

Clinical Trial Protocol

Iranian Registry of Clinical Trials

09 Jun 2026

The effectiveness of recovery-oriented cognitive therapy with and without cognitive remediation in individuals with schizophrenia disorder.

Protocol summary

Study aim

The effectiveness of recovery-oriented cognitive therapy with and without cognitive remediation in individuals with schizophrenia disorder.

Design

A concealed, randomized, single-blinded controlled clinical trial with a parallel-group design

Settings and conduct

the participants from Razi hospital will complete the basic assessment such as the Schizophrenic Negative and Positive Symptoms Scale, attitude Scale, Neuropsychologic assessment, Quality of Life Mental Health Recovery, Hope Scale. then Intervention groups A will undergo recovery-oriented cognitive therapy and Intervention B groups will also be treated recovery-oriented cognitive therapy with cognitive rehabilitation after intervention phase post-treatment assessment will conduct.

Participants/Inclusion and exclusion criteria

1. 20-60 years 2. most of the symptoms are negative. 4. Ability to reading and writing Exclusion: 1. Brain damage, mental retardation, a physical disability 2. Neurological disease 3. seriously affected by the side effects of psychiatric medications. 4. Dissatisfaction with participating in the study. 5. The patient is in the acute phase of the disease. 6. Receive electroshock at least 6 months before the study and during treatment. 7. Be currently undergoing other psychological interventions. 8. The patient is severely affected by abuse, poisoning, or deprivation.

Intervention groups

The intervention of recovery-oriented cognitive therapy with and without cognitive rehabilitation (comparison group as control) for patients with schizophrenia and to improve negative and positive symptoms, dysfunctional beliefs, neuro-cognitive, disability, quality of life, improvement of mental health, recovery, hope will be conducted.

Main outcome variables

Negative and positive symptoms, dysfunctional beliefs, neurocognitive, quality of life, improved mental health, recovery, hope.

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20200701047974N1**

Registration date: **2020-07-02, 1399/04/12**

Registration timing: **prospective**

Last update: **2020-07-02, 1399/04/12**

Update count: **0**

Registration date

2020-07-02, 1399/04/12

Registrant information

Name

Ali Ebrahimi

Name of organization / entity

Country

Iran (Islamic Republic of)

Phone

+98 21 3307 7230

Email address

alipsychologist69@gmail.com

Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2020-07-22, 1399/05/01

Expected recruitment end date

2020-10-22, 1399/08/01

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date
empty

Scientific title
The effectiveness of recovery-oriented cognitive therapy with and without cognitive remediation in individuals with schizophrenia disorder.

Public title
The effectiveness of recovery-oriented cognitive therapy with and without cognitive remediation in individuals with schizophrenia disorder.

Purpose
Education/Guidance

Inclusion/Exclusion criteria
Inclusion criteria:
20-60 years Diagnosis of schizophrenia based on DSM-5 criteria At least two years have passed since the onset of schizophrenia and most of the symptoms are negative. The score of positive and negative symptoms of the patient in SAPS tests (positive symptom assessment scale) and SANS (negative symptom evaluation scale) should be above the cutting point of 35 and 24, respectively. Ability to reading and writing
Exclusion criteria:
Brain damage, mental retardation, physical disability that interferes with the assessment process. Neurological disease (epilepsy, Alzheimer's, dementia, Parkinson's, MS, etc.) due to which the person being under treated The patient is seriously affected by the side effects of psychiatric medications that interfere with the study process. Dissatisfaction with participating in the study The patient is in the acute phase of the disease Receive an electroshock at least 6 months before the study and during treatment Be currently undergoing other psychological interventions The patient is severely affected by abuse, poisoning or deprivation of any kind of drug (smoking is not considered a exclusion criterion).

Age
From **20 years** old to **60 years** old

Gender
Both

Phase
N/A

Groups that have been masked

- Outcome assessor

Sample size
Target sample size: **30**

Randomization (investigator's opinion)
Randomized

Randomization description
In the present study, adaptive randomization methods used based on the variables of the aid of the thesis that provide decisions based on the division of individuals into two groups of interventions on a case-by-case basis. (Woman) Age (in four age groups: 30-20, 40-31, 50-41, 60-51), and severity of negative and positive symptoms (in 5 levels: (1) suspicious, (2) mild, (3) Medium, (4) obvious, (5) severe) are important variables in this study. The characteristics of the first patient in the mentioned

variables are recorded. the next individuals are assigned to the groups in such a way that the difference between the groups is minimal in terms of these important variables, and thus the imbalance can be minimized.

