Original Article

Effect of Iranian Honey bee (Apis mellifera) Venom on Blood Glucose and Insulin in Diabetic Rats

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Abstract

Background: Diabetes is an important disease. This disease is a metabolic disorder characterized by hyperglycemia resulting from perturbation in insulin secretion, insulin action or both. Honey bee venom contains a wide range of polypeptide agents. The principle components of bee venom are mellitin and phospholipase A2. These components increase insulin secretion from the β-cells of pancreas. This study was conducted to show the hypoglycemic effect of honey bee venom on alloxan induced diabetic male rats.

Methods: Eighteen adult male rats weighting 200±20 g were placed into 3 randomly groups: control, alloxan monohydrate-induced diabetic rat and treated group that received honey bee venom daily before their nutrition for four months. Forty eight hours after the last injection, blood was collected from their heart, serum was dissented and blood glucose, insulin, triglyceride and total cholesterol were determined.

Results: Glucose serum, triglyceride and total cholesterol level in treated group in comparison with diabetic group was significantly decreased (P< 0.01). On the other hand, using bee venom causes increase in insulin serum in comparison with diabetic group (P< 0.05).

Conclusion: Honeybee venom (apitoxin) can be used as therapeutic option to lower blood glucose and lipids in diabetic rats.

Keywords: Alloxan, Glucose, Honeybee Venom, Rat

Introduction

Diabetes type 1 is mainly results from autoimmune beta cell destruction, while viral infections and chemical agents seem to be the triggers of the disease. The well-documented effect of insulin is to mediate carbohydrates, proteins and lipids storage. Therefore diabetes is considered as a defect of lipids, proteins and carbohydrates metabolism in which all the body systems and organs are affected (Williams and Pickup 2000). Natural toxins have been traditionally used to heal diseases and honeybee venom (apitoxin) is of a great importance in this regard.

The venom is composed of varieties of peptides (mellitin, apamin, secapin, tertiapin, adolapin, and MCD peptid), enzymes (phospholipase A2, hyaluronidase, acid phosphomonoesterase, lysophospholipase), active amines (histamine, dopamine, norepinephrine, serotonin) and many other substances (Son et al. 2007). Bee venom and Bee sting had signifi-
Materials and Methods

Honeybee venom preparation procedure

Bee venom samples were collected from beehives using an electric shocker, on February 2010 in Khuzestan, Iran. The electric shoker is composed of two components: one component as shoker and the other to collect the venom and concomitant material. The collecting unit is wooden and composed of a network of wires with small gaps between them. A glass plane is inserted under the network. The shoker was supplied by a transformer to produce a light electric shock. The shoker is designed to produce a light electric shock once every few seconds. The collector panel was first located on bottom of the beehives and then on the top, to collect the desired amount of the bee venom. When the shoker was turned on, the honeybees stroked the wires and received a light electric shock and were stimulated them to sting and discharged their bee venom. Alarm pheromone produced by the exited bees to crowd and discharge toxin. The shoker was turned off after 25 minutes and the collecting panel was removed from the beehive, and the dried bee venom material scratched and transferred to a proper container. To evaluate the toxin production efficiency after collecting the samples, the crystallized bee venom material weighed using a sensitive scale.

Animals

Healthy adult male Lewiss rats weighting 200±20 g were maintained in Darou Paksh Pharmaceutical Mfg Co, Tehran (Iran). All animals left to acclimatize for one week before the experiment. The animal room was maintained under a constant 12-h light: 12-h dark cycle and temperature of 23±3 °C and relative humidity of 70±10% throughout the experimental period. The rats were given free access to standard pellets and water. For bees to be adapted to the new environment
condition, all the experiments were carried out 10 days after their first residence.

The research was approved by Ethical Committee of the university.

**Experimental design**

Eighteen rats were placed randomly into three groups (n=6):

- Control group: normal saline 0/9% injected intraperitoneally.
- Diabetic group: this group became diabetic by injection Alloxan monohydrate at 150 mg/kg intraperitoneally.
- Bee venom-treated group: at first received Alloxan monohydrate to induce diabetes and Iranian bee venom (apitoxin) at 0/5 mg/kg (the best dose chosen after pretest) after diabetes confirmation intraperitoneally at fasting condition every day for four consecutive weeks (Kim et al. 1999, Dong et al. 2007).

