

Preliminary results of effect of barley (*Hordeum vulgare* L.) extract on liver, pancreas, kidneys and cardiac tissues in streptozotocin induced diabetic rats

Mohsen Naseri (1,2,3), Zahra Khalaj Sereshki (4), Behnaz Ghavami (5), Bagher Minaii Zangii (6), Mohammad Kamalinejad (7), Parvaneh Mohseni Moghaddam (8), Majid Asghari (9), Seyed Abbas Hasheminejad (1,2), Fatemeh Emadi (1,2), Farzaneh Ghaffari (10)

(1) Traditional Medicine Clinical Trial Research Center, Shahed University, Tehran, Iran; (2) Department of Traditional Persian Medicine, School of Medicine, Shahed University, Tehran, Iran; (3) Hikmat, Islamic and Traditional Medicine Department, The Academy of Medical Sciences, Tehran, Iran; (4) Department of Gynecology, School of Medicine, Yas Hospital, Tehran University of Medical Sciences, Tehran, Iran; (5) Department of Gynecology, School of Medicine, Arash Hospital, Tehran University of Medical Sciences, Tehran, Iran; (6) Department of Histology, School of Medicine, Tehran University of Medical Sciences, Tehran, Iran; (7) School of Pharmacy, Shahid Beheshti University of Medical Sciences, Tehran, Iran; (8) Department of Physiology, Faculty of Medicine, Tehran Medical Sciences, Islamic Azad University, Tehran, Iran; (9) School of Persian Medicine, Qom University of Medical Sciences, Qom, Iran; (10) School of Traditional Medicine, Shahid Beheshti University of Medical Sciences, Tehran, Iran.

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Abstract

Diabetes mellitus (DM) and its complications impose a significant burden on patients and the health care system. In the Traditional Persian Medicine (TPM), barley is recommended for treatment of DM. This study sought to assess the effect of barley seed aqueous extract on hepatic, pancreatic, renal and cardiac tissues in normal (non-diabetic) and Streptozotocin-induced diabetic rats. Twenty-one male Wistar rats were randomly divided into diabetic and non-diabetic groups. Diabetes was induced by single intraperitoneal injection of Streptozotocin. After one week, the diabetic and non-diabetic groups were randomly divided into control and barley seed extract subgroups namely N group (non-diabetic control rats), S group (seed extract treated non-diabetic rats), D group (diabetic control rats) and DS group (seed extract treated diabetic rats). After 6 weeks, all rats were sacrificed for histopathological analysis and specimens were stained routinely for histological studies. The abnormal histological signs significantly decreased in the DS group compared to D group. Also, protective effects of barley seed extract against histopathological changes were seen in S group compared to N group. These findings suggest that barley seed extract exerts a protective effect on different tissues in diabetes.

Key Words: Barley; diabetes; traditional persian medicine; Streptozotocin; diabetes mellitus.

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Diabetes mellitus (DM) is a group of common metabolic disorders with the common phenotype of hyperglycemia. This condition can lead to adverse consequences for the patient and impose an heavy burden on the health care system.¹ DM type 1 is a form of DM that results from destruction of beta cells in the pancreas which leads to absolute insulin insufficiency.² It is responsible for approximately 5–10% of all cases of diabetes; however, its incidence is increasing worldwide

and it has serious short-term and long-term complications.³ Some tissue-specific injuries also occur. In the liver histological changes can vary from steatosis to steatohepatitis and liver fibrosis.^{4,5} DM contributes greatly to mortality due to the high incidence of nephropathy leading to end stage renal disease. Indeed it is a major cause of dialysis and kidney transplantation.⁶ Diabetes facilitates pathological changes in the cardiovascular system of these patients, increasing risks

of cardiovascular complications (acute coronary syndromes, stroke, heart failure and arrhythmias).⁷ High cost of treatment and side effects of medications, namely increased risk of cardiovascular diseases, hypoglycemia and weight gain, are among the common complications of pharmacologic therapies.⁸⁻¹¹

Traditional Persian Medicine (TPM) has gained attention in the recent years and some of the new medications originate from traditional medicine.^{12,13} TPM, one of the traditional healthcare systems, has been practiced since thousand years.¹⁴⁻¹⁶ Their theoretical backgrounds may offer the bases for development of new drugs.¹⁷⁻¹⁹

According to Persian traditional medicine barley grain (*Hordeum vulgare* L.) is recommended for treatment of diabetes. Outstanding books such as *Al-Havi* (*Liber continens*; written by Rhazes in 10th century AD)²⁰ and *Al-Qanun fi al-Tib* (*Canon of Medicine*; written by Avicenna in 1025 AD)²¹ recommended the use of medicinal beer to quench thirst. This drink is made by boiling barley in water and is claimed to be beneficial for diabetic patients.

