

Study registered at the country of origin: Specify

Date of registration in national regulatory agency

An Open-label Study Evaluating Ofatumumab Treatment Effectiveness and PROs in Subjects With RMS Transitioning From Dimethyl Fumarate or Fingolimod to Ofatumumab

Protocol number

COMB157G23101

N/A

Acronym

Acronym

Type of registration: Justify

Primary sponsor: Country of origin

Novartis Pharmaceuticals

04/07/2025 17:19:40

Main Information

Primary registry identifying number

LBCTR2021034775

MOH registration number

Study registered at the country of origin

Type of registration

Prospective

Date of registration in national regulatory agency

Primary sponsor

Novartis Pharma Services Inc.

Date of registration in primary registry

23/05/2021

Public title

An Open-label Study Evaluating Ofatumumab Treatment Effectiveness and PROs in Subjects With RMS Transitioning From Dimethyl Fumarate or Fingolimod to Ofatumumab

Scientific title

A Single-arm, Prospective, Multicentre, Open-label Study to Evaluate Ofatumumab Treatment Effectiveness and Patient Reported Outcomes in Patients With Relapsing Multiple Sclerosis Transitioning From Dimethyl Fumarate or Fingolimod Therapy

Brief summary of the study: English

The open label study to evaluate effectiveness of treatment with ofatumumab in patients transitioning from commonly used oral MS therapies - fingolimod or dimethyl fumarate, due to breakthrough disease.

Brief summary of the study: Arabic

دراسة تقدّميّة متعددة المراكز مفقوحة اللصاقة ذات مجموعة واحدة لتقييم فعاليّة العلاج بأوفاتوموماب والنتائج التي يفيد عنها المرضى المصابين بالتصلب اللويحي الانتكاسي الذين ينتقلون من العلاج بثنائي ميثيل الفومارات أو بفينغوليمود

Health conditions/problem studied: Specify

Relapsing Multiple Sclerosis

Interventions: Specify

Biological: Ofatumumab

Patients in the ofatumumab will receive injections of ofatumumab provided in an autoinjector (AI) for subcutaneous administration containing 20

mg ofatumumab (50 mg/ml, 0.4 ml content)

Other Name: OMB157



Key inclusion and exclusion criteria: Inclusion criteria

Inclusion Criteria:

Diagnosis of multiple sclerosis (MS) Relapsing MS (RRMS or SPMS) course Subject transitioning from either fingolimod or dimethyl fumarate, following min 6 months treatment with either drug Breakthrough disease as evidence by clinical relapses or MRI EDSS score of 0 to 4

Key inclusion and exclusion criteria: Gender Key inclusion and exclusion criteria: Specify gender

Key inclusion and exclusion criteria: Age minimum Key inclusion and exclusion criteria: Age maximum

Key inclusion and exclusion criteria: Exclusion criteria

Primary progressive MS or SPMS without disease activity

Disease duration of more than 10 years since diagnosis

Patients with an active chronic disease of the immune system other than MS

Patients at risk of developing or having reactivation of hepatitis

Patients with active systemic infections or with neurological findings consistent with PML Other protocol-defined inclusion/exclusion criteria may

N/A

N/A

N/A

Type of study

Interventional

Treatment

Type of intervention Type of intervention: Specify type

Pharmaceutical

Trial scope Trial scope: Specify scope

Therapy

Study design: Allocation Study design: Masking

Single Arm Study Open (masking not used)

Study design: Control Study phase N/A

Study design: Purpose Study design: Specify purpose

Study design: Assignment Study design: Specify assignment

IMP has market authorization IMP has market authorization: Specify

Yes, Worldwide US, UAE, Albania, Argentina, Canada, Singapore, Switzerland

Name of IMP Year of authorization Month of authorization

Ofatumumab

Pharmaceutical class Monoclonal antibodies

Type of IMP Immunological

Therapeutic indication

Patients with: relapsing multiple sclerosis



Therapeutic benefit

potential efficacy of ofatumumab in patients with relapsing MS.