Blinding (investigator's opinion)
Single blinded

Blinding description
Those assessing outcomes and data Analyzer not being able to check the patient status and blinded to the research condition so this study will be a single-blind study.

Placebo
Not used

Assignment
Parallel

Other design features

Secondary Ids
empty

Ethics committees
1
Ethics committee
Name of ethics committee
Ethics committee of University of Social Welfare and Rehabilitation Sciences
Street address
kodakyar Ave., daneshjo Blvd.,Evin
City
Tehran
Province
Tehran
Postal code
1985713871
Approval date
2020-06-22, 1399/04/02
Ethics committee reference number
IR.USWR.REC.1399.103

Health conditions studied
1
Description of health condition studied
schizophrenia disorder
ICD-10 code
F20
ICD-10 code description
Schizophrenia

Primary outcomes
1
Description
Negative Symptoms
Timepoint
Per-intervention, mid intervention and post-intervention and follow up

Method of measurement

Scale for Assessment of Negative Symptoms (SANS)

2

Description

Positive Symptoms

Timepoint

Per-intervention, mid intervention and post-intervention and follow up

Method of measurement

Scale for Assessment of Positive Symptoms (SAPS)

3

Description

Neurocognitive impairment

Timepoint

Per-intervention, mid intervention and post-intervention and follow up

Method of measurement

MATRICES

4

Description

Dysfunctional Attitude

Timepoint

Per-intervention, mid intervention and post-intervention and follow up

Method of measurement

Dysfunctional Attitude Scale (DAS)

5

Description

Real-World Functioning

Timepoint

Per-intervention, mid intervention and post-intervention and follow up

Method of measurement

WHO DAS .2

6

Description

Quality of life

Timepoint

Per-intervention, mid intervention and post-intervention and follow up

Method of measurement

Self-report quality of life measure for people with schizophrenia

7

Description

Mental Health Recovery

Timepoint

Per-intervention, mid intervention and post-intervention and follow up

Method of measurement

Mental Health Recovery Measure

8

Description

Recovery

Timepoint

Per-intervention, mid intervention and post-intervention and follow up

Method of measurement

Recovery Self-Assessment-Revised (RSA-R)

9

Description

Hope

Timepoint

Per-intervention, mid intervention and post-intervention and follow up

Method of measurement

The Schizophrenia Hope Scale

Secondary outcomes

empty

Intervention groups

1

Description

Intervention group1: Intervention groups A will undergo recovery oriented cognitive therapy (CT-R). The program is designed to run individually or in groups, and will run for 14 sessions, averaging 30 minutes and 2 sessions per week.

Category

Treatment - Other

2

Description

Intervention group2: Intervention B groups will also be treated for 14 minutes, an average of 30 minutes, and 2 sessions per week with recovery oriented cognitive therapy (CT-R) with cognitive rehabilitation (CT-R + CR).

Category

Treatment - Other

Recruitment centers

1

Recruitment center

Name of recruitment center

Razi psychiatric hospital

Full name of responsible person

Ali Ebrahimi

Street address

kodakyar Ave., daneshjo Blvd.,Evin,

City

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1985713871

Phone

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Email

Alipsychologist69@gmail.com

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

University of social welfare and rehabilitation sciences

Full name of responsible person

Ali Ebrahimi

Street address

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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

University of social welfare and rehabilitation sciences

Proportion provided by this source

10

Public or private sector

Private

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

University of social welfare and rehabilitation sciences

Full name of responsible person

Ali Ebrahimi

Position

PhD candidate

Latest degree

Ph.D.

Other areas of specialty/work

Psychology

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Person responsible for scientific inquiries

Contact

Name of organization / entity

University of social welfare and rehabilitation sciences

Full name of responsible person

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Position

PhD candidate

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

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Position

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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