Post-authorisation Safety Study of Rimegepant in Patients with Migraine and History of Cardiovascular Disease in European Countries

First published: 24/03/2023 Last updated: 03/02/2025



### Administrative details

#### **EU PAS number**

EUPAS103990

#### **Study ID**

104168

#### DARWIN EU® study

No

#### **Study countries**

Denmark

Netherlands

∣Spain

#### **Study description**

As part of the risk management plan for rimegepant in Europe, this postauthorisation safety study (PASS) is being conducted to evaluate whether there is an increased risk of major adverse cardiovascular events (MACE) among patients with migraine and history of cardiovascular disease (CVD) initiating treatment with rimegepant compared with that among patients with migraine, with history of CVD, and being treated with other treatments for migraine, either continuing the current treatment or initiating a new one, other than rimegepant. The study will also describe the use of rimegepant in the initial years after approval in the same population.

#### Study status

Planned

### Research institutions and networks

### Institutions

### Pfizer

First published: 01/02/2024

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Institution

### University of Southern Denmark (SDU)

🗌 Denmark



### The PHARMO Institute for Drug Outcomes Research (PHARMO Institute)

☐ Netherlands

First published: 07/01/2022

Last updated: 24/07/2024

Institution

Institution

Laboratory/Research/Testing facility

(ENCePP partner

ENCePP partner

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France				
Spain				
Sweden				
United Kingdom				
United Kingdom (Northern Ireland)				
United States				
First published: 21/04/2010				
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(Not-for-profit )

# Fundació Institut Universitari per a la Recerca a l'Atenció Primària de Salut Jordi Gol i Gurina, IDIAPJGol

Spain

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Institution Educational Institution	Laboratory/Research/Testing facility
Not-for-profit ENCePP partner	

# Contact details

### Study institution contact Caroline White Caroline.White@pfizer.com

Study contact

Caroline.White@pfizer.com

Primary lead investigator

Joan Forns

Primary lead investigator

# Study timelines

**Date when funding contract was signed** Planned: 01/03/2023 Actual: 01/03/2023

Study start date

Planned: 01/10/2025

Date of final study report Planned: 01/04/2029

## Sources of funding

• Pharmaceutical company and other private sector

### More details on funding

Pfizer 100%

# Study protocol

C4951017\_bhv3000-408 protocol v2\_17 Nov 2022.pdf(1.29 MB)

C4951017\_PROTOCOL- RIMEGEPANT CV PASS\_V3.0\_26AUG2024.pdf(2.29 MB)

### Regulatory

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

Yes

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

EU RMP category 3 (required)

### Methodological aspects

### Study type

**. . . . .** 

#### Study topic:

Human medicinal product

#### Study type:

Non-interventional study

#### Scope of the study:

Safety study (incl. comparative)

#### Data collection methods:

Secondary use of data

#### Study design:

This is a non-interventional population-based prospective cohort study using a prevalent newuser design.

#### Main study objective:

The study has 2 primary objectives:

1. To evaluate whether treatment initiation with rimegepant versus treatment with other preventive treatment for migraine (either continuing the current treatment or initiating a new one) increases the risk of MACE in patients with migraine, with history of CVD, and who are being treated with preventive migraine therapies.

2. To evaluate whether treatment initiation with rimegepant versus treatment with other acute treatment for migraine (either continuing the current treatment or initiating a new one) increases the risk of MACE in patients with migraine, with history of CVD, and who are being treated with acute migraine therapies.

### Study Design

#### Non-interventional study design

Cohort

# Study drug and medical condition

#### Name of medicine, other

Vydura

Study drug International non-proprietary name (INN) or common name RIMEGEPANT

### Anatomical Therapeutic Chemical (ATC) code

(N02CD06) rimegepant rimegepant

## **Population studied**

#### Short description of the study population

The study population will include adults with migraine (Protocol Section 9.3.3.1) and history of CVD ( Protocol Section 9.3.3.2) registered in each electronic health care data source who are on treatment with a qualifying acute or preventive migraine medication ( Protocol Table 3) during the study period.

#### Age groups

Adult and elderly population ( $\geq$ 18 years)

#### Estimated number of subjects

2500

### Study design details

#### Outcomes

Major adverse cardiovascular event (MACE), Individual components of major adverse cardiovascular event (MACE) including acute myocardial infarction, stroke, coronary heart disease death, cerebrovascular death, coronary bypass surgery, and coronary revascularization.

#### Data analysis plan

Each research partner will conduct analyses separately within each data source, and results will be pooled via meta-analytic methods, if appropriate. The analysis will comprise 4 different steps: select the study population, assign exposure and define follow-up, describe the study cohorts and patterns of rimegepant use, and estimate exposure propensity scores. Stabilised propensity score weights will be used in the comparative analyses. Crude and adjusted incidence rates of MACE with their 95% CIs will be estimated using a Poisson regression model with robust estimation of variance. Cumulative incidence of MACE will be estimated using the Kaplan-Meier estimator for each of the 4 exposure groups. Finally, for each comparison, crude and adjusted RRs and risk differences will be estimated using the Kaplan-Meier estimator, and 95% CIs will be derived using bootstrap methods. Adjusted HRs will be estimated with a Cox model.

### Documents

#### **Study report**

C4951017\_PROGRESS REPORT 1\_V1.0\_09Oct2024.pdf(169.57 KB)

### Data management

#### Data source(s)

The Information System for Research in Primary Care (SIDIAP) Danish registries (access/analysis) Clinical Practice Research Datalink PHARMO Data Network

#### Data sources (types)

Administrative healthcare records (e.g., claims) Drug dispensing/prescription data Electronic healthcare records (EHR) Other

**Data sources (types), other** Prescription event monitoring

## 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