The effects of leptin on F-actin remodelling in type-I diabetes

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The effects of leptin on F-actin remodelling in type-I diabetes

Running Title: F-actin remodeling by diabetes and leptin treatment

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ABSTRACT

Background: The aim of the current study is to investigate the effect of leptin on cytoskeleton structures in both in vivo and in vitro model of diabetes.

Materials and methods: For in vivo studies, leptin in different doses (240, and 480 mg/kg) was injected to the diabetic rats after one-week of Streptozotocin (STZ, 55 mg/kg) treatment. Leptin levels were analyzed in serum, liver, and pancreas samples. Hepatic and pancreatic F- and G- actin expressions were determined by Western blotting. For in vitro studies, hepatic and pancreatic primary cell lines were obtained from the control rats. To these cultures, STZ (15 and 30 mM), leptin (50, 60 and 100 ng/ml), and their combinations were applied for one, three, and four weeks. After the treatment period, F-actin was visualized by the Alexa-fluor fluorescent dye.

Results: STZ decreased the G-actin in both tissues in vivo. However, leptin caused a dose-dependent increase in G-actin levels while F-actin decreased in both tissues. Moreover, leptin caused the perimembranous condensation of actin filaments and amelioration of F-actin structures in vivo. A dose-dependent corruption of F-actin filament structures was observed in leptin-treated primary cells in vitro, while STZ also caused corruption of these filaments. Co-exposure of STZ and leptin caused the amelioration of F-actin filaments, while the perimembranous condensation was also observed as was in vivo study.
Conclusions: Leptin therapy could be a candidate for diabetes, but it should not be ruled out as being important the severity of diabetes and leptin doses.

Keywords: cytoskeleton, diabetes, F-actin, G-actin, hepatic cells, pancreatic β-cells, leptin therapy

INTRODUCTION

Diabetes mellitus (DM) is one of common disease worldwide. In 2014, it was reported that 7.5% of women and 5% of men face to DM. The prevalence is expected to reach 439 million by 2030 meaning of total 7.7% adult population between 20-79 years [1]. There are two types of DM. One of this called Type I associated with the increased blood glucose due to insufficient production of insulin in the body. Insulin insensitivity of the cells causes to type II, and the body produces sufficient insulin in this cases. Therefore, it is essential to find effective strategies to manage or control the prevalence of DM. Herein, leptin takes great attention in the control of hyperglycemia and/or hyperglycemia-induced dysfunctions.

Leptin, a protein hormone, affects the body weight, energy metabolism, reproductive system, and the neuroendocrine system [2]. Furthermore, the hormone plays a significant role in insulin-glucose homeostasis in the pancreas, liver, muscle, and hypothalamus as well. Insulin stimulates leptin secretion [3]. Leptin has affected the centrally in the hypothalamus and peripherally in the pancreas, muscle, and liver. Leptin also improve glucose-insulin metabolism via attenuating glycemia, insulinemia, and insulin resistance.

To develop leptin-based therapies against diabetes, and other insulin resistance syndromes, leptin’s effect on the glucose-insulin homeostasis should be understood well [3]. There is a complicated relationship between insulin resistance and leptin. For example, leptin may increase insulin sensitivity and improve the glucose metabolism[4]. Also, a previous study indicated that insulin resistance was ameliorated by administration of leptin in mice [5]. Hence, it was proposed that the leptin plays a possible role in insulin resistance and it can improve insulin production [5]. Insulin resistance can be described as insufficiency in the insulin pathway at target tissues which generally response, like adipose, liver, skeletal, and heart muscle [6].
The cytoskeleton is essential to keep cell shape and signaling pathways for the beta (β) cells. The perturbation of actin structures might give rise to enhance β cell dysfunction, including inflammation and fibrosis [7]. Actin plays a vital function in morphology, migration, adhesion, and permeability [8]. Actin is abundant and highly conserved proteins among species. It is a member of structural superfamily sugar kinases hexokinases and Hsp70 proteins [9]. Actin is one of the cytoskeleton element and has many essential roles in cells such as cell shape, polarity, endocytosis, contractility, motility, intracellular trafficking, etc. Actin has two form of monomeric G-actin and filamentous polymeric F-actin [10]. G-actin can be converted to F-actin by the presence of ATP, ions, actin-binding proteins. F-actin and myosin interaction cause muscle contraction. The primary control factor of transformation is nucleotide hydrolysis [9]. Conversion of G actin to the filamentous form participates in pathological processes, such as cancer metastasis as well [11]

