

Clinical Trial Protocol

Iranian Registry of Clinical Trials

29 Jul 2026

Comparison of efficacy of internet- based cognitive behavior therapy alone versus combination CBT with well-being therapy for improvement of depressive symptoms of pregnant women

Protocol summary

Study aim

Determination of the effect of internet-based cognitive behavior therapy (iCBT) alone vs a combination of iCBT with Well being therapy (WBT) on depressive symptoms of pregnant women

Design

A clinical trial with randomized, four parallel groups on 408 patients. R software version 3,6,1 will be used for randomization

Settings and conduct

In a multicenter randomized clinical trial of equality type, 408 pregnant women with depressive symptoms are divided into four groups of psychotherapy models (iCBT with reminder sessions - iCBT without reminder sessions - iCBT combination with iWBT without reminder sessions - iCBT and iWBT combination therapy with reminder sessions). The content of four types of treatment is prepared online in 8 sessions. The validity of the content will be confirmed by the research team. Reminder groups will receive a reminder dose during pregnancy monthly until the end of pregnancy, and at postpartum every two months until the first year of birth. Outcomes of this study are Edinburgh Depression, Psychiatric Symptoms (BSI-18), Specific Pregnancy Stress (NuPDQ), Psychological well-being, Maternal-Child Postnatal Attachment, Ages and Stages Questionnaire (ASQ).

Participants/Inclusion and exclusion criteria

Inclusion: Gestational age less than 30 weeks, Edinburgh depression score higher than 10 for low-risk pregnancies and higher than 7 for high-risk pregnancies, access to computers and the Internet, Educational level higher than primary school, consent to enter treatment
Exclusion: Current substance abuse or dependence, currently using depressant drugs, recently receiving psychotherapy (<6 months), having a severe psychiatric, having suicidal ideations.

Intervention groups

Group 1: Only iCBT Group 2: Combined iCBT with WBT
Group 3: iCBT with reminder doses. Group 4: Combined iCBT and iWBT with reminder doses.

Main outcome variables

Symptoms of depression and anxiety

General information

Reason for update

Acronym

iCBT: Internet-based Cognitive behavior therapy iWBT: Internet-base well-being treatment

IRCT registration information

IRCT registration number: **IRCT20110228005931N12**

Registration date: **2022-06-05, 1401/03/15**

Registration timing: **prospective**

Last update: **2022-06-05, 1401/03/15**

Update count: **0**

Registration date

2022-06-05, 1401/03/15

Registrant information

Name

Mahbobeh Faramarzi

Name of organization / entity

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Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2022-08-23, 1401/06/01
Expected recruitment end date
 2026-03-21, 1405/01/01
Actual recruitment start date
 empty
Actual recruitment end date
 empty
Trial completion date
 empty

Scientific title
 Comparison of efficacy of internet- based cognitive behavior therapy alone versus combination CBT with well-being therapy for improvement of depressive symptoms of pregnant women

Public title
 Effect of Internet cognitive-behavioral psychotherapy with psychological well-being on depression of pregnant women

Purpose
 Treatment

Inclusion/Exclusion criteria
Inclusion criteria:
 Gestational age of less than 30 weeks Edinburgh depression score higher than 10 for low-risk pregnancies and higher than 7 for high-risk pregnancies Access to computers and the Internet Educational level higher than primary school Consent to enter treatment
Exclusion criteria:
 Current substance abuse or dependence Current use of antidepressant drugs, or recent psychotherapy for depression/anxiety (<4 weeks) Women with severe depression disorders who require medical treatment Severe psychiatric disorders or suicidal ideations Childbirth before the end of the treatment period

Age
 From **18 years** old to **45 years** old

Gender
 Female

Phase
 N/A

Groups that have been masked
No information

Sample size
 Target sample size: **408**

Randomization (investigator's opinion)
 Randomized

Randomization description
 After checking inclusion and exclusion criteria, participants are allocated randomly into four groups by permuted block randomization method. The block size is 4 and each block of each intervention group will be repeated once. Then, will be generated 102 blocks with 4 sizes and a total of 408 sequences using the statistical program in R software environment version 3,6,1. The participants will be assigned to four groups of the study by a randomly generated list.

Blinding (investigator's opinion)
 Not blinded

Blinding description
Placebo

Not used
Assignment
 Parallel
Other design features

Secondary Ids

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Ethic Committee of Babol University of Medical Sciences

Street address

Babol University of Medical Sciences, Ganjafroz Avenue, Babol, Mazandaran, Babol, Iran

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Postal code

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Approval date

2021-10-06, 1400/07/14

Ethics committee reference number

IR.MUBABOL.HRI.REC.1400.084

Health conditions studied

1

Description of health condition studied

Depressive Disorder

ICD-10 code

F32.9

ICD-10 code description

Depression NOS

Primary outcomes

1

Description

Depression

Timepoint

Before intervention, immediately after intervention, follow-ups 6 and 12 months after intervention

Method of measurement

Edinburgh Postnatal Depression Inventory

2

Description

Anxiety

Timepoint

Before intervention, immediately after intervention, follow-ups 6 and 12 months after intervention

Method of measurement

Secondary outcomes

1

Description

Pregnancy stress

Timepoint

Before the intervention, at end of the intervention, and follow-ups of 1 and 6 months after the end of intervention

Method of measurement

Revised Prenatal Distress Questionnaire (NuPDQ)

2

Description

Mother and child attachment

Timepoint

After the birth of a child at 4 and 12 months

Method of measurement

Mother-child attachment questionnaire

3

Description

Child development

Timepoint

After the birth of a child at 4, and 12 months

Method of measurement

Ages and Stages Questionnaire (ASQ)

4

Description

Psychological well-being

Timepoint

Before the intervention, at end of the intervention, and follow-ups of 1 and 6 months after the end of intervention

Method of measurement

Ryff psychological well-being questionnaire

Intervention groups

1

Description

Intervention group 1: Participants in this group receive online cognitive behavioral therapy in 8 sessions of 90 minutes, once a week.

Category

Treatment - Other

2

Description

Intervention group 2: Participants in this group receive combined online cognitive-behavioral therapy with psychological well-being in 8 sessions of 90 minutes, once a week.

Category

Treatment - Other

3

Description

Intervention group 3: Participants in this group receive Internet cognitive behavioral therapy in 8 sessions of 90 minutes, once a week, with reminder sessions.

Category

Treatment - Other

4

Description

Fourth intervention group: Participants in this group receive combined online cognitive-behavioral therapy with psychological well-being in 8 sessions of 90 minutes, once a week, with reminder sessions.

Category

Treatment - Other

Recruitment centers

1

Recruitment center

Name of recruitment center

Rouhani and Yahya Nejad Educational and Medical Center and private offices in Babol

Full name of responsible person

D.zahra basirat

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

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Full name of responsible person

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Grant name
Research Project

Grant code / Reference number
724133475

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

Title of funding source
Babol 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
Babol University of Medical Sciences

Full name of responsible person
Mahbobeh Faramarzi

Position
Associate professor

Latest degree
Ph.D.

Other areas of specialty/work
Psychology

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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