Subcutaneous Interferon beta therapy in multiple sclerosis patients and characterization of injection site reactions and flu-like symptoms under daily practice setting (PERFECT)

First published: 22/11/2017 Last updated: 02/04/2024



## Administrative details

#### **EU PAS number**

EUPAS21013

#### **Study ID**

36739

#### DARWIN EU® study

No

#### **Study countries**

Germany

#### **Study description**

The purpose of this study is to investigate the safety of subcutaneous (SC) interferon beta therapies with regard to frequency of injection site reactions (ISR) and flu-like symptoms (FLS) as reported by the relapsing-remitting multiple sclerosis (RRMS) participants.

#### Study status

Finalised

## Research institutions and networks

## Institutions

## Biogen

First published: 01/02/2024

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Institution

# Multiple centres: 54 centres are involved in the study

## Contact details

Study Institution contact Study Director Biogen CTRR@biogen.com CTRR@biogen.com

Primary lead investigator Study Director Biogen

Primary lead investigator

# Study timelines

Date when funding contract was signed Planned: 31/12/2017 Actual: 31/12/2017

**Study start date** Planned: 31/12/2017 Actual: 16/11/2017

Data analysis start date Planned: 30/08/2019

Date of final study report Planned: 31/12/2019 Actual: 16/06/2020

# Sources of funding

• Pharmaceutical company and other private sector

## More details on funding

Biogen

## Regulatory

Was the study required by a regulatory body?

No

Is the study required by a Risk Management Plan (RMP)?

Not applicable

# Other study registration identification numbers and links

GER-PEG-16-10988

## Methodological aspects

## Study type

# Study type list

#### Study topic:

Disease /health condition Human medicinal product

#### Study type:

Non-interventional study

#### Scope of the study:

Safety study (incl. comparative)

#### Data collection methods:

Primary data collection

#### Main study objective:

The purpose of this study to investigate the safety of subcutaneous (SC) interferon beta therapies with regard to frequency of injection site reactions (ISR) and flu-like symptoms (FLS) as reported by the relapsing-remitting multiple sclerosis (RRMS) participants.

## Study Design

#### Non-interventional study design

Cross-sectional Other

#### Non-interventional study design, other

Post-authorization safety study

# Study drug and medical condition

## Study drug International non-proprietary name (INN) or common name INTERFERON BETA-1A INTERFERON BETA-1B

#### Medical condition to be studied

Relapsing-remitting multiple sclerosis

## **Population studied**

#### Short description of the study population

Adult relapsing-remitting multiple sclerosis (RRMS) patients receiving a Subcutaneous (SC) Interferon beta therapy.

Inclusion criteria

- Ability to understand the purpose of the study and provide signed and dated informed consent
- At least 18 years at time of informed consent
- Diagnosed relapsing-remitting multiple sclerosis
- Currently receiving a SC Interferon beta treatment (label conform)
- Stable on SC Interferon beta treatment for three months or longer (switch

between SC Interferon beta treatments possible)

Exclusion criteria

- Contraindications according to the "Fachinformation" (German equivalent to Summary of Product Characteristics [SmPC])
- Therapy with Glatiramer acetate or intramuscular (IM) Interferon beta-1a
- Participation in a non-interventional or interventional clinical study of Biogen

#### Age groups

Adults (18 to < 46 years) Adults (46 to < 65 years) Adults (65 to < 75 years) Adults (75 to < 85 years) Adults (85 years and over)

#### Special population of interest

Other

#### Special population of interest, other

Relapsing-remitting multiple sclerosis patients

626

# Study design details

#### Outcomes

• Percentage of Participants with at Least one Injection Site Reaction (ISR) as Reported by the Participants• Percentage of Participants with at Least one Flulike Symptoms (FLS) as Reported by the Participants, Participants with at Least 1 ISR (FLS), Types of ISR, Duration and Time of Occurrence of ISR Reported by Participants, Nurses, and PhysicianInterference Score of ISR With Participant's Daily ActivitiesParticipants Taking Actions to Relieve the ISR (FLS) and Resulting in Relief of ISR/FLSParticipants With Reducing/Increasing Frequency and Intensity of ISR/FLS Compared to Previous Therapy

#### Data analysis plan

All documented data are analyzed by descriptive statistics, that is, no formal statistical hypothesis will be formulated and no statistical tests will be carried out. All analyses will be performed based on the data set of patients who were eligible for participation according to inclusion criteria and completed the questionnaire. If a patient omits a specific question, he will be set to missing in all analyses referring to this question. No data imputation will be performed.

## Documents

Study results

GER-PEG-16-10988\_CSR Synopsis\_16-Jun-2020.pdf(267.14 KB)

Data management

Data sources

#### Data sources (types)

Other

#### Data sources (types), other

Prospective patient-based data collection

## Use of a Common Data Model (CDM)

#### **CDM** mapping

No

## Data quality specifications

#### **Check conformance**

Unknown

#### **Check completeness**

Unknown

#### **Check stability**

Unknown

#### **Check logical consistency**

Unknown

## Data characterisation

#### **Data characterisation conducted**

No