

Clinical Trial Protocol

Iranian Registry of Clinical Trials

04 Aug 2026

The evaluation of the efficacy and safety of oral fingolimod in relapsing remitting multiple sclerosis

Protocol summary

Summary

In order to investigate efficacy and safety of oral fingolimod in relapsing-remitting multiple sclerosis (RRMS), a clinical trial will be done for 3 months. Participants will be recruited from RRMS patients living in Kermanshah province, referred to Emam Raza hospital MS clinic in Kermanshah, who are eligible to the study inclusion and exclusion criteria, until study sample size of 60 cases is reached. After describing study goals to the patients and providing written informed consent, patients will be admitted to hospital for at least 6 hours to receive first dose oral fingolimod (0.5 mg fingolid ® capsules, Osve pharmaceuticals) to rule out probable bradycardia and heart block. During admission heart rate, blood pressure and electrocardiogram will be monitored. Afterward patients will receive fingolimod capsules at home, once daily orally at least for 12 months. Patients will be visited at 2 weeks, 1 month, 2 month, 3 month, 9 month and 12 month after intervention. In each visit patients will be investigated for MS relapse, drug complication and safety and heart rate and blood pressure will be recorded. In order to investigate disease progression, expanded disability scale score (EDSS) will be recorded every 3 months.

General information

Acronym

IRCT registration information

IRCT registration number: **IRCT201406018323N10**
Registration date: **2015-01-03, 1393/10/13**
Registration timing: **registered_while_recruiting**

Last update:

Update count: **0**

Registration date

2015-01-03, 1393/10/13

Registrant information

Name

Nazanin Razazian

Name of organization / entity

Neurology Department of Kermanshah Faculty of Medicine

Country

Iran (Islamic Republic of)

Phone

+98 83 1427 3409

Email address

nrزازian@kums.ac.ir

Recruitment status

Recruitment complete

Funding source

kermanshah university of medicine

Expected recruitment start date

2014-12-22, 1393/10/01

Expected recruitment end date

2016-02-20, 1394/12/01

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

The evaluation of the efficacy and safety of oral fingolimod in relapsing remitting multiple sclerosis

Public title

The evaluation of the efficacy and safety of oral fingolimod in relapsing remitting multiple sclerosis

Purpose

Treatment

Inclusion/Exclusion criteria

Inclusion criteria: 1. Patients with Relapsing- Remitting Multiple Sclerosis (RRMS) based on McDonald's Criteria (2010) 2. age between 18 to 45 years 3. EDSS between 0

to 5.5 4. at least one relapse in previous year while receiving first line disease modifying therapy 5.at least 2 relapses in previous 2 years while receiving first line disease modifying therapy 6. no relapses in 30 days previous to fingolimod initiation 7. negative pregnancy tests in child bearing women 8. positive history of varicella zoster virus vaccination or positive anti varicella zoster antibody in serum Exclusion criteria: 1. primary or secondary progressive MS 2. history of chronic systemic disease other than MS (lung, liver, kidney,...disease) 3. history of or active malignancy 4. presence of macular edema at baseline 5. acute or chronic active bacterial, viral or fungal infection 6. treatment with IV corticosteroids during 30 days previous to fingolimod initiation 7. history of treatment with cyclophosphamide or mitoxantrone at any time 8. history of treatment with immunoglobulins or monoclonal antibodies in previous 6 months 9. history of treatment with methotrexate or azathioprine in previous 6 months 10. pregnancy or lactation 11. inoculation of live attenuated vaccines in previous 2 months 12. history of unstable angina, MI, TIA or CVA in previous 6 months 13. history of hospitalization due to decompensated heart failure in previous 6 months 14. Canada heart association class 3 or 4 heart failure 15. history or presence of second degree mobitz type AV block, third degree AV block or sick sinus syndrome, except having functional cardiac pace maker 16. QT interval more than 500 millisecond on baseline EKG 17. treatment with antiarrhythmic drugs class 1a or class 3 18. AST or ALT two folds normal upper limit 19. ALP 1.5 folds normal upper limit 20. direct or total Billi more than normal upper limit 21. creatinin more than 1.7 mg/dl 22. WBC less than 3500/ mm3 23. lymphocyte count less than 800/ mm3 24. presence of uveitis

Age

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

Gender

Both

Phase

4

Groups that have been masked

No information

Sample size

Target sample size: **60**

Randomization (investigator's opinion)

N/A

Randomization description

Blinding (investigator's opinion)

Not blinded

Blinding description

Placebo

Not used

Assignment

Single

Other design features

Secondary Ids

empty

Ethics committees

1

Ethics committee

Name of ethics committee

Kermanshah University of Medical Sciences

Street address

Kermanshah University of Medical Sciences, Shahid Beheshti boulder, Kermanshah

City

Kermanshah

Postal code

Approval date

2014-11-11, 1393/08/20

Ethics committee reference number

35220

Health conditions studied

1

Description of health condition studied

multiple sclerosis

ICD-10 code

G35

ICD-10 code description

Disseminated Multiple Sclerosis

Primary outcomes

1

Description

expanded disability score scale,EDSS

Timepoint

before intervention and every 3 months after intervention

Method of measurement

EDSS questionair

Secondary outcomes

1

Description

annual relapse rate

Timepoint

before intervention and monthly after intervention

Method of measurement

physical examination by neurologist and EDSS questionair

2

Description

disability progression

Timepoint

before intervention and monthly after intervention

Method of measurement

one point increment in EDSS that remains for 3 months,

without new relapse at the time of evaluation

3

Description

drug side effects

Timepoint

before intervention and monthly after intervention

Method of measurement

patient reported complications and designed drug side effect questionair

Intervention groups

1

Description

fingolimod capsule, chemical formulation FTY -720 , (0.5 mg/ day, oral , every day) for 12 months ,made in Osvah Pharmaceutical Company in Islamic Republic Of Iran, generic name Cap. Fingolide®

Category

Treatment - Drugs

Recruitment centers

1

Recruitment center

Name of recruitment center

Emam Reza Hospital of Kermanshah

Full name of responsible person

Dr. Nazanin Razazian

Street address

Emam Reza Hospital, Parastar bulvard, Sorkhelijeh, Kermanshah

City

Kermanshah

Sponsors / Funding sources

1

Sponsor

Name of organization / entity

Vice-chancellor for research, Kermanshah University of Medical Sciences

Full name of responsible person

Mr. Shirzadian

Street address

Kermanshah University of Medical Sciences, building no.2, Shahid Beheshti boulder, Kermanshah

City

Kermanshah

Grant name

Grant code / Reference number

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

Yes

Title of funding source

Vice-chancellor for research, Kermanshah University of Medical Sciences

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

Kermanshah University of Medical Sciences

Full name of responsible person

Dr. Nazanin Razazian

Position

neurologist/ Associate Professor

Other areas of specialty/work

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Person responsible for updating data

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Full name of responsible person

Dr. Elham Ouspid

Position

MD, resident of Neurology

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Web page address**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