There is little knowledge available on the relationship between insulin and leptin. The current study was aimed to evaluate how F actin structure can be affected by leptin therapy in STZ-induced diabetic rats in both in vivo and in vitro conditions. To our knowledge, it is the first study to investigate the effects of leptin on cytoskeleton structures of the STZ-treated cells and animals.

**MATERIAL AND METHODS**

**Experimental Animals**

Male Wistar rats weighing between 140–250 g were purchased from the Faculty of Veterinary Medicine, Department of Reproduction and Artificial Insemination, Istanbul University. The animals were individually housed in stainless cages at 20±2°C, 60-70% humidity with 12:12 h photoperiod. Water and food were given ad libitum. All processes were approved by the Istanbul University Animal Welfare Committee.

**Induction of Experimental Diabetes**

Diabetes was induced by Streptozotocin (STZ). Streptozotocin was freshly prepared in a citrate buffer (50 mM, pH 4.5). Control group received only citrate buffer as a vehicle. 55 mg/kg of STZ was injected the intraperitoneally to rats [12]. One week
after STZ injection, animal having blood glucose levels ≥ 250 mg/dL was accepted as diabetic [13]. The blood glucose levels were measured by using a glucometer (OneTouch Ultra; LifeScan, Milpitas, CA, USA) from orbital vein blood samples.

**In vivo experimental design**

Four groups were randomly carried out as (I) control (C): non-diabetic rats treated with the vehicle, (II) diabetes (D): diabetic rats treated with the vehicle, (III) diabetes+leptin 240 (DL240): diabetic rats treated with 240 ng/kg dose of leptin, (IV) diabetes+leptin 480 (DL480): diabetic rats treated with 480 ng/kg dose of leptin (modified from [14]). Both doses are physiologic doses of leptin [15] Experimental animals were treated with the leptin every other day for one week. C group received intraperitoneally physiological saline as the same volume of DL group. Four experimental animals were used for each group. After treatments periods, liver and pancreas were removed from experimental animals for F and G actin analysis by Western blotting. Body weights were measured at two time points: at the beginning of the first injection and 24 h after the last injection. Then, body weight gain was calculated by subtracting from the last day to the first day of the experiment.

**In vitro experimental design**

For in vitro studies, primary hepatic (modified from [16]) and pancreatic cells (modified from [17]) were isolated from the corresponding tissues of the C group. Hepatic primary cells were recognized as parenchyma and Kupffer. The liver was removed aseptically and washed with Hanks' balanced salt solution (HBSS). Then, the tissues were minced into one-mm³ pieces in HBSS. The cells were suspended in William’s Medium E (WME) and centrifuged for 5 min at 3000 x g and the supernatant discarded. Cells were dissolved in WME and transferred into an incubator with 5% CO₂ atmosphere at 37 °C. For pancreatic acinar primary cell culture, each pancreas was minced into one mm³ piece and incubated with two mL of HBSS with 2 M EDTA at 37 °C for 15 min. The chelated mixture was centrifuged for 5 min at 3000 x g, and the supernatant was discarded. The pellet was rinsed with ten mL Ham's F-12 medium, centrifuged for 5 min at 3000 x g, and the supernatant discarded. The tissue pellet digested with ten mL of 1 mg/mL collagenase type II, 1 mg/mL hyaluronidase and 20 %
heat-inactivated calf serum in Ham's F-12 medium at 37 °C for 30 min in a shaking water bath. Following centrifugation at 3000 x g for 5 min, the supernatant discarded, pellet rinsed with ten mL of HBSS, and this procedure was replicated. The cells were rinsed with 5% heat-inactivated calf serum in Ham's F-12 medium and centrifuged at 3000 x g for 10 min. The pellet rinsed with ten mL of 5% heat-inactivated calf serum in Ham's F-12. Cells allocated to a well containing two mL of Waymouth's serum-free medium and incubated at 37 °C with 5 % humidified CO₂ for 48 h.

