# EU-Wide Cross-Sectional Observational Study of Lipid-Modifying Therapy Use in Secondary and Primary Care DA VINCI

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### Administrative details

EU PAS number	
EUPAS22075	
Study ID	
31479	
DARWIN EU® study	
No	
Study countries	
Austria	
Belgium	
Czechia	
☐ Denmark	

France
Germany
Greece
Hungary
Ireland
Italy
Netherlands
Poland
Romania
Slovakia
Spain
Sweden
Ukraine
United Kingdom
Study status
Study status Finalised
Finalised
Research institutions and networks
Research institutions and networks Institutions
Research institutions and networks Institutions  Amgen
Research institutions and networks Institutions  Amgen United States

### **Amgen**

Multiple centres: 153 centres are involved in the study

### Contact details

### **Study institution contact**

Global Development Leader Amgen Inc. medinfo@amgen.com

Study contact

medinfo@amgen.com

### Primary lead investigator

Global Development Leader Amgen Inc.

**Primary lead investigator** 

## Study timelines

### Date when funding contract was signed

Planned: 17/11/2016 Actual: 17/11/2016

### Study start date

Planned: 01/06/2017

Actual: 01/06/2017

### Data analysis start date

Planned: 22/02/2019 Actual: 22/02/2019

### **Date of final study report**

Planned: 19/09/2019 Actual: 16/08/2019

## Sources of funding

• Pharmaceutical company and other private sector

### More details on funding

Amgen Ltd.

## Study protocol

EUPAS22075-23671.pdf(832.58 KB)

## Regulatory

Was the study required by a regulatory body?

No

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

Not applicable

## Methodological aspects

### Study typo

#### **Study topic:**

Human medicinal product

Disease /health condition

### Study type:

Non-interventional study

### Scope of the study:

Other

#### If 'other', further details on the scope of the study

Characterise unmet medical need

#### **Data collection methods:**

Secondary use of data

### Main study objective:

To estimate the proportion of subjects in EU primary and secondary care, with or without established ASCVD and receiving LMT, with LDL-C above 2016 Joint ESC Guideline recommended levels

### Study Design

### Non-interventional study design

Cross-sectional

## Study drug and medical condition

#### Name of medicine

#### REPATHA

#### Medical condition to be studied

Ischaemic stroke

Myocardial infarction

Peripheral vascular disorder

Hypercholesterolaemia

## Population studied

### Short description of the study population

- 1. Low density lipoprotein-cholesterol (LDL-C) measurement within 14 months of enrolment, obtained independently of participation in a clinical trial
- 2. Use of any LMT (may include statin/ezetimibe/fibrate/PCSK9 inhibitor/bile acid absorption inhibitor/nicotinic acid/other) at time of enrolment, or any LMT prescribed within 12 months prior to date of enrolment, or any LMT prescribed at date of enrolment
- 3. Age 18 years or older at enrolment
- 4. Provided informed consent/notified according to local requirements
- 5. Subject expected to survive for at least 1 year after enrolment

#### 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**

6000

## Study design details

#### **Outcomes**

LDL-C measurement most recent to enrolment, Lipid levels (total cholesterol, non-HDL-C, HDL, triglycerides, Lp(a), apo B100, apo A1) most recent to enrolment, use of Lipid-modifying therapy (type, dose, frequency), clinical characteristics at time of enrolment

#### Data analysis plan

All summaries of data will be descriptive in nature: categorical variables in frequency and percentage with 95% confidence intervals and continuous variables in mean (standard deviation) and/or median with percentiles, minimum and maximum.

### **Documents**

#### Study results

20150333\_DA VINCI FINAL ORSR abstract 16AUG19 incl PI signature - 120919\_redacted.pdf(407.3 KB)

### Data management

### Data sources

<b>Data source</b> Other	s (types)
Data source	s (types), other
Medical notes	(primary and secondary care)
Use of a	Common Data Model (CDM)
CDM mappir	ıg
No	
Data qua	ality specifications
Check confo	rmance
Unknown	
Check comp	leteness
Unknown	
Check stabil	ity
Unknown	
Check logica	al consistency
Unknown	

## Data characterisation

### **Data characterisation conducted**

No