A Post-marketing Safety Study to Investigate the Risk of Serious CV Events among Galcanezumab (Emgality®) Users Using the Japan Medical Data Center Claim Database (I5Q-JE-B007)

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## Administrative details

### **EU PAS number**

EUPAS104412

#### **Study ID**

104467

#### DARWIN EU® study

No

### **Study countries**

Japan

### **Study description**

The primary objective of this study is to describe the incidence rate of serious CV events among migraine patients treated with galcanezumab. This objective is descriptive and aims to describe the incidence and distribution of time to first serious CV events in adult patients treated with galcanezumab, overall, by age, gender, and treatment duration of galcanezumab.

### Study status

Planned

## Research institutions and networks

## Institutions

## Eli Lilly and Company

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# Contact details

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Study contact

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Primary lead investigator

Machiko Minatoya

Primary lead investigator

## Study timelines

**Date when funding contract was signed** Planned: 25/04/2023

Study start date Planned: 01/04/2020

Date of final study report Planned: 30/09/2027

## Sources of funding

• Pharmaceutical company and other private sector

## More details on funding

Eli Lilly Japan K.K

## Regulatory

### Was the study required by a regulatory body?

Yes

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

Non-EU RMP only

# Methodological aspects

### Study type:

Non-interventional study

### Scope of the study:

Assessment of risk minimisation measure implementation or effectiveness

### Main study objective:

The primary objective of this study is to describe the incidence rate of serious CV events among migraine patients treated with galcanezumab. This objective is descriptive and aims to describe the incidence and distribution of time to first serious CV events in adult patients treated with galcanezumab, overall, by age, gender, and treatment duration of galcanezumab.

# Study Design

### Non-interventional study design Cohort

# Study drug and medical condition

### Anatomical Therapeutic Chemical (ATC) code

(N02CD02) galcanezumab galcanezumab

### Medical condition to be studied

Migraine

# Population studied

#### 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)

#### **Estimated number of subjects**

6599

## Study design details

#### Outcomes

Serious CV events

#### Data analysis plan

Calculate the descriptive statistics of the patient characteristics at baseline among galcanezumab group. The incidence and distribution of time to serious CV events among migraine patients treated with galcanezumab, overall, by age, and gender will be described. Incidence of serious CV events of the special populations of interest including with CV events during 12 months prior to initiation of galcanezumab will be calculated. Subgroup analysis for duration of the galcanetumab treatment (e.g. ?18 mo) will be conducted to investigate incidence of serious CV events of galcanetumab users with long-term exposure. Time-to-event analysis using the Kaplan-Meier method will be performed. The patient will be either to have a event or censored at the end of follow-up period.

### Data management

## **ENCePP** Seal

The use of the ENCePP Seal has been discontinued since February 2025. The ENCePP Seal fields are retained in the display mode for transparency but are no longer maintained.

## Data sources

Data source(s), other JMDC Japan

Data sources (types) Administrative healthcare records (e.g., claims)

# 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