

Clinical Trial Protocol

Iranian Registry of Clinical Trials

29 Aug 2026

Determination of the melatonin effect on gastrointestinal symptoms and quality of life in patients with irritable bowel syndrome with and without sleep disorders

Protocol summary

Study aim

Comparison of the melatonin effect on gastrointestinal symptoms and quality of life in irritable bowel syndrome patients with and without sleep disorders

Design

Double blind randomised _ paralalled clinical trial which participants randomly allocated to control (n=68) and melatonin supplementation groups(n=68)

Settings and conduct

In this study, patients referred to Imam Reza Hospital, after the definitive diagnosis of irritable bowel syndrome, using the ROME 4 diagnosis index, enter the study if they complete the informed consent form. According to the PSQI sleep questionnaire, patients were divided into two groups with and without sleep disorders. Each group are randomly divided into control and paralalled groups, and the paralalled group will receive 2 melatonin 3 mg tablets and the control group received two placebo.

Participants/Inclusion and exclusion criteria

Inclusion criteria: Patients diagnosed with irritable bowel syndrome based on ROME 4 diagnostic index. According to the PSQI sleep questionnaire, they are with and without sleep disorders. Exclusion criteria: depression, history of bowel surgery, dietary supplements and herbal remedies

Intervention groups

Paralalled group will receive 2 melatonin 3 mg tablets every 12 hours. The control group also will receive two placebo pills every 12 hours.

Main outcome variables

In this study, the effects of melatonin tablets on the severity of abdominal pain, bloating intensity, stool form, quality of life (Based on a valid quality of life questionnaire in patients with irritable bowel syndrome), sleep quality (Based on PSQI sleep quality questionnaire) in patients with irritable bowel syndrome will be evaluated in comparison with the control group.

General information

Reason for update

Acronym

IBS

IRCT registration information

IRCT registration number: **IRCT20220104053626N2**

Registration date: **2022-02-13, 1400/11/24**

Registration timing: **registered_while_recruiting**

Last update: **2022-02-13, 1400/11/24**

Update count: **0**

Registration date

2022-02-13, 1400/11/24

Registrant information

Name

Masood Dinevari

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 41 3334 7054

Email address

dinvarim@tbzmed.ac.ir

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2022-01-20, 1400/10/30

Expected recruitment end date

2022-04-18, 1401/01/29

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty	will be in. Placebo tablets are similar in appearance, color, and size to melatonin tablets.
Scientific title Determination of the melatonin effect on gastrointestinal symptoms and quality of life in patients with irritable bowel syndrome with and without sleep disorders	Placebo Used
Public title comparison of the melatonin effect in irritable bowel syndrome patients	Assignment Parallel
Purpose Supportive	Other design features
Inclusion/Exclusion criteria Inclusion criteria: being satisfied to participate in the study Patients with definitive diagnosis of IBS based on ROME 4 questionnaire, with sleep disorders diagnosed based on PSQI questionnaire. Patients with definitive diagnosis of IBS based on ROME 4 questionnaire, without sleep disorders Exclusion criteria: Previous history of diagnosed gastrointestinal diseases including IBD Intestinal surgery Depression Receive herbal medicines	Secondary Ids empty
Age No age limit	Ethics committees
Gender Both	1
Phase 2-3	Ethics committee Name of ethics committee Ethics Committee of Tabriz University of Medical Sciences Street address Tabriz University of Medical Sciences, Daneshgah St, Golgasht St. City Tabriz Province East Azarbaijan Postal code 5166614766
Groups that have been masked • Participant • Investigator	Approval date 2021-08-01, 1400/05/10 Ethics committee reference number IR.TBZMED.REC.1400.387
Sample size Target sample size: 136	Health conditions studied
Randomization (investigator's opinion) Randomized	1
Randomization description Among the patients who volunteered to participate in the study, 136 people will be selected by simple random sampling. Randomization method: block Randomization unit: individual Random layers: In each block, people will be matched based on age and gender. Random allocation tool: Random allocation software How to build a random sequence: Using Random allocation software Concealment: A random sequence created in a safe place and performed by an independent person not involved in the trial during the study. Random allocation of hidden individuals, patients and researchers will not be aware of it.	Description of health condition studied Irritable Bowel Syndrome ICD-10 code K58 ICD-10 code description Irritable bowel syndrome
Blinding (investigator's opinion) Double blinded	Primary outcomes
Blinding description This study is a double-blind study in which the researcher of this study and the patients participating in the study will be unaware of the type of supplement received. The supplements will be provided to patients by another person who has no role in completing the questionnaire. They will also be informed of the existence of two types of supplements (melatonin and placebo) when obtaining consent, but will be unaware of which study groups they	1 Description Abdominal pain severity based on a questionnaire on a percentage scale, From zero with an increase of 25 units Timepoint Before and after the intervention Method of measurement Irritable Bowel Syndrome Symptoms Severity Questionnaire
	2 Description Quality of life based on the quality of life questionnaire