The hepatic and pancreatic cells from non-diabetic control rats were divided into four groups as control (c), STZ (S), leptin (L), and STZ-leptin (SL) for one and four weeks. Like hepatic cells, the pancreatic cells were also divided into four groups. To observe the effect of severity of hyperglycemia, the cell lines were treated with two different doses of STZ as 15 or 30 mM. To determine the effect of different doses of leptin on cell lines, 50, 60 and 100 ng/mg of leptin were injected into the cells. The hepatic cells were treated with the leptin for one and four weeks. The pancreatic cells were treated with the leptin for three weeks. The data from dl groups in the pancreatic cells were not obtained because of lower cell numbers although the medium was replaced two times a week.

**F-actin organization**

Immunofluorescence microscopy was used to determine F-actin organization in both *in vitro* (the primary cell cultures) and *in vivo* (cryo-sections of liver and pancreas) [18]. Briefly, primary culture cells were fixed with 4% paraformaldehyde in PBS for 10 min at room temperature and then permeabilized and blocked in PBS containing 0.1% Triton X-100 and 5% fetal calf serum for 30 min. Actin filaments were visualized using rhodamine-phalloidin or Alexa 497, and nuclei were labeled with 4′,6-diamidino–2–phenylindole dichloride (DAPI) (Invitro Molecular Probes, OR, USA). All images were obtained using an Olympus BX51 Microscope equipped with a DP72 camera controlled by Olympus DP2-TWAIN software.

**Western Blotting**

*In vivo*, F and G actin proteins were isolated from the pancreas and liver tissues. Tissues were homogenized by a MagNA lyser (Roche Diagnostics). Aliquots of liver
and pancreas homogenates were centrifuged at 3000x g for 15 and 4500x g for 20 min, respectively. F and G actin were separated by a kit (Cytoskeleton, Cat #BK037) according to the manufacturer’s instruction [19]. Then, 50 µg F-actin or G-actin were loaded to SDS–PAGE and transferred to a PVDF membrane (Millipore Corporation, Bedford, MA, USA) at 100 V for 45 min. The membranes were blocked by using 5% bovine serum albumin. Membranes were incubated with anti-actin rabbit polyclonal antibody (1/500) at +4 ºC for overnight (Abcam, USA), and then incubated with a horseradish peroxidase-conjugated goat anti-rabbit IgG antibody for one h at room temperature. 5-Bromo-chloro-3-indol phosphate (BCIP) was used for visualization for protein band. After membranes were photographed with Canon camera, ImageJ was used to analyze the band.

**Leptin Analyzing**

Blood and homogenates of liver and pancreas were centrifuged at 10000 g for 15 min at +4ºC. Leptin levels were measured by ELISA using a commercial assay kit according to the manufacturer’s directions (Millipore, USA) [20].

**Statistical Analysis**

The analyses of statistical were performed by SPSS version 23.0. Data are expressed as mean ±SEM. One-way analysis of variance (One-Way ANOVA) with post doc Tukey for equal variance assumed and post hoc Dunnett’T3 for equal variance not assumed were used to compare mean between the groups. \( P < 0.05 \) was accepted as the statistical significance.