Modern studies also shows positive effects of drinking barley aqueous extract. In a study by Naseri et al. in diabetic rats, they caused a significant reduction in serum glucose level after 4 weeks of consumption.²² Furthermore, antioxidant properties of phenolic compounds in barley add to its nutritional value.²³⁻²⁵ Antioxidants play an important role in prevention of chronic diseases by elimination of reactive oxygen species (ROS). It appears that the antioxidant properties of barley are related to the flavan-3-ol, hydroxycinnamic acid derivatives and flavonols.²⁶

Considering increased amount of free radicals in diabetic patients,²⁷⁻²⁹ and presence of abundant antioxidants in barley,²³⁻²⁶ this study was designed to evaluate the effect of barley seed extract on liver, pancreas, kidney and heart in control and diabetic rats.

Materials and Methods

Extract Preparation

Ready for harvest barley grains 4 months after sowing were purchased from an agricultural farm in Mehrshahr Karaj. After ensuring the apparent health of grains the voucher specimen was deposited at herbarium of School of Shahid Beheshti University with voucher number 580. The method of extract preparation in this study was adapted from the traditional medicine instructions on how to prepare a medicinal beer.^{20,30-33} For extract preparation, 500 g barley seeds in 1000ml of distilled water were heated in a beaker. After the contents of the beaker were boiled, the boiling process were continued for 10 minutes and after cooling down, were filtered using a laboratory filter paper. The obtained solution was concentrated, weighed 97 g (19.4%) and stored in a freezer at -20°C (1 %, weight per weight, W/W). Barely aqueous extract was mixed with standard food to prepare the food for the treatment groups.

Total phenolic content of barley extract

Total phenolic content of barely aqueous extract was detected using spectrophotometric method. To determine phenolic concentration, the reaction mixture was use by mixing 1 ml barely aqueous extract (2 mg/ml), Folin-Ciocalteu's reagent 10% (5 ml) and 4 ml NaHCO₃ (7.5 mg/ml). Blank was prepared using water without extract. Samples were incubated at room temperature for 30 min. The absorbance was detected using spectrophotometer at λ_{max} = 765 nm. All samples for analysis were prepared in triplicate. The same procedure was repeated for the gallic acid as standard in concentrations of 10, 20, 40, 80 µg/ml. To achieve phenolic concentration the gallic acid (GA) calibration curve based on the detected absorbance was construed.^{34,35}

Animal Ethics

All experimental protocols were approved by the local animal care committee in accordance with Tehran University of Medical Sciences guidelines and Use of Laboratory Animals of the National Institute of Health Bethesda United State of America for the care and use of laboratory animals.³⁶

Animals

This study was conducted on 21 white male Wistar rats weighing 200 to 300g in the age range of 6 to 8 months. The rats had been transferred from the Razi Institute to the animal room of the School of Medicine, Shahed University, 2 weeks earlier. During the study, animals had free access to water and food. Animals were fasted for 12h and weighed using a digital scale. Blood was taken from the caudal artery for the measurement of fasted blood glucose level.³⁷ All study phases followed the guide for the care and use of laboratory animals.³⁸

Diabetes induction

For induction of type I diabetes, 7 rats were fasted for 12h. Streptozotocin powder (Sigma, USA) was completely dissolved in freshly prepared 0.05 molar citrate buffered solution with a pH of 4.5³⁹. The final concentration of Streptozotocin in citrate buffered solution was 13.5 mg/ml. In less than 15min,⁴⁰ the prepared solution was intraperitoneally injected to rats as single dose using an insulin syringe in such a way that the pure dose of injected Streptozotocin was 45 mg/kg for each rat.^{39,41-43} After one week, diabetic rats were weighed, fasted. Symptoms of diabetes including weight loss, polydipsia and polyuria were seen in the diabetic animals. Blood was taken and fasted blood glucose level was assessed to confirm the induction of diabetes. Fasted serum glucose level over 200 mg/dL was considered as the criterion for diabetes.⁴⁴

