diabetic rats and after camel milk administration for 2 months

| Group                        | Glucose (mg/dL) | Insulin (µU/ml) | Leptin (ng/ml) |
|------------------------------|-----------------|-----------------|----------------|
| Control (C)                  | 83.0±9.7        | 35.3±2.3        | 12.6±3.1       |
| Diabetes (D)                 | 462.3±37.8*     | 8.3±0.5*        | 9.1±5.1*       |
| Diabetes + metformin (D+MET) | 120.0±10.1#     | 27.2±3.4#       | 17.2±5.3#      |
| Control + Camel milk (CM)    | 85.7±5.8        | 39.3±6.3        | 10.0±1.6       |
| Diabetes + Camel milk (D+CM) | 96.7±11.1#      | 33±2.3#         | 10.4±1.3#      |

Values are means ± standard error (SEM) for 3 independent experiments per each treatment. Values are statistically significant at *\( p<0.05 \) Vs. control; #\( p<0.05 \) Vs. diabetic group #\( p<0.05 \) Vs. diabetic metformin group.

Effect of camel milk supplementation on kidney and liver biomarkers in diabetic rats

Due to oxidative stress induced by diabetes, diabetic rats showed significant an increase in serum levels of urea, creatinine, GPT and GOT (Table 3). Administration of camel milk to diabetic rats (D+CM) normalized these changes. The results reported for camel milk in diabetic rats are the same reported for control positive metformin administered rats (D+MET).
Table (3): Serum changes in kidney and liver function parameters in diabetic rats and after camel milk administration for 2 months.

| Group                  | Urea (mg/dL) | Creatinine (mg/dL) | GPT (U/L) | GOT (U/L) |
|------------------------|--------------|--------------------|-----------|-----------|
| Control (C)            | 43.0±9.2     | 0.8±0.1            | 103.6±7.7 | 90.6±7.6  |
| Diabetes (D)           | 111.7±14.8*  | 2.4±0.2*           | 169.1±15.1* | 181.1±5.1* |
| Diabetes + metformin (D+ME) | 69.0±22.4# | 1.1±0.3# | 99.2±25.3# | 121.7±14.9# |
| Control + Camel milk (C+CM) | 40.7±5.8 | 0.9±0.2    | 86.0±2.5 | 96.0±16.3 |
| Diabetes + Camel milk (D+CM) | 66.2±8.5# | 0.7±0.02# | 103.7±24.2# | 70.7±13.0# |

Values are means ± standard error (SEM) for 3 independent experiments per each treatment. Values are statistically significant at *p<0.05 Vs. control; #p<0.05 Vs. diabetic group #p<0.05 Vs. diabetic metformin group.

Effect of camel milk supplementation on oxidative stress and antioxidants levels of diabetic rats

The changes in malondialdehyde (MDA) levels in diabetic rats showed an increase in MDA levels and CM administration normalized MDA levels in CM administered diabetic rats (Table 4). Catalase (CAT), glutathione reductase (GR), and superoxide dismutase (SOD) were decreased in diabetic rats. The percent decrease in antioxidant enzymes were about 71% compared to control group (C). Antioxidants levels were normalized in diabetic rats after CM supplementation (D+CM; table 4).

Table (4): Serum changes in MDA and antioxidants levels in diabetic rats and after camel milk administration for 2 months.

| Group                  | MDA (nmol/g tissue) | CAT (U/g tissue) | GR (U/g tissue) | SOD (U/g tissue) |
|------------------------|---------------------|------------------|-----------------|-----------------|
| Control (C)            | 4.5±0.1             | 31.4±2.0         | 9.4±0.4         | 11.5±0.3        |
| Diabetes (D)           | 28.5±1.8*           | 8.7±0.5*         | 3.1±0.1*        | 2.9±0.2*        |
| Diabetes + metformin (D+ME) | 14.1±3.8#     | 26.4±2.4#        | 7.5±1.0#        | 9.8±1.2#        |
| Control + Camel milk (C+CM) | 11.6±4.0  | 31.4±6.1         | 9.4±0.4         | 10.7±1.0        |
| Diabetes + Camel milk (D+CM) | 8.3±0.3    | 32.3±1.8#        | 9.8±0.3#        | 11.4±0.5#       |

Values are means ± standard error (SEM) for 3 independent experiments per each treatment. Values are statistically significant at *p<0.05 Vs. control; #p<0.05 Vs. diabetic group #p<0.05 Vs. diabetic metformin group.