Blood sugar test was performed in all groups. Blood glucose level was measured using Acua Check (Germany). To measure the blood glucose level, a small incision was made on the animals’ tail using a lancet and a drop of fresh blood was extracted and used for glucometry. These samples were collected in fasting condition and expressed in mg/dl.

Alloxan monohydrate (Sigma-Aldrich Germany) was used to induce diabetes in rats. The drug was administrated intraperitoneally at the rate of 150 mg/kg (Viana et al. 2004, Antia et al. 2005). The drug administration leads to pancreas β-cells apoptosis and necrosis. The method is preferred to induce diabetes in many other animals (Soto et al. 2001). Following the administration a condition of hyperglycemia like diabetes type 1 appeared in rats (Byung-Hyun and Jin-Woo 2001). Seventy two hours after the administration, blood glucose level was measured using Acua Check to determine diabetic condition. In this study, blood glucose level elevation over 280 mg/dl is supposed to reveal diabetic condition (Zhang and Tan 2003). Usual signs of diabetes including polydipsia, polyuria and weight loss were observed 6–7 days following the administration (Nuraliev et al. 1992, El-demerdash et al. 2005).

To measure glucose level, fasting blood samples were collected 2 weeks following bee venom administration using "Stone" method. In this method, venous blood was collected from infraorbital sinus using a hematocrit tube. The animal is kept between forefinger and thumb while the tube is inserted to the orbital foramen with a rotational movement. The capillaries are usually very sensitive and fragile here and burst following a light pressure. When a few almost large drops of blood were collected, the hematocrit tube is removed. The method is suitable for repetitive blood sampling especially for primary analysis following 2 weeks of administration. The blood glucose level was determined using enzymatic kits.

**Biochemical analysis**

The animals were anesthetized with ether after 16 h of fasting to collect blood for analysis. Forty eight hours after the last injection, blood samples were collected from the heart and centrifuged (3000 rpm for 15 min at 4 °C) for separating the serum. The serum was then frozen at -70 °C for the biochemical analysis. Then the amount of blood glucose, insulin, cholesterol and triglyceride in blood serum were determined. Serum glucose level was measured by kinetic (enzymatic) and colorimetric methods using Glucose Estimation Kit (Pars Azmoon, Iran). These serum insulin levels were assayed with an ELISA, IRMA (Biosource Europe SA), serum glucose, triglyceride and total cholesterol levels were determined using commercial kits and enzymatic assays.

**Statistical analysis**

The values were expressed as mean ±S.E.M Statistical analyses were performed by one
way analysis of variance (ANOVA) followed by Tukey multiple comparison test. P< 0.05 were considered as significant.

Results

Bee venom preparation results: the bee venom samples were collected and panel scratched using a sterile blade and kept in clean dark vials. Average amount of venom collected from three beehives (each containing estimated 10000 bees) was 147.7 mg. The average for each beehive was calculated 49.56 mg. Alloxan administration result showed a significant increase in blood glucose level in (diabetic) group compared with control group after 2 and 4 weeks (P< 0.05). In addition a significant increase was observed in blood glucose level in diabetic group after 4 weeks compared with 2-week period (P< 0.05). There was a significant decline in blood glucose level in be venom-treated group compared with diabetic group (P< 0.05). Also there was a significant decline in blood glucose level in diabetic treated group after 4 weeks compared with 2-week period (P< 0.05) (Fig. 1).

The effect of bee venom administration on serum insulin level is illustrated in Figure 2. Our results showed that serum insulin level significantly decreased in diabetic group compared with controls (P< 0.05). Also there was a significant increase in bee venom-treated group compared with controls (P< 0.05) (Fig. 2).

The effect of bee venom administration on serum triglyceride (TG) content is illustrated in Figure 3. There was a significant increase in serum TG content in diabetic group compared with controls (P< 0.05). A significant decline was observed in bee venom-treated group compared with diabetic animals (P< 0.05). But no significant difference was observed in serum TG content in bee venom-treated group compared with control (Fig. 3).

A significant increase was observed in serum total cholesterol in diabetic group compared with controls (P< 0.05). There was a significant decline in bee venom-treated group compared with diabetic animals (P< 0.05). But no significant difference was observed in cholesterol content in bee venom-treated group compared with control group (P= 0.552) (Fig. 4).

