

Clinical Trial Protocol

Iranian Registry of Clinical Trials

17 Jun 2026

Evaluating of the effect of educational intervention based on the developed parallel process model on the adherence to the proper diet, glycemic factors, and lipid profile in middle-aged people with type 2 diabetes

Protocol summary

Study aim

Effect of educational intervention based on the developed parallel process model on the adherence to proper diet, glycemic control, and lipid profile in middle-aged people with type 2 diabetes

Design

A 4-arm randomized, controlled, single-blind, factorial field trial (with control group) on 88 participants was designed. Centers were randomly allocated to the control or intervention groups and in each center, 22 were selected by simple random sampling. Nutrition education including Gain Frame Messages, Loss Frame Messages, or a combination of both types of messages versus usual diabetic nutritional education were conveyed to 88 type 2 diabetic patients aged 30 to 59 years.

Settings and conduct

Nutritional education including Gain Frame Messages, Loss Frame Messages, or their combination versus usual nutritional education were conveyed to diabetic patients in Bownat city for 3 months to assess their effects on dietary adherence, biochemical, anthropometric, and glycemic control markers. Data analyzer was not informed of study groupings (single-blind).

Participants/Inclusion and exclusion criteria

Age range: 30-59 years old (middle age); having type 2 diabetes and not using insulin (FBS higher than 125 and 2-hr glucose higher than 200 mg/dl); not having chronic diseases including cancer; kidney or liver diseases not using special drugs; not smoking; having no special diet; not being pregnant or lactating; not having sudden disturbances of blood glucose including sudden and dangerous hypo- or hyperglycemia;

Intervention groups

1) Gain Frame Messages (GFM) 2) Loss frame Messages (LFM) 3) Combination of both types of

messages (G\LFM) 4) Control (CG)

Main outcome variables

Hb A1C, blood glucose

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20230104057040N1**

Registration date: **2023-05-08, 1402/02/18**

Registration timing: **retrospective**

Last update: **2023-05-08, 1402/02/18**

Update count: **0**

Registration date

2023-05-08, 1402/02/18

Registrant information

Name

Tayebe Dehghan

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 917 566 2643

Email address

dehghant89@yahoo.com

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2021-12-22, 1400/10/01

Expected recruitment end date

2022-05-20, 1401/02/30

Actual recruitment start date

2022-01-21, 1400/11/01

Actual recruitment end date

2022-04-19, 1401/01/30

Trial completion date

2022-08-21, 1401/05/30

Scientific title

Evaluating of the effect of educational intervention based on the developed parallel process model on the adherence to the proper diet, glycemic factors, and lipid profile in middle-aged people with type 2 diabetes

Public title

Nutrition education with extended parallel model in diabetic patients for improving glycemic indices and lipid profile control and adherence to the proper diet

Purpose

Prevention

Inclusion/Exclusion criteria**Inclusion criteria:**

Age range: 30-59 years old (middle age) Having type 2 diabetes and not using insulin (FBS higher than 125 and 2-hr glucose higher than 200 mg/dl) Not having chronic diseases including cancer, kidney or liver diseases Not using special drugs, not smoking Having no special diet Not being pregnant or lactating

Exclusion criteria:

Incidence of any chronic or acute diseases or incidence of some diseases such as kidney diseases Sudden disturbances of blood glucose including sudden and dangerous hypo- or hyperglycemia Acute or respiratory infections Any surgery Following any special diet or physical activity regiment

Age

From **30 years** old to **59 years** old

Gender

Both

Phase

3

Groups that have been masked

- Data analyser

Sample size

Target sample size: **88**

Actual sample size reached: **80**

Randomization (investigator's opinion)

Randomized

Randomization description

For data gathering, first, name and address of all health centers of Bowanat city were extracted and from 8 centers, four centers were selected by random multi-stage method. After selecting 4 centers, each center was allocated to a group of intervention or control group by lottery, in order not to let the patients visit each other and not to distort the results. Then, from each center, 22 patients were selected from eligible middle-aged diabetic patients with simple random sampling.

Blinding (investigator's opinion)

Single blinded

Blinding description

the study was single blind. Data analyzer was not also

informed of groupings. At the time of analysis, the analyzer was informed that 4 groups with the names of A,B,C,D were available and he/she was not informed of group characteristics.

Placebo

Not used

Assignment

Factorial

Other design features**Secondary Ids**

empty

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

Shiraz University of Medical Sciences

Street address

Razi Avenue

City

Shiraz

Province

Fars

Postal code

7134814336

Approval date

2021-07-18, 1400/04/27

Ethics committee reference number

IR.SUMS.SCHEANUT.REC.1400.012

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

Type 2 Diabetes

ICD-10 code**ICD-10 code description****Primary outcomes****1****Description**

Glycosylated hemoglobin

Timepoint

At baseline and after 3 months

Method of measurement

High performance liquid chromatography (HPLC)