Effect of camel milk supplementation on total cholesterol, triglycerides and high density lipoproteins in diabetic rats

A significant increase in the total cholesterol (TC), triglycerides (TG) and a decrease in high-density lipoproteins (HDL) levels were reported in diabetic rats. Camel milk administration ameliorated and normalized the increase in TC and TG levels in diabetic rats (D+CM). In parallel, CM supplementation increased HDL levels compared to diabetic group alone. The ameliorated effect induced by camel milk is relatively the same reported for metformin administered rats (D+MET) as seen in table 5.

Table (5): Serum changes in lipid parameters in diabetic rats and after camel milk administration for 2 months.

| Group                  | TC (mg/dL) | TG (mg/dL) | HDL (mg/dL) |
|------------------------|------------|------------|-------------|
| Control (C)            | 93.7±10.2  | 53.3±2.8   | 19.7±0.9    |
| Diabetes (D)           | 252.7±18.2* | 128.3±11.8* | 8.9±4.1*   |
| Diabetes + metformin (D+ME) | 161.2±21.8# | 85.7±9.1# | 16.2±0.6#  |
| Control + Camel milk (C+CM) | 104.7±19.2 | 77.7±9.8  | 20.3±0.9    |
| Diabetes + Camel milk (D+CM) | 100.0±6.6# | 72.3±0.5# | 15.7±2.2#  |

Values are means ± standard error (SEM) for 3 independent experiments per each treatment. Values are statistically significant at *p<0.05 Vs. control; #p<0.05 Vs. diabetic group #p<0.05 Vs. diabetic metformin group. (TC) total cholesterol, (TG) triglycerides, (HDL) high density lipoprotein.
Effect of camel milk supplementation on hepatic gene expression of carbohydrates and lipids metabolism in diabetic rats

The effect of camel milk on the expression of some genes related to carbohydrate and lipid metabolism altered during diabetes are shown in Figs (1 and 2). As shown in figure 1, CM administration alone did not affect the expression of PEPCK gene in diabetic group (D+CM). The mRNA expression of IRS-2 gene was down-regulated in diabetic rats (D) and increased in positive metformin diabetic rats (D+MET). CM administration increased significantly mRNA expression of IRS-2 in CM administered diabetic rats (D+CM) in parallel with results reported for metformin diabetic group. A similar change in mRNA expression of PK occurred and was the same like in IRS-2 gene. On the other hand, figure 2 shows that the expression of CPT-I was increased due to diabetic rats (D). Metformin administration failed to inhibit the up-regulation occurred in diabetic rats. Interestingly, CM administration restored the expression level of CPT-I to normal levels as compared to control group (C). Moreover, CM supplementation up-regulated the transcription of FASN mRNA in diabetic rats (D+CM) in same pattern reported for metformin diabetic rats (Figure 2).
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Figure 1: Semi-quantitative RT-PCR analysis of PEPCK, PK and IRS-2 mRNA expressions and their corresponding GAPDH in the rat liver tissue. RNA was extracted and reverse-transcribed (3 μg), and RT-PCR analysis was carried out for examined genes as described in the Materials and methods. Densitometric analysis was carried for 15 different rats per each group. Values are means ±SEM obtained from 5 rats in triplicate per group. *P<0.05 vs. control group, and #P<0.05 vs. diabetic group.

Figure 2: Semi-quantitative RT-PCR analysis of CPT-1 and FASN mRNA expressions and their corresponding GAPDH in the rat adipose tissue. RNA was extracted and reverse-transcribed (3 μg), and RT-PCR analysis was carried out for examined genes as described in the Materials and methods. Densitometric analysis was carried for 15 different rats per each group. Values are means ± SEM obtained from 5 rats in triplicate per group. *P<0.05 vs. control group, and #P<0.05 vs. diabetic group.

Immunohistochemical analysis of pancreatic insulin after camel milk supplementation in diabetic rats

The pancreas of control rats showed normal tissue architecture with regular insulin expression in the pancreatic β-cells (Fig. 3A). CM and metformin administered control rats (C+CM) and (C+MET) showed normal insulin expression in pancreatic beta cells (Figs. 3B & 3C respectively). The pancreatic tissue from diabetic rats (D) exhibited a reduction in the expression of insulin, with atrophy of the pancreatic β-cells (Fig. 3 D). Finally, the pancreas of CM administered diabetic rats (D+CM) exhibited regenerative restoration of normal expression of insulin in the pancreatic β-cells (Fig. 3 E). On the same line, pancreas of control rats (C) presented normal GLUT4 expression in the pancreatic tissues (Fig. 3F). CM and metformin administered control rats (C+CM) and (C+MET) showed normal GLUT-4 expression in pancreatic tissue (Figs. 3G & 3H respectively). However, the pancreatic tissue from diabetic rats (D) exhibited a reduction in the expression of...
GLUT4 (Fig. 3I) while the pancreas of the group (D+CM) exhibited restoration of normal expression of GLUT4 in the pancreatic tissue (Fig. 3J).