![Fig. 1. Changes of blood glucose level after injection of bee venom in diabetic rats](image-url)
Fig. 2. Serum insulin levels after injection of bee venom in diabetic rats. Difference between control and treated group are significant with *P<0.05.

Fig. 3. Triglyceride levels after injection of bee venom in diabetic rats. Difference between control and treated group are significant with *P<0.05.

Fig. 4. Cholesterol levels after injection of bee venom in diabetic rats. Difference between control and treated group are significant with *P<0.05.

**Discussion**

In our study, blood glucose level increased following alloxan monohydrate administration which led to pancreas B-cells destruction (Byung-Hyun and Jin-Woo 2001). Blood glucose level decreased following bee venom treatment. This may be contributed to substances like mellitin and phospholipase A2 contained in the venom. They may play a role in diminishing inflammation of Islets of Langerhans and thus elevating blood insulin level. With regard to the fact that insulin regulates blood glucose level, bee venom could decrease glucose content via increasing insulin secretion (Morgan and Montague 1984, Fujimoto and Metz 1987, Kim et al. 1999, Simonson et al. 2000). Following alloxan administration in diabetic
rats, blood glucose level and triglyceride (TG) content were elevated, which indicate insulin role in regulating lipid metabolism (Zhang and Tan 2003). Insulin activates the enzyme lipoprotein lipase and hydrolysis triglycerides (Frayn 1993). According to the obtained results, bee venom decreased blood TG content. One could explain the observed decline as follows: bee venom improves glycemic control and decreases blood glucose level. Also glucose consumption is increased instead of lipids. Acetyl coA derived from pyrolic acid enters Krebs cycle which finally leads to glucose metabolism, however Acetyl coA can enter TG synthesis pathway in usual condition (Zhang and Tan 2003). Blood cholesterol level increased following Alloxan administration. (Yadav et al. 2004). A decline was observed in cholesterol content of bee venom-treated group (Kim et al. 1999). Probably cholesterol lowering effect is largely due to inhibition of its absorption in small intestine and promoting its hepatic release. The liver plays a critical role in discharging cholesterol via bile secretion (Reinner et al. 1989).

Alloxan administration led to destruction of Islets of Langerhans and diminished insulin secretion in diabetic rats. Treating the rats with honeybee venom (apitoxin) increased insulin secretion up to control levels. According to the published reports, mellitin polypeptide and phospholipase A₂, which are two main component of the venom, promote insulin secretion. According to the literature, the observed effect is mediated by extracellular calcium and calcium channels. When these channels are opened large amounts of calcium enters the β-cells and excite them to secrete insulin (Morgan and Montague 1984, Fujimoto and Metz 1987, Kim et al. 1999, Simonson et al. 2000). In a study on the effects of bee venom on Islets of Langerhans inflammation and onset of insulin dependent diabetics, Kim et al. (1999) showed that intensity of inflammation and onset of diabetes declined following bee venom treatment. They also found that insulin, TG and cholesterol levels decreased in diabetic rats compared with non-diabetic animals (Kim et al. 1999). According to the experiments of Morgan et al. (1984), mellitin polypeptide promotes insulin secretion from Islets of Langerhans in vitro. The obtained results suggest that mellitin as a valuable candidate for further studies on β-cells plasma membrane role in regulating insulin secretion. The findings also indicate that mellitin can depolarize plasma membranes of β-cells and acts as a calcium transporter in the cell, which in turn promotes insulin granules secretion. The effect of mellitin on insulin secretion depends on extracellular calcium (Morgan and Montague 1984). Simonson et al. (2000) found that mellitin may promote insulin secretion via activating phospholipase A₂ in Islets of Langerhans. Their results indicate that phospholipase A₂ activation plays a role in compensating insulin resistance response in Islets of Langerhans (Simonson et al. 2000). Treatment with exogenous phospholipase A₂ or mellitin promotes arachidonic acid and lysophospholipids production and insulin secretion. Produced arachidonic acid and lysophospholipids may corporate in two-step insulin production mechanism. Arachidonic acid produced by phospholipase A₂ induction may act as a calcium transporter in to β-cells and promote insulin secretion (Fujimoto and Metz 1987).

Conclusion

In this study, our results indicate that honeybee venom (apitoxin) can be used as therapeutic option to lower blood glucose and lipids in diabetic rats, however further biochemical and pharmacological studies are necessary to provide more detailed understanding of the issue.

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References