Study model

N/A

Study model: Specify model

N/A

Time perspective

N/A

Time perspective: Specify perspective

Target follow-up duration

Number of groups/cohorts

Biospecimen retention

Samples without DNA

Target sample size

10

Date of first enrollment: Type

Anticipated

Date of study closure: Type

Anticipated

Recruitment status

Pending

Date of completion

31/08/2022

IPD sharing statement plan

Yes

Study model: Explain model

N/A

Time perspective: Explain time perspective

N/A

Target follow-up duration: Unit

Biospecimen description

Covance Central lab: Ambient and Frozen conditions

Actual enrollment target size

Date of first enrollment: Date

29/04/2021

Date of study closure: Date

25/06/2025

Recruitment status: Specify

IPD sharing statement description

Novartis is committed to sharing with qualified external researchers, access to patient-level data and supporting clinical documents from eligible studies. These requests are reviewed and approved by an independent review panel on the basis of scientific merit. All data provided is anonymized to respect the privacy of patients who have participated in the trial in line with applicable laws and regulations.

This trial data availability is according to the criteria and process described on www.clinicalstudydatarequest.com

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Additional data URL

https://clinicaltrials.gov/ct2/show/record/NCT04353492? term=ofatumumab+treatment+effectiveness&cond=relapsing+multiple+sclerosis+transitioning&draw=2&rank=1

Admin comments

Trial status

Approved

| Secondary Identifying Numbers | | |
|--------------------------------|------------------------------|--|
| Full name of issuing authority | Secondary identifying number | |
| NCT04353492 | Clinical trials.gov | |

Sources of Monetary or Material Support

Name

Novartis Pharma services Inc.

Secondary Sponsors

Halim Abboud

Name

Public

NA

| Contact for Public/Scientific Queries | | | | | | |
|---------------------------------------|-------------------|----------|---------|------------------|-----------------------------------|--|
| Contact type | Contact full name | Address | Country | Telephone | Email | Affiliation |
| Public | Taghrid El Hajj | Beirut | Lebanon | 961349400 8 | taghridelhajj@gm ail.com | Rafik Hariri University Hospital |
| Scientific | Hind Khairallah | Sinelfil | Lebanon | 01512002# 271 | Hind.khairallah@ fattal.com.lb | Khalil Fattal et Fils s.a.l |

Beirut

| Centers/Hospitals Involved in the Study | | | |
|---|---|-----------|----------|
| Center/Hospital name | Name of principles investigator Principles investigator Speciality Ethical approval | | |
| Rafik Hariri University Hospital | Taghrid El Hajj | Neurology | Approved |
| Hotel Dieu De France | Halim Abboud | Neurology | Approved |

Hotel Dieu

De France

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Lebanon



| Ethics Review | | | | |
|-------------------------------------|---------------|---------------|--------------------------|--------------------|
| Ethics approval obtained | Approval date | Contact name | Contact email | Contact phone |
| Rafic Hariri University Hospital | 02/12/2020 | Rawan Yammout | rawan.yamout@crurhuh.com | 018300000 ext 2037 |
| Hotel Dieu de France | 03/11/2020 | Sami Richa | cue@usj.edu.lb | 961421229 |

| Countries of Recruitment |
|--------------------------|
| Name |
| Lebanon |
| Australia |
| Austria |
| Belgium |
| Bulgaria |
| Czech Republic |
| Germany |
| Greece |
| Hungary |
| Norway |
| Poland |
| Portugal |
| Russian Federation |
| Slovakia |
| Spain |
| Switzerland |
| Turkey |
| United States of America |



| Health Conditions or Problems Studied | | |
|---------------------------------------|--------------------------|----|
| Condition Code Keyword | | |
| Relapsing Multiple sclerosis | Multiple sclerosis (G35) | MS |

| Interventions | | |
|--|--|----------|
| Intervention | Description | Keyword |
| Informed Consent form , IMP administration , Visit assessment and schedule | Informed Consent form , IMP administration , Visit assessment and schedule | ICF, IMP |

| Primary Outcomes | | |
|---------------------|-------------|---|
| Name | Time Points | Measure |
| Annual Relapse Rate | 96 weeks | number of confirmed relapses in a year calculated based on cumulative number of relapses by patient adjusted for time-in-study by patient |

| Key Secondary Outcomes | | |
|------------------------|-------------|---|
| Name | Time Points | Measure |
| Safety evaluation | 96 weeks | Proportion of patients with adverse events, including injection related reactions, abnormal laboratory results or vital signs as well as proportion of patients discontinuing treatment due to insufficient effectiveness or safety |



| Trial Results | |
|--------------------------------------|--|
| Summary results | |
| Study results globally | |
| Date of posting of results summaries | Date of first journal publication of results |
| Results URL link | |
| Baseline characteristics | |
| Participant flow | |
| Adverse events | |
| Outcome measures | |
| URL to protocol files | |
| | |