for patients with irritable bowel syndrome

Timepoint

Before and after the intervention

Method of measurement

the quality of life questionnaire for patients with irritable bowel syndrome

3

Description

Intensity of bloating based on a questionnaire as a percentage scale, From zero with an increase of 25 units

Timepoint

Before and after the intervention

Method of measurement

Irritable Bowel Syndrome Symptoms Severity Questionnaire

4

Description

Stool form

Timepoint

Before and after the intervention

Method of measurement

Irritable Bowel Syndrome Symptoms Severity Questionnaire

Secondary outcomes

empty

Intervention groups

1

Description

Intervention group: Melatonin. Patients in the intervention group who will take two melatonin 3 mg tablets daily (manufactured by Zahravi Pharmaceutical Company) for two months during the study.

Category

Treatment - Other

2

Description

Control group: Control group: placebo. Patients in the placebo group who will take two placebo tablets daily (manufactured by Zahravi Pharmaceutical Company) during two months of study.

Category

Placebo

Recruitment centers

1

Recruitment center

Name of recruitment center

Emam reza hospital

Full name of responsible person

Farzaneh Jafarzadeh

Street address

Emam reza hospital, Golgast st, Tabriz.

City

Tabriz

Province

East Azarbaijan

Postal code

566614766

Phone

+98 41 3334 4918

Email

farzanehjafarzadeh998@gmail.com

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Tabriz University of Medical Sciences

Full name of responsible person

Masood dinevari

Street address

Tabriz University of Medical Sciences, Golgasht St, Tabriz

City

tabriz

Province

East Azarbaijan

Postal code

5165669531

Phone

+98 41 3335 8257

Email

masood.dinevari@gmail.com

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

Masood Dinevari

Position

Assistant Professor

Latest degree

Subspecialist

Other areas of specialty/work

Internal Medicine

Street address

Emam reza hospital, Gogash st, Tabriz

City

Tabriz

Province

East Azarbaijan

Postal code

5165665931

Phone

+98 41 3334 2500

Email

masood.dinevari@gmail.com

Person responsible for scientific inquiries

Contact

Name of organization / entity

Tabriz University of Medical Sciences

Full name of responsible person

Masood dinevari

Position

Assistant Professor

Latest degree

Subspecialist

Other areas of specialty/work

Internal Medicine

Street address

Emamreza hospital, Golgasht st, Tabriz

City

Tabriz

Province

East Azarbaijan

Postal code

5165665931

Phone

+98 41 3334 2669

Email

masood.dinevari@gmail.com

Person responsible for updating data

Contact

Name of organization / entity

Tabriz University of Medical Sciences

Full name of responsible person

Masood Dinevari

Position

Assistant Professor

Latest degree

Subspecialist

Other areas of specialty/work

Internal Medicine

Street address

Emamreza hospital, Golgasht st, Tabriz

City

Tabriz

Province

East Azarbaijan

Postal code

5165665931

Phone

+98 41 3336 4665

Email

masood.dinevari@gmail.com

Sharing plan

Deidentified Individual Participant Data Set (IPD)

Undecided - It is not yet known if there will be a plan to make this available

Study Protocol

No - There is not a plan to make this available

Statistical Analysis Plan

Undecided - It is not yet known if there will be a plan to make this available

Informed Consent Form

No - There is not a plan to make this available

Clinical Study Report

No - There is not a plan to make this available

Analytic Code

No - There is not a plan to make this available

Data Dictionary

Not applicable