

Clinical Trial Protocol

Iranian Registry of Clinical Trials

25 Jun 2026

Comparison of the Effectiveness of the Unified Protocol for Transdiagnostic Treatment (UP) and Cognitive Behavioral Therapy (CBT) on Emotional, Behavioral and Cognitive Components in Patients with Illness Anxiety Disorder Comorbid with Depression

Protocol summary

Study aim

The aim of the study is to compare the effectiveness of the Unified Protocol for Transdiagnostic Treatment and Cognitive Behavioral Therapy (CBT) on cognitive, behavioral and emotional components in patients with illness anxiety disorder comorbid with depression.

Design

This research compares two groups receiving unified transdiagnostic treatment and the group receiving cognitive-behavioral therapy. The sample includes 50 patients with an illness anxiety disorder who are selected as available and randomly assigned to intervention groups.

Settings and conduct

After the invitation was presented at Rasoul Akram Hospital in Tehran city, 50 eligible patients were randomly divided into two groups after explaining the treatment and obtaining informed consent. Then a pre-test and a post-test are conducted after the intervention and the subjects of both groups are followed up three months later. This study will be conducted in a double-blind manner, which means that the type of treatment will be unknown to the patients and the data analyst. Participants will also be unaware of which group they belong to.

Participants/Inclusion and exclusion criteria

Inclusion criteria: primary diagnosis of illness anxiety disorder, presence of a comorbid diagnosis of major depressive disorder (MDD) and/or persistent depressive disorder (PDD) based on DSM-5 criteria; Exclusion criteria: receiving psychological treatments in the last 3 years.

Intervention groups

This study has two intervention groups. In the first group, in order to check the effectiveness of the unified transdiagnostic treatment method, Barlow's new unified

transdiagnostic treatment protocol is used. Patients in the second group receive the treatment protocol called "cognitive-behavioral therapy for disease anxiety". Both protocols are 12 weekly sessions for 60-90 minutes.

Main outcome variables

Health anxiety symptoms and Beck depression symptoms

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20170104031765N2**

Registration date: **2023-11-14, 1402/08/23**

Registration timing: **prospective**

Last update: **2023-11-14, 1402/08/23**

Update count: **0**

Registration date

2023-11-14, 1402/08/23

Registrant information

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Name of organization / entity

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Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2023-11-22, 1402/09/01

Expected recruitment end date

2024-01-21, 1402/11/01

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Comparison of the Effectiveness of the Unified Protocol for Transdiagnostic Treatment (UP) and Cognitive Behavioral Therapy (CBT) on Emotional, Behavioral and Cognitive Components in Patients with Illness Anxiety Disorder Comorbid with Depression

Public title

Comparison of the Effectiveness of the Unified Protocol for Transdiagnostic Treatment and Cognitive Behavioral Therapy (CBT) in Improving Severe Health Anxiety

Purpose

Treatment

Inclusion/Exclusion criteria**Inclusion criteria:**

Primary diagnosis of illness anxiety disorder according to the Diagnostic and Statistical Manual of Mental Disorders 5 (DSM-5) Diagnosis of major depressive disorder (MDD), persistent depressive disorder (PDD) based on DSM-5 criteria Ages 18 to 65 years Able to read and write Not taking psychiatric drugs or in the case of taking antidepressants and anti-anxiety drugs for at least two months before the onset of interventions, the drug dose must be constant and the patient must agree to the drug dose remaining constant during the study

Exclusion criteria:

Patients with psychotic disorders Patients with bipolar disorder Patients with depressive disorder with psychotic symptoms Patients with a history of drug use in the last 6 months Patients at risk of suicide Personality disorder making the treatment procedure very difficult Patients with eating disorders Having a history of incurable physical illness such as cancer The need to change, stop or increase the drug dose or start a new drug after pre-test evaluations Receiving other psychological treatments in the past 5 years

Age

From **18 years** old to **65 years** old

Gender

Both

Phase

N/A

Groups that have been masked

- Participant
- Outcome assessor
- Data analyser

Sample size

Target sample size: **50**

Randomization (investigator's opinion)

Randomized

Randomization description

In this study, 50 people are selected according to similar research, in an accessible way. Then random replacement is done by lottery method (one number will be assigned to each sample person). We write down the numbers on paper, fold them and put them in a container. We mix and stir the papers completely. Then we take out the papers one by one and put one number in the first group and one number in the second group. Then random application will be done. In this way, one group is known as "cognitive behavioral therapy group" and the other group is known as "Unified transdiagnostic therapy group". The lottery will be done by putting the two groups obtained from the previous step (random replacement step) into two separate envelopes and placing them in a container. Then we randomly take one envelope out of the container and assign it to the "Unified transdiagnostic treatment group" and the other envelope to the "cognitive behavioral therapy group".

