

Clinical Trial Protocol

Iranian Registry of Clinical Trials

03 Jun 2026

Evaluation of the effects of Chia seeds (*Salvia hispanica*) on nutritional status, metabolic factors, liver function, inflammatory hematologic and atherogenic indices in obese patients with non-alcoholic fatty liver disease

Protocol summary

Study aim

To determine the effects of Chia seeds on nutritional status, metabolic factors, liver function, inflammatory hematologic and atherogenic indices in obese patients with non-alcoholic fatty liver disease

Design

A single-blind, controlled, randomized clinical trial using random assignment software (RAS) and randomized block method on 38 patients.

Settings and conduct

Study location: Tabriz University of Medical Sciences Nutrition Science Research Center; Study population: patients with non-alcoholic fatty liver disease; blinding: single-blind, in this study, the person who analyzes is blinded; A research assistant not otherwise involved in the study, will randomly allocate the participants into one of the two experimental groups (1:1), using the Random allocation software (RAS) and randomized block procedure. The person (completely unrelated to the study) who will prepare the supplement sachets will assign a 3-digit code for each treatment.

Participants/Inclusion and exclusion criteria

Inclusion criteria: both genders; age 18 to 60 years; body mass index 30- 39.9 Kg/m; hepatic steatosis; willing to participate in the study. Exclusion criteria: athletes; pregnancy; breastfeeding and menopause; following a special diet; taking medications for weight-reducing or any other medications affecting liver function for 3 months before or during the study; suffering from certain diseases

Intervention groups

The intervention group will receive two- 20 gr sachets of chia seeds per day along with weight loss diet while the control group will only receive weight loss diet

Main outcome variables

Nutritional status (intake of energy and macronutrients,

appetite status); Anthropometric indices and body composition; Metabolic factors; Liver function and liver fibrosis score; Hematological indicators of inflammation

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20100209003320N23**

Registration date: **2024-01-30, 1402/11/10**

Registration timing: **prospective**

Last update: **2024-01-30, 1402/11/10**

Update count: **0**

Registration date

2024-01-30, 1402/11/10

Registrant information

Name

Mehrangiz Ebrahimi mamagani

Name of organization / entity

Health & Nutrition faculty of Tabriz university of medical sciences

Country

Iran (Islamic Republic of)

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Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2024-02-20, 1402/12/01

Expected recruitment end date

2024-07-22, 1403/05/01

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Evaluation of the effects of Chia seeds (*Salvia hispanica*) on nutritional status, metabolic factors, liver function, inflammatory hematologic and atherogenic indices in obese patients with non-alcoholic fatty liver disease

Public title

The effect of chia seeds on non-alcoholic fatty liver disease

Purpose

Treatment

Inclusion/Exclusion criteria**Inclusion criteria:**

Both genders Age range of 18 to 60 years body mass index= 30- 39.9 Kg/m² Hepatic steatosis based on ultrasound findings (grade 1 and 2) willingness to participate in the study

Exclusion criteria:

Athletes pregnancy breastfeeding menopause Under infertility treatment taking birth control pills estrogen therapy Smoking history of alcohol consumption Taking medications including chemical or herbal medications for weight loss anti-hypertensive Fat-lowering drugs Sugar-lowering drugs any nutritional supplements from 3 months before the study Having cardiovascular diseases Liver dysfunction Kidney dysfunction Intestinal dysfunction Thyroid dysfunction Parathyroid dysfunction known autoimmune diseases polycystic ovary syndrome cancers Following a special diet in the last three months performing weight loss surgery in the last one year Taking Phenytoin medicine Taking amoxifene medicine Taking lithium medicine Performing or being a candidate for liver transplantation

Age

From **18 years** old to **60 years** old

Gender

Both

Phase

3

Groups that have been masked

- Data analyser

Sample size

Target sample size: **38**

Randomization (investigator's opinion)

Randomized

Randomization description

A research assistant not otherwise involved in the study, will randomly allocate 38 patients into one of the two experimental groups (1:1), using the Random allocation software (RAS) and randomized block procedure (age (18-39 vs 39-60 years)- gender (female vs male) and BMI (<35 kg/m² vs. ≥35 kg/m²)).