Conduction of study

Two weeks' time was allowed for rats to become familiar with the environment and the eating habits. Male Wistar rats were randomly divided into 2 groups of 14 non-diabetic and 7 diabetic rats. In the diabetic rats, diabetes

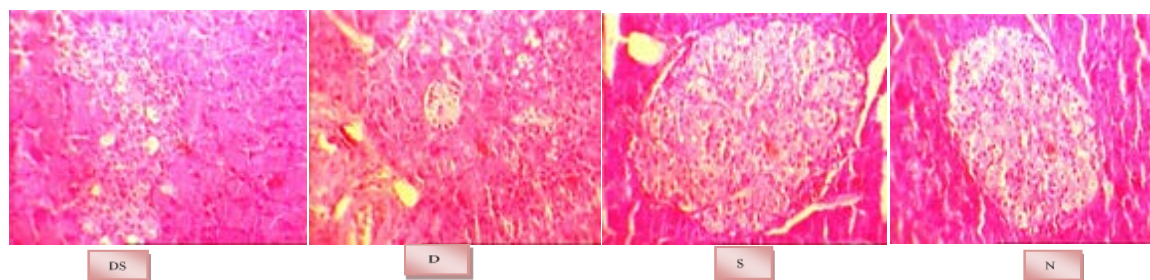


Fig 1. Histological views of the pancreas. N: Non-diabetic control group, S: Non-diabetic group receiving barley extract, D: Diabetic control group and DS: Diabetic group receiving barley extract.

was induced by single dose intraperitoneal injection of 45 mg/kg Streptozotocin. Fasted serum glucose level of rats in the diabetic group was measured one week later and induction of diabetes was confirmed by a serum glucose level over 200 mg/dL. Diabetic and non-diabetic groups were then randomly divided into 2 subgroups as follows:

1. Non-diabetic control group (N) with 7 rats
2. Non-diabetic group receiving barley extract (S) with 7 rats
3. Diabetic control group (D) with 4 rats
4. Diabetic group receiving barley extract (DS) with 3 rats

It should be mentioned that the control groups received regular food (Pars, Iran); while the test groups received a combination of extract and the standard food. Two weeks after the completion of the 4-week experiment, animals were sacrificed and their liver, pancreas, kidneys and heart were removed and used for histological studies. For this purpose, organs (tissues) were fixed in 10% formaldehyde for 24h and dehydrated using sequential ethanol concentrations. Specimens were then embedded in paraffin blocks and cut into 4-5µm thick sections for staining. Then haematoxylin-eosin staining (H&E) was done. After staining, specimens were histologically evaluated under a light microscope. For determination of tissue changes, a 0-2 scoring system was used compared to the control group.⁴⁵⁻⁴⁷

Results

Total Phenols contents of barely aqueous extract was determined as 3.61 ± 7 mg/g GA equivalent using the spectrophotometric method. In histomorphological study of pancreas in the diabetic group (D), atrophy of the islets of Langerhans, reduction in their size and number, significant reduction in cell differentiation, destruction and necrosis of beta cells and infiltration of mononuclear cells in the islets of Langerhans were observed. Such abnormal histological changes significantly decreased in the diabetic group that consumed barley extract (DS) compared to the diabetic control group. The non-diabetic group receiving barley extract (S) showed healthier pancreatic tissue compared to the non-diabetic control group (N). Histological evaluation of pancreatic tissue revealed that the S group had increased number of islets of Langerhans, increased number of undifferentiated cells around the islets of Langerhans and increased capillaries (Figure 1 and Table 1).

Histomorphological study of the liver: In the diabetic control group (D), fatty liver, local infiltration of mononuclear cells, increased number of Kupffer cells, fibrosis of the portal spaces and sinusoidal dilatation and hemorrhage were seen. Such abnormal histological changes showed a significant reduction in the DS group compared to D group. The comparison of N and S groups

Table 1. Comparison of the different characteristics of pancreatic tissues in the understudy groups after hematoxylin and eosin staining.

Pancreatic tissue	Reduction of Island size	Reduction of Island number	Reduction of the number of granules in beta cells	Increment of capillary in Island	Increase in differentiation	Reduction in differentiation	Infiltration of Mononuclear cells
N	-	-	-	-	-	-	+
D	+	+	+	-	-	++	++
DS	+	+	+	+	+	-	+
S	-	-	-	++	++	-	+

The scores were calculated by a pathologist from zero to 2. N: Non-diabetic control group, S: Non-diabetic group receiving barley extract, D: Diabetic control group, and DS: Diabetic group receiving barley extract.