2**Description**

Fasting blood glucose

Timepoint

At baseline and after 3 months

Method of measurement

Calorimetric method by using auto-analyzer

3

Description

2 hour- blood glucose

Timepoint

At baseline and after 3 months

Method of measurement

Calorimetric method by using auto-analyzer

Secondary outcomes

1

Description

Low density lipoprotein-cholesterol

Timepoint

At baseline and after 3 months

Method of measurement

Calorimetric method by using auto-analyzer

2

Description

High density lipoprotein-cholesterol

Timepoint

At baseline and after 3 months

Method of measurement

Calorimetric method by using auto-analyzer

3

Description

Total cholesterol

Timepoint

At baseline and after 3 months

Method of measurement

Calorimetric method by using auto-analyzer

4

Description

Triglyceride

Timepoint

At baseline and after 3 months

Method of measurement

Calorimetric method by using auto-analyzer

5

Description

Weight

Timepoint

At baseline and after 3 months

Method of measurement

With scale

6

Description

Height

Timepoint

At baseline and after 3 months

Method of measurement

With tape meter

7

Description

Body mass index

Timepoint

At baseline and after 3 months

Method of measurement

With calculation from weight and height

8

Description

Waist circumference

Timepoint

At baseline and after 3 months

Method of measurement

With tape meter

9

Description

Hip circumference

Timepoint

At baseline and after 3 months

Method of measurement

With tape meter

10

Description

Waist to hip ratio

Timepoint

At baseline and after 3 months

Method of measurement

Calculate from waist and hip measurements

11

Description

Physical activity

Timepoint

At baseline and after 3 months

Method of measurement

With questionnaire

12

Description

Knowledge level

Timepoint

At baseline and after 3 months

Method of measurement

With questionnaire

13

Description

Perceived sensitivity

Timepoint

At baseline and after 3 months

Method of measurement

With questionnaire

14

Description

Perceived severity

Timepoint

At baseline and after 3 months

Method of measurement

With questionnaire

15

Description

Perceived effectiveness

Timepoint

At baseline and after 3 months

Method of measurement

With questionnaire

16

Description

Self-efficacy

Timepoint

At baseline and after 3 months

Method of measurement

With questionnaire

17

Description

Intention

Timepoint

At baseline and after 3 months

Method of measurement

With questionnaire

18

Description

Food intake of macro nutrients and calorie

Timepoint

3-day dietary recall

Method of measurement

With questionnaire

Intervention groups

1

Description

1- Control group:routine educations for 3 months (82 messages in the form of images and clips, almost 5 messages per day via virtual media such as messengers and by formation of virtual groups)

Category

Prevention

2

Description

1- Intervention group: gain-frame messages group for 3 months (82 messages in the form of images and clips, almost 5 messages per day via virtual media such as

messengers and by formation of virtual groups)

Category

Prevention

3

Description

2- Intervention group:loss-frame messages group for 3 months(78 messages in the form of images and clips, almost 5 messages per day via virtual media such as messengers and by formation of virtual groups)

Category

Prevention

4

Description

3- Intervention group: combined messages group for 3 months (157 messages in the form of images and clips, almost 5 messages per day via virtual media such as messengers and by formation of virtual groups)

Category

Prevention

Recruitment centers

1

Recruitment center

Name of recruitment center

Bowanat city health centers

Full name of responsible person

Tayebe Dehghan

Street address

Razi avenue

City

Shiraz

Province

Fars

Postal code

۱۴۳۳۶۷۱۳۴۸

Phone

+98 917 566 2643

Email

dehghant89@yahoo.com

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Shiraz University of Medical Sciences

Full name of responsible person

Zahra Sohrabi

Street address

Razi avenue

City

Shiraz

Province

Fars

Postal code

۱۴۳۳۶۷۱۳۴۸

Phone

+98 917 711 3086

Email

Zahra_2043@yahoo.com

Grant name

Grant code / Reference number

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

Yes

Title of funding source

Shiraz 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

Shiraz University of Medical Sciences

Full name of responsible person

Tayebe Dehghan

Position

Msc Student

Latest degree

Bachelor

Other areas of specialty/work

Nutrition

Street address

Razi avenue

City

Shiraz

Province

Fars

Postal code

۱۴۳۳۶۷۱۳۴۸

Phone

+98 917 566 2643

Email

dehghant89@yahoo.com

Person responsible for scientific inquiries

Contact

Name of organization / entity

Shiraz University of Medical Sciences

Full name of responsible person

Zahra Sohrabi

Position

Associate Professor

Latest degree

Ph.D.

Other areas of specialty/work

Nutrition

Street address

Razi avenue

City

Shiraz

Province

Fars

Postal code

۱۴۳۳۶۷۱۳۴۸

Phone

+98 917 711 3086

Email

Zahra_2043@yahoo.com

Person responsible for updating data

Contact

Name of organization / entity

Shiraz University of Medical Sciences

Full name of responsible person

Tayebe Dehghan

Position

Msc student

Latest degree

Bachelor

Other areas of specialty/work

Nutrition

Street address

Razi Avenue

City

Shiraz

Province

Fars

Postal code

۱۴۳۳۶۷۱۳۴۸

Phone

+98 917 566 2643

Email

dehghant89@yahoo.com

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

No - There is not a plan to make this available

Informed Consent Form

No - There is not a plan to make this available

Clinical Study Report

Yes - There is a plan to make this available

Analytic Code

No - There is not a plan to make this available

Data Dictionary

No - There is not a plan to make this available

Title and more details about the data/document

The final results will be shared.

When the data will become available and for how long

After publishing the article.

To whom data/document is available

Now it is available for the analyzer and the main investigators.

Under which criteria data/document could be used

Only with permission from the investigator

From where data/document is obtainable

By contacting the investigator

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

By asking from e-mail (email address: Sohrabi@sums.ac.ir)

Comments