Blinding (investigator's opinion)

Double blinded

Blinding description

Our study will be conducted in a double-blind manner, which means that the type of treatment was unknown to the patients and the data analyst. Participants will not know the group type. Also, at first, they will be unaware of which group they belong to. The questionnaires will be completed under the supervision of an individual unaware of the research objectives.

Placebo

Not used

Assignment

Parallel

Other design features**Secondary Ids**

empty

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

Ethics committee of Iran University of Medical Sciences

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Iran University of Medical Sciences, Shahid Hemmat Highway

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1449614535

Approval date

2023-11-12, 1402/08/21

Ethics committee reference number

IR.IUMS.REC.1402.676

Health conditions studied

1

Description of health condition studied

illness anxiety disorder

ICD-10 code

F45.2

ICD-10 code description

Hypochondriacal disorders

Primary outcomes

1

Description

Health anxiety score in Salkowski questionnaire

Timepoint

Before and after interventions and three month follow-up

Method of measurement

Health Anxiety Questionnaire - Short Form (SV-HAI)

2

Description

Depression score in Beck questionnaire

Timepoint

Before and after interventions and three month follow-up

Method of measurement

Beck Depression Inventory - Second Edition (BDI-II)

Secondary outcomes

1

Description

The score of cognitions related to health anxiety

Timepoint

Before and after interventions and three month follow-up

Method of measurement

Health Cognition Questionnaire (HCQ)

2

Description

Anxiety Sensitivity Score

Timepoint

Before and after interventions and three month follow-up

Method of measurement

Anxiety sensitivity index-3 (ASI-3)

3

Description

Distress Tolerance Score

Timepoint

Before and after interventions and three month follow-up

Method of measurement

Distress Tolerance Scale (DTS)

4

Description

Emotion Regulation Score

Timepoint

Before and after interventions and three month follow-up

Method of measurement

Emotion regulation Questionnaire (ERQ)

5

Description

Beck anxiety score

Timepoint

Before and after interventions and three month follow-up

Method of measurement

Beck Anxiety Inventory (BAI)

6

Description

Cyberchondria Score

Timepoint

Before and after interventions and three month follow-up

Method of measurement

Short-Form- Cyberchondria Severity Scale (12-CSS)

7

Description

Score different aspects of illness behavior

Timepoint

Before and after interventions and three month follow-up

Method of measurement

The Scale for the Assessment of Illness Behaviour (SAIB)

8

Description

Interoceptive Attention Score

Timepoint

Before and after interventions and three month follow-up

Method of measurement

Interoceptive Attention Scale

9

Description

Amount of credibility/expectancy of improvement

Timepoint

After interventions

Method of measurement

Credibility/Expectancy Questionnaire (CEQ)

10

Description

The rate of therapeutic alliance

Timepoint

After interventions

Method of measurement

Working Alliance Inventory (WAI) For assessing therapeutic alliance

Intervention groups

1

Description

Intervention group: the people of the experimental group or integrated meta-diagnostic therapy group during 12 group sessions of approximately 60 to 90 minutes. All sections are performed on the subjects in a fixed order and with equal sessions. The sessions of this treatment protocol include the first session: increasing motivation; The second session: Providing psychological training; The third and fourth sessions: Emotional awareness training; The fifth session: cognitive reassessment; The sixth session: Identifying avoidance patterns; the Seventh session: Emotion-induced behaviors (EDBs); Eighth session: awareness and tolerance of physical feelings; 9th and 11th sessions: interoception and situational expouser and 12th session: relapse prevention. Treatment sessions will also be offered to clients by targeting treatment goals and focusing on different parts of the protocol.

Category

Treatment - Other

2

Description

Control group: Patients assigned to the control group receive the treatment protocol called "cognitive-behavioral therapy for illness anxiety". This protocol consists of 12 weekly sessions for 60-90 minutes, and was designed by Axelsson in 2020 based on the cognitive-behavioral model. The content of the cognitive-behavioral therapy sessions is as follows. The first session: Introduction to CBT and health anxiety. Second session: The CBT model of health anxiety. The third session: Interoceptive exposure. Fourth session: Response prevention. Fifth session: Exposure in vivo. The sixth session: Imaginal exposure. Seventh session: Continued imaginal exposure and the fear of death. Eighth session: Common obstacles to exposure. 9th and 10th session: Continued exposure and response prevention. The eleventh session: summary and values. Session 12: Continuous improvement and healthcare utilization.

Category

Treatment - Other

Recruitment centers

1

Recruitment center

Name of recruitment center

Hazrate Rasoole Akram Hospital

Full name of responsible person

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

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Grant name

Grant code / Reference number

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

Yes

Title of funding source

Iran 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

Iran University of Medical Sciences

Full name of responsible person

Hamid Mohsenabadi

Position

PhD student

Latest degree

Master

Other areas of specialty/work

Psychology

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

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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

Undecided - It is not yet known if there will be 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

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

Clinical Study Report

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

Analytic Code

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

Data Dictionary

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