Table 2. Comparison of different characteristics of liver tissue in the understudy groups after H & E staining.

Liver tissue	Fatty liver	Congestion	Infiltration of Mononuclear in Lobule	Kupffer cell Hyperplasia	Fibrosis in portal space	Hemorrhage in Sinusoidal dilation
N	-	-	+	-	-	-
D	+	+	+	+	+	+
DS	-	-	++	-	-	-
S	-	-	+	-	-	-

The scores were calculated by a pathologist from zero to 2. N: Non-diabetic control group, S: Non-diabetic group receiving barley extract, D: Diabetic control group, and DS: Diabetic group receiving barley extract.

revealed that these two groups had similar histology and a normal liver structure (Table 2).

Histomorphological study of the kidneys: In group D, bleeding in the cortex of kidneys, hyperemia, glomeruli to capsule adhesion, increased number of glomerular cells, increased thickness of glomerular endothelium, destruction of distal convoluted tubule (DCT) and proximal convoluted tubule (PCT) epithelial cells and infiltration of mononuclear cells were seen. The comparison of D and DS groups revealed that DS group had healthier organs. Groups N and S were found to be histologically similar (Table 3).

Histomorphological study of the heart: In group D, atrophy of the cardiac muscle cells in the apex of the heart, hypertrophy of the atrial cardiac muscle cells, hyperemia, bleeding, hypochromic atrial muscle cells and infiltration of mononuclear cells were noted. Group DS had healthier tissues than group D. Groups N and S were histologically similar (Table 4).

Discussion

DM leads to severe metabolic imbalance and changes in tissues. Oxidative stress seems to play an important role in this imbalance. Diabetic animal models show significant oxidative stress due to chronic hyperglycemia

that decreasing the antioxidant defense system increases the production of free radicals.^{48,49} Free radicals play an important role in the DM complications.⁵⁰ Plants are among the antioxidant sources, specifically grains due to their phenolic compounds have antioxidant properties.²⁵ Grains of barley are a rich source of phenolic compounds.²⁴ Persian traditional medicine has emphasized the beneficial effects of barley on DM. The efficacy of medicinal beer, made by boiling barley in water, for treatment of DM has also been documented.^{20,21} Furthermore, in a study conducted in 2010 by Naseri et al.,²² it was demonstrated that barley aqueous extract decreased hyperglycemia in diabetic rats. In the present study, we evaluated the effect of barley extract on pancreatic, hepatic, renal and cardiac tissues in Streptozotocin-induced diabetic rats. The data show that barley aqueous extract significantly decreases the histological signs of disease in the diabetic group receiving the extract compared to the diabetic control group. Also, our findings show that the pancreatic tissue in the non-diabetic group receiving the extract seemed healthier histologically compared to the non-diabetic control group. Our preliminary data may indicate the protective and therapeutic effects of barley aqueous extract in diabetic rats. Evaluation of oxidative stress

Table 3. Comparison of different characteristics of renal tissues in the understudy groups after H & E staining.

Renal tissue	Hemorrhage in cortex	Congestion	Glomeruli to capsule adhesion	Hyperplasia of internal mesangial	Thickness of glomerular endothelium	Degeneration of PCT & DCT Epithelia	Infiltration of mononuclear cells
N	-	-	-	-	-	-	+
D	++	+	++	++	++	++	++
DS	+	+	+	+	+	+	+
S	-	-	-	-	-	-	+

The scores were calculated by a pathologist from zero to 2. N: Non-diabetic control group, S: Non-diabetic group receiving barley extract, D: Diabetic control group, and DS: Diabetic group receiving barley extract.

Table 4. Comparison of different characteristics of liver tissue in the understudy groups after H & E staining.

Cardiac tissue	Atrophy in apex	Hypertrophy in atrium	Congestion	Hemorrhage	Pale of cytoplasm	Infiltration of mononuclear cells
N	-	-	-	-	-	+
D	+	+	+	++	+	+
DS	-	-	-	-	-	+
S	-	-	-	-	-	+

The scores were calculated by a pathologist from zero to 2. N: Non-diabetic control group, S: Non-diabetic group receiving barley extract, D: Diabetic control group, and DS: Diabetic group receiving barley extract.