**RESULTS**

**Effect of Leptin on Body Weight and Blood Glucose level**

Diabetes decreased significantly body weight gain in D groups when compared to the C group \( (p < 0.05) \). The treatment of leptin failed to control body weight gain in diabetic rats. Interestingly, 240 ng/kg of leptin non-significantly worsen the body weight gain in DL240 group compared to the D group. But, there were no significant differences in body weight gain among D, DL240, and DL480 groups (Figure 1).
Blood glucose level is a primary parameter to determine diabetes. The delta was calculated by subtraction of glucose levels before and after experiments. As expected, all diabetes groups had high blood glucose level and the delta compared to C group after STZ injection ($p < 0.05$). Leptin treatment did not improve the high blood glucose when compared to D group (Figure 2).

The leptin levels were measured in serum, liver and pancreas samples. Diabetes did not influence the leptin levels in serum, pancreas, and liver in contrast to C group. Also, leptin treatment in all dose significantly increased the serum leptin levels compared to D group as expected ($p < 0.01$). The doses of leptin higher than 240 ng/kg significantly increased the serum leptin levels in contrast to D and DL240 groups ($p < 0.01$). However, the treatment of diabetic rats with leptin did not affect the level of leptin in liver and pancreas when compared to D group (Figure 3).

**The Effect of Leptin on Actin Structure**

The actin structure is so crucial for glucose metabolism. Therefore, it was evaluated the F-actin distribution of liver and pancreas tissues in vivo and pancreas primary acinar, Kupffer and parenchymal also called as hepatocyte cells in vitro, as well.

For in vivo studies, diabetes induction gave rise to disrupt F-actin structure in liver tissues. The treatment of diabetic rats with 240 ng/kg dose of leptin improved F-actin in DL240 group when compared to D group; however, as the dose of leptin (480 ng/kg, DL480) increases, it worsens the effect of diabetes on F-actin (Figure 4a-d). In pancreatic tissue, while diabetes also has a destructive impact on the F-actin structure in D group compared to C group, the leptin treatment in all doses had the protective effect on F-actin structure compared to D group (Figure 4e-h). To evaluate the effect of leptin treatment on different cell types, parenchymal and Kupffer cells were also isolated from rats in C, D and DL240 groups. Based on cell types, the effects of leptin on F-actin showed differences. In the parenchymal cells, F-actin disappeared in the cytosol (Figure 5b, c) whereas slight destruction was observed in Kupffer cells (Figure 5e, f).

Isolated primary cell cultures were preferred to reduce the number of animals used in the experiment. In primary liver cells, only 100 ng/ml dose of leptin (higher dose) was destructive on F-actin structure (Figure 6 g, h, i); therefore, the treatment of
the cell with the leptin was not continued after three weeks. When compared to 100 ng/ml dose of leptin, 50 ng/ml dose of leptin was not so destructive to the F-actin structure in the isolated liver cells. In pancreas cells, the detrimental effect of leptin on F-actin was only observed at the dose of 60 ng/ml (Figure 7c, d). In the present study, the effect of leptin on F-actin structure in STZ-induced pancreatic cells (SL group) were not evaluated due to high toxicity of STZ on pancreatic cells (Figure 7e, f) because the cells were exposed to STZ at the beginning of the study and F-actin destruction was measured after three and/or four weeks. That indicates the highly destructive effects of STZ on the pancreas.

Also, the effects of leptin on F-actin structures in the primary hepatic cells were studied based on the severity of hyperglycemia or diabetic condition. At the mild diabetic condition (15 mM STZ), filopodia formation by hyperglycemia (Figure 8 B1 and B2) was reversed by leptin treatment without any protection on the F-actin structure after four weeks in both at parenchymal and Kupffer (Figure 8 D1 and D2). Leptin failed to improve F destruction and filopodia formation (Figure 8 E1 and E2); both induced by severe diabetic condition (30 mM STZ; Figure 8 C1 and C2).

**The Effect of Leptin Treatment on F and G Actin Protein Expression**

Leptin treatment showed a protective effect on disproportional F and G actin protein expressions compared to diabetes groups in both liver and pancreas tissues *in vivo* (Figure 9 A, B, C, and D).

