

Clinical Trial Protocol

Iranian Registry of Clinical Trials

02 Jul 2026

The Effect of Fluxetine on Disease Clinical & Radiologic activity of Multiple sclerosis

Protocol summary

Summary

The aim of the present study is to determine the anti-inflammatory effects of fluoxetine on clinical course and new lesion formation in patients with relapsing MS. In a triple-blind, placebo-controlled exploratory study, 58 nondepressed patients with relapsing remitting MS will be randomised to oral fluoxetine 20 mg or placebo daily for 24 weeks. Clinical and Radiologic activity of MS will be studied by assessing EDSS (Expanded Disability Status Scale) and the cumulative number of lesions on different phases of brain MRI performed on weeks 1 and 24

General information

Acronym

IRCT registration information

IRCT registration number: **IRCT2014052417811N2**
Registration date: **2014-06-01, 1393/03/11**
Registration timing: **retrospective**

Last update:

Update count: **0**

Registration date

2014-06-01, 1393/03/11

Registrant information

Name

Shayan Amjadi

Name of organization / entity

Students Research Committee, Arak University of Medical Science

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Recruitment status

Recruitment complete

Funding source

Arak university of medical science

Expected recruitment start date

2013-07-02, 1392/04/11

Expected recruitment end date

2014-06-01, 1393/03/11

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

The Effect of Fluxetine on Disease Clinical & Radiologic activity of Multiple sclerosis

Public title

Fluxetine AND Multiple Sclerosis

Purpose

Treatment

Inclusion/Exclusion criteria

inclusion criteria: age 18–50 years; clinically definitive relapsing remitting MS (according to the 2005 revised McDonald criteria) confirmed by a neurologist; Additional inclusion criteria were an Expanded disability Status Score (EDSS) of less or equal to 5.5; exclusion criteria: progressive forms of multiple sclerosis; an onset of relapse or receipt of any glucocorticoid treatment during trial period; use of immunomodulatory, immunosuppressive, IVIg or plasmapheresis or antidepressant drugs in the previous 6 months; the use of corticosteroids in the previous 8 weeks; depression defined as a score of 19 or higher on Beck's Depression Inventory II; contraindication to MRI; other neurological or systemic disorder that would interfere with the assessments; pregnancy or unwillingness to use acceptable birth control. severe suicidality including

ideation, plan, and intentclinically significant : unstable medical or surgical conditions that would preclude safe and complete participation in the study.

Age

From **18 years** old to **50 years** old

Gender

Both

Phase

3

Groups that have been masked

No information

Sample size

Target sample size: **58**

Randomization (investigator's opinion)

Randomized

Randomization description

Blinding (investigator's opinion)

Triple blinded

Blinding description

Placebo

Used

Assignment

Parallel

Other design features

Secondary Ids

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Arak University of medical science

Street address

Arak University of medical science,sardasht,arak,iran

City

arak

Postal code

Approval date

2013-02-02, 1391/11/14

Ethics committee reference number

92-143-10

Health conditions studied

1

Description of health condition studied

multiple sclerosis

ICD-10 code

G35

ICD-10 code description

Demyelinating diseases of the central nervous system

Primary outcomes

1

Description

EDSS (Expanded Disability Status Scale)

Timepoint

WEEK 0 AND24

Method of measurement

evaluate by neurologist

2

Description

the cumulative number of lesions on different phases of brain MRI

Timepoint

WEEK 0 AND24

Method of measurement

counting the plaques by researchers

Secondary outcomes

empty

Intervention groups

1

Description

oral flouxetine 20 mg for 24 weeks will be given to patient whom randomised to intervention group

Category

Treatment - Drugs

2

Description

patients whom randomized to placebo group will received placebo capsules during 24 weeks of clinical trial course

Category

Treatment - Drugs

Recruitment centers

1

Recruitment center

Name of recruitment center

Neurology clinic of valiasr hospital-Arak MS society-private offices

Full name of responsible person

Dr.Fardin Faraji

Street address

City

ARAK

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Arak University of Medical Science,Office of Vice
Chancellor for Research

Full name of responsible person

Davoud Hekmatpo

Street address

Arak university of medical science , Arak , Iran

City

Arak

Grant name

22100000 جلا

Grant code / Reference number

907

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

Yes

Title of funding source

Arak University of Medical Science,Office of Vice
Chancellor for Research

Proportion provided by this source

100

Public or private sector

empty

Domestic or foreign origin

empty

Category of foreign source of funding

empty

Country of origin

Type of organization providing the funding

empty

Person responsible for general inquiries

Contact

Name of organization / entity

Students Research Committe , Medicine faculty ,
Aarak University of Medical Science

Full name of responsible person

Shayan Amjadi

Position

Medical student

Other areas of specialty/work

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Person responsible for scientific inquiries

Contact

Name of organization / entity

Neurology Department , Medicine faculty, Arak
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Full name of responsible person

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Sharing plan

Deidentified Individual Participant Data Set (IPD)

empty

Study Protocol

empty

Statistical Analysis Plan

empty

Informed Consent Form

empty

Clinical Study Report

empty

Analytic Code

empty

Data Dictionary

empty