Figure 3: (A): Pancreatic tissue of a healthy control rat showing a normal architecture with normal expression of insulin in the β cells (magnification, x1200). (B): Pancreatic tissue of camel milk administered rat showing normal expression of insulin in the β cells. (C): Pancreatic tissue of metformin administered rat showing normal insulin expression in the β cells. (D): Pancreatic tissue of streptozotocin administered rat showing a reduction in the expression of insulin in the β cells, with atrophy of the β cells (magnification, x1200). (E): Pancreatic tissue of Streptozotocin administered rat treated with camel milk showing the restoration of normal expression of insulin in the pancreatic β cells (magnification, x1200). (F): the pancreas of the healthy control rats presented normal GLUT-4 expression in the pancreatic tissue (magnification, x150). (G): The pancreas of camel milk administered rats showed normal expression of GLUT-4 in the pancreatic tissue (magnification, x150). (H): The pancreas of metformin administered rats presented normal GLUT-4 expression. (I): the pancreatic tissue from the diabetic rats exhibited a reduction in the expression of GLUT-4 (magnification, x150). (J): the pancreas of the diabetic treated with camel milk exhibited restoration of normal expression of GLUT-4 in the pancreatic tissue (magnification, x150). Scale bar for photos from A to E is 50 μm & for photos from F to J is 100 μm.

Discussion

Traditional medicine is a common way used in public remedy; the desert Bedouins use different varieties of natural materials including animal products in treating diseases. A large number of articles mentioned the benefits of camel milk (Malik et al 2012; Hanieh et al 2015; Laila et al 2015). Our results showed that administration of camel milk reduced the levels of serum glucose and restored the average level of leptin in the treated diabetic rats because camel milk contains insulin nanoparticles which are protected from digestive tract proteases and absorbed directly in the intestine (Malik et al 2012; Al-Hashem 2009; Kamal et al 2007; Abd El-Aziz et al 2012). As known, the insulin contained in camel milk has same features as human insulin (Malik et al 2012). Moreover, the camel milk keeps its liquidity form in the environment of the low stomach pH. Camel milk has an advantage, that it contains 'insulin-like' small molecule substances that mimic insulin interaction with its receptor, for that, it can be suitable for treating the patients who have insulin resistance (T2D) (Malik et al 2012). Diabetes has a strong relationship with renal and liver diseases (Mazen and John 2015). Camel milk protected the liver and kidney function from failure; we suppose that camel milk contains insulin nanoparticles that safeguard the role of kidney and liver by restoring the normal glucose levels in the blood. Previous studies (Sharma et al 2012; Saleh et al 2016; Merecz et al 2015), reported that diabetes increases the induction of oxidative stress by elevating the level of MDA and decreasing antioxidant concentrations in the serum, and that coincides with our results. Zeineb et al
2014, investigated the antioxidant activity of camel milk and found that the casein of camel milk inactivates the reactive oxygen species by scavenging the free radicals. Mohamad et al. (2009) and Hanieh et al. (2015) evaluated the effects of camel milk on the TC, HDL and TG levels in T1D and T2D respectively, their findings agreed with our results that, CM normalized the alteration in HDL and TG, while reduced the increase in TC levels. Therefore, camel milk can give promising results when used as dietary supplement for patients of T1D. Insulin plays a main role in the regulation of PK, IRS-2, CPT-1, PK and FASN gene expression (Quinn and Yeagley 2005; O’Brien et al. 1990). PK enzyme catalyzes the rate-controlling step of gluconeogenesis. Insulin is a dominant negative regulator of PK gene transcription (Kaushik and Richard 2007). Our results showed that camel milk did not reduce the expression of PK gene. Probably the increase in both insulin and leptin levels are the cause for this effect. As both hormones can regulate glucose levels, therefore, no need for the gluconeogenesis. Insulin resistance is the leading cause of T2D, which is correlated with reduction in expression of both major insulin receptor substrate (IRS) proteins, (IRS-1) and (IRS-2) (Fisher and Kahn 2003). Our results showed that camel milk provoked the activity of IRS-2 while metformin failed to achieve this goal. Metformin works directly on the reduction of glucose production from non-carbohydrate carbon substrates in the liver through gluconeogenesis metabolic pathway (Michael and Deborah 2001). The reduction of IRS-2 expression causes severe reduction of β-cells (Kubota et al 2000). Camel milk contains zinc, which has a key role in the insulin maturation and secretion in pancreatic β-cells (Gisela 2005), in addition to the presence of camel milk immunoglobulin may offer massive potentials by interaction with the host cell protein that cause an enhancement of regulatory cells and leads to down-regulation of the immune system and salvage the β-cells (Agrawal et al 2007). Various published data reported that insulin stimulates expression of the PK gene which is responsible for the increase in hepatic glycolysis to help in peripheral glucose utilization (Yuuki et al. 2003; Yamada and Tanaka 1999). Administration of camel milk normalized the expression of CPT-1 level in the diabetic rats, reduced the TG levels to normal levels and elevated the levels of high-density lipoprotein cholesterol (HDL). There is an inverse relationship between CPT-1 expression and triglycerides content in the cell. Insulin reduces the expression of CPT-1 and increases glyceroneogenesis and triacylglycerides synthesis. Furthermore administration of camel milk inhibited lipolysis associated with diabetes because the used camel milk is already non-defatted. The FASN gene is the enzyme involved in converting acetyl-CoA into fatty acids (lipogenesis). Insulin plays a key role in the regulation of fatty acid synthase (FAS). It does increase the rate of FASN gene transcription and FAS enzymatic activity. Conversely, in the case of insulin resistance FASN is markedly inactivated (Jose et al. 2010). The results showed that administration of camel milk enhanced the expression of FASN gene. This enhancement is referred to the insulin particles included in the camel milk. We boosted our investigation by the immunohistochemical test. STZ induced diabetes by destroying the pancreatic β-cells (Eleazu et al. 2013; Sharma et al. 2012). Checking the amount of produced insulin is a good indicator of the normal case of Langerhans islets β-cells because the active insulin is secreted from secretory granules in the β-cells (Agrawal et al. 2007).

