Mavacamten Real-World Safety: a postauthorisation multi-national, long-term, observational study in patients with obstructive Hypertrophic Cardiomyopathy (oHCM) in the real-world setting in Europe (CV027-013)

First published: 03/01/2024

**Last updated:** 05/02/2025





### Administrative details

#### **EU PAS number**

EUPAS108178

#### **Study ID**

108179

### **DARWIN EU® study**

No

Study countries  Denmark  France  Germany  Sweden  United Kingdom
Study description  The aim of this study is to evaluate the risk of clinically important cardiovascular safety-related outcomes among adult patients with symptomatic oHCM receiving mavacamten compared to non-mavacamten oHCM treatment in a real-world setting in Europe.
Study status Planned Research institutions and networks Institutions
Bristol-Myers Squibb (BMS)  First published: 01/02/2024  Last updated: 01/02/2024  Institution
IQVIA  United Kingdom

**First published:** 12/11/2021

**Last updated:** 22/04/2024

Institution

Non-Pharmaceutical company

**ENCePP** partner

### Contact details

### **Study institution contact**

Transparency and Disclosure Lead ctt.group@bms.com

Study contact

ctt.group@bms.com

### Primary lead investigator

Sophie Shen

**Primary lead investigator** 

# Study timelines

Date when funding contract was signed

Planned: 30/06/2024

Study start date

Planned: 31/03/2026

Data analysis start date

Planned: 31/03/2026

**Date of final study report** 

Planned: 31/12/2034

# Sources of funding

• Pharmaceutical company and other private sector

# More details on funding

Bristol-Myers Squibb

# 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 type list

Study type:

Non-interventional study

### Main study objective:

The main objective of this study is to estimate the incidence rates of heart failure (HF) associated with exposure to mavacamten and with exposure to non-mavacamten oHCM treatment, among adult patients with symptomatic oHCM and to estimate the risk of HF associated with exposure to mavacamten compared with non-mavacamten oHCM treatment, among adult patients with symptomatic oHCM.

# Study Design

#### Non-interventional study design

Cohort

## Study drug and medical condition

#### Name of medicine

**CAMZYOS** 

## 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**

3648

## Study design details

#### **Outcomes**

Incidence rate (IR) of Heart Failure (HF) Risk of HF, HF with systolic dysfunction Hospitalisation due to HF Non-fatal myocardial infarction (MI) Non-fatal stroke Arrhythmia Hospitalisation for arrhythmia Atrial fibrillation Atrial flutter Ventricular tachycardia Implanted cardioverter defibrillator therapy 3-point MACE 4-point MACE eMACE Cardiovascular mortality All-cause mortality

#### **Data analysis plan**

Data analysis will be performed separately for each registry or database, using statistical packages SAS® Version 9.4 or higher (SAS Institute, Cary, NC, USA) or R (Version 3.5.0 or above), as appropriate.

In the study reports, study results will be presented separately for each registry or database, as appropriate, per data availability.

The complete study results for all registries/databases, including descriptive, comparative, exploratory, and sensitivity analyses, as well as the meta-analysis results (if feasible) will be provided in the final report.

A full description of the analytical approach, data derivations, category definitions, analyses and presentation of the study results will be detailed in the SAP.

The SAP will be finalised prior to the conduct of the study analyses.

# 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

Register of Hypertrophic Cardiomyopathy (REMY) France, Translational Registry for Cardiomyopathies (TORCH) Germany, Statutory Health Insurance (SHI) Claims Germany, Clinical Practice Research Datalink (CPRD) Aurum United Kingdom, Danish National Health and Socioeconomic registries Denmark

#### **Data sources (types)**

Administrative healthcare records (e.g., claims)

Disease registry

## 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