Blinding (investigator's opinion)

Single blinded

Blinding description

A research assistant not otherwise involved in the study, randomly allocated the participants into one of the two experimental groups (1:1), using the Random allocation software (RAS) and randomized block procedure. The person (completely unrelated to the study) who will prepare the supplement sachets will assign a 3-digit code for each treatment.

Placebo

Not used

Assignment

Parallel

Other design features**Secondary Ids**

empty

Ethics committees**1****Ethics committee****Name of ethics committee**

Ethics Committee of Tabriz University of Medical Sciences

Street address

Faculty of Nutrition and Food Sciences, Tabriz University of Medical Sciences, Attar Neishaburi St, Golgasht St.

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Tabriz

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East Azarbaijan

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Approval date

2024-01-01, 1402/10/11

Ethics committee reference number

IR.TBZMED.REC.1402.751

Health conditions studied**1****Description of health condition studied**

Non-alcoholic fatty liver disease

ICD-10 code

K76.0

ICD-10 code description

Fatty (change of) liver, not elsewhere classified

Primary outcomes**1****Description**

Insulin

Timepoint

Baseline and after 8 weeks

Method of measurement

ELISA method

2**Description**

Hemoglobin A1C

Timepoint

Baseline and after 8 weeks

Method of measurement

chromatography

3**Description**

total cholesterol

Timepoint

Baseline and after 8 weeks

Method of measurement

Enzymatic method

4**Description**

Triglyceride

Timepoint

Baseline and after 8 weeks

Method of measurement

Enzymatic method

5**Description**

HDL cholesterol

Timepoint

Baseline and after 8 weeks

Method of measurement

Enzymatic-colorimetric method using spectrophotometer

6**Description**

LDL cholesterol

Timepoint

Baseline and after 8 weeks

Method of measurement

Based on the Friedelwald formula

7**Description**

Insulin resistance indices (HOMA-IR, HOMA-B and QUICKI)

Timepoint

Baseline and after 8 weeks

Method of measurement

Based on the formula

8**Description**

Body mass index(BMI)

Timepoint

Baseline and after 8 weeks

Method of measurement

Based on the formula

9**Description**

Waist circumference

Timepoint

Baseline and after 8 weeks

Method of measurement

tape measure

10**Description**

Waist to hips ratio (WHR)

Timepoint

Baseline and after 8 weeks

Method of measurement

Based on the formula

11**Description**

Waist to Height ratio (WHTR)

Timepoint

Baseline and after 8 weeks

Method of measurement

Based on the formula

12**Description**

Fat mass

Timepoint

Baseline and after 8 weeks

Method of measurement

Body analyzer

13**Description**

Fat Free mass

Timepoint

Baseline and after 8 weeks

Method of measurement

Body analyzer

14**Description**

Energy, macronutrients and micronutrients intake

Timepoint

Baseline and after 8 weeks

Method of measurement

24- hr food recall questionnaire for 3 days and Nutritionist 4 software

15**Description**

Physical activity level

Timepoint

Baseline and after 8 weeks

Method of measurement

16

Description

Appetite status

Timepoint

Baseline and after 8 weeks

Method of measurement

Validated appetite questionnaire

17

Description

Body weight

Timepoint

Baseline and after 8 weeks

Method of measurement

Scale

18

Description

Fasting glucose

Timepoint

Baseline and after 8 weeks

Method of measurement

Enzymatic method

19

Description

Complete blood cell count (CBC)

Timepoint

Baseline and after 8 weeks

Method of measurement

Coulter counter

20

Description

Serum Albumin

Timepoint

Baseline and after 8 weeks

Method of measurement

using BROMOCRESOL GREEN

21

Description

Platelets count

Timepoint

Baseline and after 8 weeks

Method of measurement

Coulter counter

22

Description

Liver fibrosis score

Timepoint

Baseline and after 8 weeks

Method of measurement

based on formula

23

Description

total body water

Timepoint

Baseline and after 8 weeks

Method of measurement

Using bioelectric impedance analyzer

24

Description

Hematological indicators of inflammation (the number of monocytes, neutrophils, lymphocytes and platelets in whole blood and the estimation of the ratios of monocytes, neutrophils and platelets to lymphocytes)

Timepoint

Baseline and after 8 weeks

Method of measurement

Coulter counter

25

Description

Atherogenic indices (LDL-C/HDL-C, TC/HDL-C, TG/HDL-C, nonHDL-C/HDL-C)