**DISCUSSION**

Cytoskeleton plays a vital role in the cell morphology and the elasticity [21]. Insulin can tonically regulate actin filament network and microtubule as well [22, 23]. Actin network mediates glucose transport through insulin action [23]. Actin phosphorylation alters cytoskeleton re-orientation via polymerization and depolymerization [21]. Insulin stimulates leptin secretion [3]. Leptin is considered as an anti-diabetic hormone. It plays a significant role in glucose homeostasis in the pancreas, liver, muscle, and hypothalamus. Also, it blocks glucose production in the liver and enhances glucose consumption in the tissue [24]. It was reported that the effects of
leptin were mediated via the JAK/STAT signaling pathways. β-cells are hyperpolarized via ATP sensitive potassium channel (K_{ATP}) opening by leptin; actin cytoskeleton reorganizes through the PI3K pathway [25]. STZ causes hyperphagia, hyperleptinemia, hypoinsulinemia, and hyperglycemia [26]. Insulin can secrete leptin from adipose tissue, controversy; leptin suppresses insulin secretion from pancreatic β-cells by activation of K_{ATP}. Leptin is reported to increase K_{ATP} trafficking to enhance the channel density on the membrane by strengthening F-actin depolymerization. The trafficking of K_{ATP} by leptin is mediated through AMPK and PKA (protein kinases-A) [27].

In the present study, diabetes did not affect the serum, liver and pancreatic leptin levels. However, leptin treatment elevated only the serum leptin levels in diabetic rats without altering the pancreatic and liver levels. Our results are an inconsistency with the previous study, which indicates that increased plasma insulin levels due to increased insulin resistance could elevate the pancreatic leptin levels [28]. The result of the present study demonstrates that leptin did not improve blood glucose level in diabetes. Inconsistently, it was reported that a high dose of leptin ameliorated the both of hyperlipidemia and hyperglycemia [29, 30]. Co-treatment of leptin and insulin could be more useful to prevent hyperglycemia than insulin only therapy, which reduces blood glucose to undesirable levels. One advantage of leptin is without to gain weight in the treatment of type 1 diabetes while making a balance of blood glucose. An extensive research report that type 1 diabetes reduces body weight [29, 31] as reported in the present study. It is probably that type I diabetes causes dehydration, declining glucose utilization and protein anabolism [9]. Leptin provides a link between body weight and food intake [32, 33]. A correlation between serum leptin levels, plasma insulin, and body mass index has been reported [33]. Leptin in high concentrations causes insulin resistance [33]. In the present study, diabetic rats lose the body weight as stated in the previous study [34]. While leptin treatment seems to be protective, it was considered that weight gain in leptin-treated groups looks like developing type 2 diabetes as was explained in below. Leptin antagonizes the effect of insulin on glucokinase and phosphoenolpyruvate carboxylase enzymes to decrease blood glucose levels in the liver. A decrease in leptin efficiency is related to insulin resistance in type 2 diabetes [35]. Insulin secretion occurs in two phases. One phase called fast secretion requires glucose
entry to the β-cells, and this leads to the closure of ATP sensitive potassium channel (K\textsubscript{ATP}) resulting in depolarization [36]. Finally, insulin secretion occurs. The second phase is essential to maintain the insulin level in plasma [36]. This phase is associated with the reserved pool, which is far away from the plasma membrane [37]. Reserved insulin granules must be transported to the plasma membrane [36]. F-actin is vital to maintaining basal plasma insulin levels [38]. Therefore, actin cytoskeleton plays a significant role in both phases of insulin secretion [37]. In the current study, diabetes caused a decrease of G-actin, while any effect was observed on F-actin in vivo. This effect can be explained that glucose enhances F/G-actin ratio which means that hyperglycemia can inhibit G-actin disassociation by partial stimulation of cofillin phosphorylation. Cofilin belongs to one of actin-depolymerizing factor family and can be inactivated with phosphorylation by LIM kinase, causing F-actin stabilization [39]. Rho-A could participate in this pathway [40]. Diabetes is well known as an autoimmune disease. Macrophages inhibit Rho pathways caused to disrupt of the cytoskeleton from subcortical actin ring to near the cell periphery [41]. G-actin is needed for F-actin resembling. Therefore; F-actin maintains its proper function. It was proposed that attenuation of G-actin participate in the development of diabetes.