markers in diabetic rats indicated that ROS, increased in the islets of Langerhans.²⁹ Chanpoo et al.,²⁸ in a study in 2010 evaluated the effect of consumption of curcumin on the pancreatic tissue in rats and reported the infiltration of mononuclear cells in the islets of Langerhans and reduction in size and number of islets in the diabetic group. Our preliminary data are in accord with those findings. They also concluded that curcumin had antioxidant properties and in the diabetic groups resulted in increase of pancreatic islets after 12 weeks.²⁸ Thymoquinone is among the main ingredients of *Nigella sativa* plant and has a high antioxidant potential. By reducing oxidative stress, it preserve the islets of Langerhans and the beta cells.⁵¹ Infiltration of mononuclear cells and sinusoidal dilatation as well as signs of tissue necrosis in the liver of diabetic rats reported by Maisaa et al.,⁵² are also seen in our study. They evaluated the effect of *Trifolium alexandrinum* flower aqueous solution on the hepatic tissue of diabetic rats and showed that this extract may improve the histological changes in diabetic rats. This function may be attributed to the high flavonoid content acting as antioxidant.⁴ Moreover, in our study, adhesion of glomeruli to capsule, increased number of glomeruli cells, thickening of the glomerular endothelium and destruction of epithelial cells in DCT and PCT were seen in the renal tissue of diabetic group. All these signs were also reported by Gross et al., in their study on the renal tissue of diabetic rats.⁵³ Many similar studies have investigated the effects of different medicinal plants with antioxidant properties on the diabetic renal tissues with significant therapeutic effects.^{54,55} Histological analysis of diabetic cardiac tissue revealed changes in the myocardial cells, bleeding in the cardiac tissue, hypochromic atrial muscle cells and infiltration of mononuclear cells. Other studies have also demonstrated that cardiac ischemia is more common and more severe in diabetic patients and DM can cause significant cardiovascular diseases.⁵⁶ Studies have reported that oxidative stress in DM is the main cause of cardiovascular diseases.⁵⁷ Based on our preliminary data, we may conclude that barley aqueous extract has beneficial effects on

prevention of tissue changes in DM. This property is probably due to its antioxidant characteristics, but further studies are required to confirm by quantitative statistical analyses the here reported qualitative data and precisely determine the active ingredients of barley aqueous extract.

List of acronyms

AD: Anno Domini
DCT - destruction of distal convoluted tubule
DM - diabetes mellitus
DS - diabetic group that consumed barley extract
GA - gallic acid
N - non-diabetic control group
PCT - proximal convoluted tubule
ROS - Reactive Oxygen Species
S - non-diabetic group receiving barley extract
TPM - Traditional Persian Medicine
W/W - weight per weight

Contributions of Authors

All authors made substantial contributions to the conception, design, and analysis and interpretation of data for this work. All authors contributed substantially to the drafting and critical revision of the manuscript, provided final approval of the version submitted, and agree to be held accountable for all aspects of the work.

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Conflict of Interest

The authors declare no conflict of interests.

Ethical Publication Statement

We confirm that we have read the Journal's position on issues involved in ethical publication and affirm that this report is consistent with those guidelines.

Corresponding Author

Dr. Farzaneh Ghaffari. School of Traditional Medicine, Shahid Beheshti University of Medical Sciences, Tehran, Iran. Tel: +989123881874

ORCID iD: 0000-0002-5424-4348

E-mail: f_ghaffari@sbmu.ac.ir

E-mails and ORCID iD of co-authors

Mohsen Naseri: naseri@shahed.ac.ir

ORCID iD: . 0000-0001-9239-768X

Zahra Khalaj Sereshki: dr.zahrakhalaj@gmail.com

ORCID iD: 0000-0002-7208-7964

Behnaz Ghavami: Email: Behnaz.ghavami@gmail.com

ORCID iD: 0000-0002-5192-1745

Bagher Minaii Zangii: minaebz@sina.tums.ac.ir

ORCID iD: 0000-0001-5064-2448

Mohammad Kamalinejad: mkamalinejad@yahoo.com

ORCID iD: 0000-0002-5946-4913

Parvaneh Mohseni Moghaddam:

p.mohseni@iautmu.tums.ac.ir

ORCID iD: 0000-0002-0608-2305

Majid Asghari: asghari.rall@gmail.com

ORCID iD: 0000-0003-1155-3172

Seyed Abbas Hashemi Nejad:

abbas.hasheminejad@shahed.ac.ir

ORCID iD: 0000-0001-9635-7669

Fatemeh Emadi: fatemeh12emadi@yahoo.com

ORCID iD: 0000-0002-5053-4591

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