

Clinical Trial Protocol

Iranian Registry of Clinical Trials

02 Jul 2026

Comparison of therapeutic effectiveness based on the analysis of lived experiences of social stigma with positive psychotherapy method in mental toughness, cognitive flexibility, distress tolerance and feeling of entrapment in patients with multiple sclerosis.

Protocol summary

Study aim

Reducing the symptoms of multiple sclerosis and complementary therapies

Design

This clinical trial with a quasi-experimental design with control group on 45 patients used coin tossing method by an independent person to randomize the groups.

Settings and conduct

The method and place of study of this clinical trial was conducted in the Darya Psychology Clinic, under the supervision of the research department of the Isfahan faculty of Psychology. By calling through WhatsApp and also the clients of the supervisor professor, 45 eligible patients with MS who wished to participate were registered. Participants in two experimental groups of 15, underwent a 90 minutes session for 8 weeks as a group undergoing positive psychotherapy and treatment based on lived experiences of social stigma in patients with MS, while the control group which included 15 people, did not receive any therapy. In order to evaluate the effects of therapy, subjects were assessed before and after the therapy and one month after the intervention, respectively.

Participants/Inclusion and exclusion criteria

Inclusion criteria: Diagnosis of MS by a specialist. Willingness and signing of written consent. Form of consent to participate in the research project. Not suffering from a severe physical illness, severe neurological disorders, or having symptoms of psychosis. Exclusion criteria: relapse of illness and inability to continue sessions absence of more than two sessions

Intervention groups

The intervention group includes patients with multiple sclerosis who have received a positive psychotherapy intervention. The intervention group includes patients with multiple sclerosis who have received treatment

intervention based on lived experience.

Main outcome variables

mental toughness, cognitive flexibility, distress tolerance, feeling of entrapment

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20210808052113N1**

Registration date: **2021-09-17, 1400/06/26**

Registration timing: **retrospective**

Last update: **2021-09-17, 1400/06/26**

Update count: **0**

Registration date

2021-09-17, 1400/06/26

Registrant information

Name

Yaghoob Harooni Jamalooei

Name of organization / entity

University of Isfahan

Country

Iran (Islamic Republic of)

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Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2021-01-19, 1399/10/30

Expected recruitment end date

2021-01-29, 1399/11/10

Actual recruitment start date

2021-01-19, 1399/10/30

Actual recruitment end date

2021-01-29, 1399/11/10

Trial completion date

2021-06-10, 1400/03/20

Scientific title

Comparison of therapeutic effectiveness based on the analysis of lived experiences of social stigma with positive psychotherapy method in mental toughness, cognitive flexibility, distress tolerance and feeling of entrapment in patients with multiple sclerosis.

Public title

Comparison of therapeutic effectiveness based on the analysis of lived experiences of social stigma with positive psychotherapy method in mental toughness, cognitive flexibility, distress tolerance and feeling of entrapment in patients with multiple sclerosis.

Purpose

Treatment

Inclusion/Exclusion criteria**Inclusion criteria:**

Diagnosis of MS by a specialist. Age range between 18 and 55 years. Willingness and signing of written consent. Form of consent to participate in the research project. Not suffering from a severe physical illness, severe neurological disorders, or having symptoms of psychosis.

Exclusion criteria:

relapse of illness and inability to continue sessions
absence of more than two sessions

Age

From **18 years** old to **55 years** old

Gender

Both

Phase

N/A

Groups that have been masked

No information

Sample size

Target sample size: **45**

Actual sample size reached: **45**

Randomization (investigator's opinion)

Not randomized

Randomization description**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**

University of Isfahan

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Isfahan university, Daneshgah st

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Postal code

8174673441

Approval date

2021-01-12, 1399/10/23

Ethics committee reference number

IR.UI.REC.1399.082

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

Multiple Sclerosis

ICD-10 code

G35

ICD-10 code description

Multiple sclerosis

Primary outcomes**1****Description**

mental toughness

Timepoint

Before and after the therapy

Method of measurement

Mental toughness survey

Secondary outcomes**1****Description**

Cognitive flexibility

Timepoint

Before and after the therapy

Method of measurement

Cognitive flexibility survey(CFI)

2**Description**

Mental toughness

Timepoint

Before and after the therapy

Method of measurement

Mental toughness survey(MTQ)

3

Description

Distress tolerance

Timepoint

Before and after the therapy

Method of measurement

Distress tolerance survey(DTF)

4

Description

feeling entrapment

Timepoint

Before and after the therapy

Method of measurement

feeling entrapment survey(Gilbert and Allen)

Intervention groups

1

Description

The method of intervention is phenomenology of lived experiences in the field of qualitative research. The contents of the treatment session based on the stigmatized life experiences of MS patients to explain and effectiveness of treatment, after the information was done through in-depth interviews with patients and interpretation of data by Colaizzi method, analysis and coding and categorization during data collection and Data collection and analysis continued until theoretical saturation within the scope of the research. Then, the extracted factors were presented and developed to develop a treatment model based on the analysis of lived experiences of stigma. It was held in eight one-and-a-half hour sessions during a week.

Category

Behavior

2

Description

Content of Positive Psychotherapy Sessions has been done in eight consecutive sessions. The sessions included recording a positive self-introduction on a page and directing clients in a positive psychotherapy framework, recording each person's ability by themselves, mentioning blessings and repeating tasks which is adapted from Seligman(2006). In the review session, whether writing these three good things or three blessings and emphasizing positive memories and recollections over the past week has had a positive effect or not, using the worksheet related to gratitude, reviewing the relation of treatment of clients with their progress in writing booklets and letters of forgiveness and gratitude and using their capabilities in practice based on their activity plans which began in the second session were examined. Focusing on the topics of hope, faith and optimism, using the art of improving relationships and creating positive social relationships and happiness in life, teaching response style and training to improve relationships and completing

questionnaires by participants in eight sessions of one and a half hours per week were done.

Category

Behavior

3

Description

Control group: includes 15 people who are in the waiting list group and do not receive therapy at the time of intervention. But at the end of the therapy, for the sake of ethics, the control group can also use positive psychotherapy and lived-experience therapy if needed.

Category

N/A

Recruitment centers

1

Recruitment center

Name of recruitment center

University of Isfahan

Full name of responsible person

Yaghoob Harooni jamalooei

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Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Deputy of Research, University of Isfahan

Full name of responsible person

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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

Deputy of Research, University of Isfahan

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

University of Isfahan

Full name of responsible person

Yaghoob Harooni Jamalooei

Position

Student

Latest degree

Ph.D.

Other areas of specialty/work

Psychology

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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

Not applicable

Data Dictionary

Not applicable

Title and more details about the data/document

The group data is related to the main messages of the interview without mentioning the demographic characteristics of the buyers

When the data will become available and for how long

Five months after printing

To whom data/document is available

Academic researchers after authentication

Under which criteria data/document could be used

Request to use the data is permitted only for use in cross-cultural studies, multiple sclerosis treatment centers, or meta-combination and meta-analytic studies.

From where data/document is obtainable

send a request to email address
yaghoob.harooni2014@gmail.com

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

Data information will be provided to individuals via email or WhatsApp after evaluating the request.

Comments