Furthermore, F-actin reorientation was observed the at the plasma membrane surrounding in the current study. F-actin can be depolymerized and scavenged by the action of gelsolin, which is one of the free calcium regulators, resulting in blood glucose attenuation via the blockage of insulin vesicle transport to the membrane [9]. It was suggested that a high plasma glucose level could induce actin cytoskeleton disruption [42]. This is asserted to relate apoptotic cell death [43]. That is why F-actin could release from cell to plasma when cell injury is occurred [9]. Leptin is also reported to activate LIMK for phosphorylation of cofillin, resulting in accumulation F-actin and depletion of G-actin. The other effect of leptin on actin is up-regulation of α-skeletal actin and myosin light chain-2 (MLC-2) [44].

Insulin and leptin have anti-apoptotic and proliferative effects [45]. Furthermore, leptin was shown to have a positive impact on glucose metabolism, and it was an excellent adjuvant candidate for insulin therapy [46]. Leptin enhanced the insulin sensitivity in rats with type 2 diabetes and patients with type 2 as well. A single administration of leptin ameliorated the mortality related to high blood glucose and
insulin independent insulin deficiency [47]. Leptin is shown its anti-diabetic effect via increasing insulin receptor sensitivity. However, the involvement of F-actin in the mechanism of leptin on type-1 diabetes is not clearly evaluated. Diabetes and leptin therapy showed their destructive effect on F-actin according to cell type in the current study in vitro. F-actin disappeared from cytosol center and accumulated within the periphery of the cell membrane in primary parenchymal cells.

On the contrary, F-actin clustered within the cytosol center as multicentered radiant groups in Kupper primary cells. In primary pancreas cells, the destruction process in actin filament is similar to Kuppfer primary cell. It was shown that leptin binding to its receptor (Ob-Rb) phosphorylates JAK2 and transforms PIP$_2$ into PIP$_3$. This destroys the F-actin into the G-actin [48]. One study is reported that RhoA and ROCK are mediated leptins effect on F-actin distribution. Leptin is suggested to enhance the expression and activities of GLUT2 and GLUT5 transporters [49]. Leptin has been reported to improve insulin sensitivity in both healthy and diabetic animals. However, the hormone is also said to destroy insulin effect on hepatocytes. Leptin infusion is an essential factor for disclosing its effect on tissue. Leptin is suggested to decrease blood glucose and insulin level without altering body weight of animals by subcutan infusion. Leptin intravenously injection boosts inulin sensitivity in healthy animals, increase glucose consumption in diabetic animals. STZ destroy pancreatic β-cell and decrease adipose and plasma leptin levels. Therefore, leptin therapy can mimic insulin action by insulin-independent and an insulin-sensitizing mechanism in STZ treated animals. Leptin has been shown to activate K$_{ATP}$, resulting in decreasing insulin secretion [50].

Cell skeleton is a dynamic and sensitive system. It is pointed out as an important morphologic parameter in the study of changes that took place in the cell after getting into reaction with substance [51]. Some vesication is formed within the whole cell during the process of cell skeleton’s restructuring. This incidence of vesication that took place outside the cell goes along both with the distribution of microtubules [52] and actin filaments’ restructuring. Actin polymerization occurs at G1 and G2/M phases of the cell cycle [53]. In actin polymerization, F-actin/G-actin rates show cellular separation (differentiation) [53].
CONCLUSION

The in vivo protective effect of leptin treatment on the F-actin structure in both liver and pancreas tissues has been presented in the current study. However, leptin could not significantly improve F-actin orientation in vitro in different cell types. A slight improvement of F-actin orientation was observed in Kupffer cells compared to parenchymas. Leptin at the higher doses can contribute to the destructive effects of diabetes. Filopodia formation can be reversed by leptin treatment without changing the F-actin destruction according to the severity of diabetic conditions. In conclusion, leptin treatment could be useful when the severity of diabetes and leptin doses are taken into consideration.