Timepoint

Baseline and after 8 weeks

Method of measurement

Based on the formula

26

Description

Hip circumference

Timepoint

Baseline and after 8 weeks

Method of measurement

tape measure

27

Description

Serum ferritin

Timepoint

Baseline and after 8 weeks

Method of measurement

Electrochemiluminescence method

Secondary outcomes

1

Description

Fatty liver grade

Timepoint

Baseline and 8 weeks after intervention

Method of measurement

Ultrasonigraphy findings

2

Description

Alanine aminotransferase (ALT)

Timepoint

Baseline and 8 weeks after intervention

Method of measurement

Enzymatic method

3

Description

Aspartate aminotransferase (AST)

Timepoint

Baseline and 8 weeks after intervention

Method of measurement

Enzymatic method

Intervention groups

1

Description

"Intervention group": patients in this group will receive chia seeds along with a weight loss diet, taking into account that 100 grams of chia seeds contain 486 calories (approximately 200 calories for 40 grams of chia seeds per day) to moderate the amount of weight loss. Calories in the intervention group will be reduced by 700 kilocalories and to adjust the diet with the distribution of macronutrients as 55% carbohydrates, 30% fat, 15% protein. Daily, two 20-gram sachets of Paraguayan chia seeds will be purchased from Isfahan Herbal Company in sachet form. 20 grams will be packed, half an hour before two meals a day, and they will be consumed in a glass of lukewarm water. The duration of the study will be 8 weeks. People in the intervention group will be asked to bring unused sachets with them every two weeks to determine compliance. Weight control and distribution of sachets will be done every two weeks

Category

Treatment - Drugs

2

Description

"Control group": patients in this group will only receive a weight loss diet (500 kcal reduction) along with a glass of water before two main meals for 8 weeks.

Category

Lifestyle

Recruitment centers

1

Recruitment center

Name of recruitment center

Tabriz University of Medical Science

Full name of responsible person

Dr.Mehrangiz Ebrahimi- Mameghani

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Tabriz University of Medical Sciences

Full name of responsible person

Mohammad Samiei

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ebrahimimamagani@tbzmed.ac.ir

Grant name

Grant code / Reference number

Is the source of funding the same sponsor organization/entity?

Yes

Title of funding source

Tabriz University of Medical Sciences

Proportion provided by this source

100

Public or private sector

Public

Domestic or foreign origin

Domestic

Category of foreign source of funding

empty

Country of origin

Type of organization providing the funding

Academic

Person responsible for general inquiries

Contact

Name of organization / entity

Tabriz University of Medical Sciences

Full name of responsible person

Dr. Mehrangiz Ebrahimi- Mameghani

Position

professor

Latest degree

Ph.D.

Other areas of specialty/work

Nutrition

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Bachelor

Other areas of specialty/work

Nutrition

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Person responsible for scientific inquiries**Contact****Name of organization / entity**

Tabriz University of Medical Sciences

Full name of responsible person

Dr. Mehrangiz Ebrahimi- Mameghani

Position

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Latest degree

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Other areas of specialty/work

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Person responsible for updating data**Contact****Name of organization / entity**

Tabriz University of Medical Sciences

Full name of responsible person

Maryam Parimi

Position

student

Latest degree**Sharing plan****Deidentified Individual Participant Data Set (IPD)**

Yes - There is a plan to make this available

Study Protocol

Yes - There is a plan to make this available

Statistical Analysis Plan

Yes - There is a plan to make this available

Informed Consent Form

Yes - There is a plan to make this available

Clinical Study Report

Yes - There is a plan to make this available

Analytic Code

Yes - There is a plan to make this available

Data Dictionary

Yes - There is a plan to make this available

Title and more details about the data/document

Data collected for the primary outcomes will be shared

When the data will become available and for how long

The access period starts 12 months after the results are published

To whom data/document is available

The data will be available only to people working in scientific institutions

Under which criteria data/document could be used

The data of this study will be available to other researchers only for meta-analysis studies

From where data/document is obtainable

Maryam Parimi, email address:

maryam.parimi6@gmail.com, phone number:

09109337321

What processes are involved for a request to access data/document

The applicant should provide a brief explanation of the goals and methodology of his meta-analysis. The applicant's request will be reviewed and if approved, the data will be sent to the applicant via email. All these steps will not take more than 15 days

Comments