Our results showed that administration of camel milk showed restoration of insulin secretion in diabetic rats; this means that the Langerhans islets β-cells restored their activity. There are three possibilities to explain this result; (1) regeneration has occurred of distorted β-cells, this explanation agrees with Srinivasan and Ramaraoo (2007) who mentioned that there is a spontaneous regeneration of damaged β-cells in the chemically induced diabetes. (2) The undamaged β-cells secrete insulin with overdose to compensate the shortage caused by damaged cells, (3) camel milk reduced the STZ damage effects occurred in the β-cells; this explanation congruent with Hamers et al. (1993). Furthermore, camel milk has antitoxic effects against drugs (Afifi 2010; Khan and Alzohairy 2011; Al-Fartosi et al 2011), as well as antioxidant effects (Zeineb et al 2014) that reduced the STZ dangerous effects. GLUT-4 is insulin-responsive glucose transporter, which diffuses glucose into adipose and muscular tissues (Stenbit et al. 1997). We indicated that; the reduction of GLUT-4 appeared in the diabetic rats (under immunohistochemical examination) reflects the decrease in insulin secretion. Because the expression of GLUT-4 is stimulated by cascade gene regulation enhanced by secretion of insulin hormone (Gisela 2005). Administration of camel milk restored the expression of GLUT-4 in the pancreas tissue which is detected by the immunohistochemical staining, camel milk already contains insulin as mentioned in different articles (Malik et al. 2012; Al-Hashem 2009; Kamal et al. 2007; Abd El-Aziz et al. 2012) as well as it restored the activity of the β-cells as we mentioned previously.

Conclusion

Camel milk is a natural product that can be considered as a nutritional supplement that helps in treatment of diabetes and it metabolic associated disorders such as insulin resistance. It has dynamic effects on the regulation of CPT-I, FASN, PK and IRS-2 expression, stimulation of insulin production and secretion from the pancreas. CM ameliorated and normalized the changes in glucose, TC, TG, and HDL levels in the blood of diabetic rats.
Ethics approval and consent to participate

All procedures used in treating animals were approved by the Ethical Committee Office of the dean of scientific affairs of Taif University, KSA.

Author Contributions

All authors participated in acquisition, analysis or interpretation of the data, in the drafting of the article or critical revision and in final approval of the version to be published; all authors agreed to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the article are appropriately investigated and resolved.

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