ACKNOWLEDGE

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REFERENCES


FIGURE LEGENDS

**Figure 1:** Body weight gain. Abbreviations: C, Control; D, Diabetes; DL240, Diabetic rats treated with 240 ng/kg dose of leptin; DL480, Diabetic rats treated with 480 ng/kg dose of leptin. *P < 0.01; different from C group

**Figure 2:** Blood Glucose Level. Abbreviations: C, Control; D, Diabetes; DL240, Diabetic rats treated with 240 ng/kg dose of leptin; DL480, Diabetic rats treated with 480 ng/kg dose of leptin. *P < 0.01; different from C group.
Figure 3: Leptin levels in serum, pancreas, and liver. Abbreviations: C, Control; D, Diabetes; DL240, Diabetic rats treated with 240 ng/kg dose of leptin; DL480, Diabetic rats treated with 480 ng/kg dose of leptin. *P < 0.01; different from D group.

Figure 4: The leptin effect on F-actin distribution from liver and pancreas tissues. Abbreviations: Liver tissues were as from a to d, although pancreas tissues were from e to h. a and e) Control; b and f) Diabetes; c and g) Diabetic rats treated with 240 ng/kg dose of leptin; d and h) Diabetic rats treated with 480 ng/kg dose of leptin.

Figure 5: F-actin distribution of primary liver cells at parenchymal (upper line) and Kupffer cells (below line). Abbreviations: Red color represents F-actin, blue represents the nucleus by DAPI at x100 magnification. Parenchymal cells: a and d) Control; b and e) Diabetes; c and f) Diabetic rats treated with 240 ng/kg dose of leptin.

Figure 6: The leptin effect on F-actin distribution after three weeks treated on primary liver cells. Abbreviations: a, b, c) Control; d, e, f) cells treated with 50 ng/ml dose of leptin; g, h, i) cells treated with 100 ng/ml dose of leptin. c, f, i is at 20 magnification, and the other is at 100 magnification.

Figure 7: The leptin effect on F-actin distribution at primary pancreas cells at 100 magnification. Abbreviations: a and b) Control; c and d) cells treated with 60 ng/ml dose of leptin; e and f) cells treated with 30 doses of mM STZ (magnification x 100).

Figure 8: The leptin effect on F-actin distribution at parenchymal (A1, B1, C1, D1, E1) and Kupffer (A2, B2, C2, D2, E2) cells after four weeks treatment. Abbreviations: A1 and A2) Control; B1 and B2) cells treated with 15 mM of STZ, C1, and C2) cells treated with 30 mM of STZ, D1 and D2) 15 mM exposed cells treated with 60 ng leptin E1 and E2) 30 mM exposed cells treated with 60 ng leptin.

Figure 9: The alternation of F and G actin protein expression in pancreas and liver tissue. Abbreviations: 1, control; 2, diabetes; 3 Diabetic rats treated with 240 ng/kg dose
of leptin; 4, Diabetic rats treated with 480 ng/kg dose of leptin. A: F-actin of the liver; B: G-actin of the liver; C: F-actin of the pancreas; D: G-actin of the pancreas

**Figure 1**

**Figure 2**
Figure 3

Leptin concentration (ng/ml)

- C
- D
- DL240
- DL480

Legend:
- ■ Serum
- □ Pancreas
- □ Liver

Figure 4
Figure 7
Figure 8

Figure 9