# Romiplostim effectiveness stratified on duration of immune thrombocytopenia at initiation

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## Administrative details

EU PAS number	
EUPAS32183	
Study ID	
36382	
DARWIN EU® study	
No	
Study countries United States	

## **Study status**

**Finalised** 

Research institutions and networks

## **Institutions**

Amge	en
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☐ United States

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Institution

NoviSci, Inc. United States

## 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: 22/01/2019 Actual: 22/01/2019

#### Study start date

Planned: 01/11/2019 Actual: 17/10/2019

#### **Data analysis start date**

Planned: 17/10/2019 Actual: 17/10/2019

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

Planned: 12/06/2020 Actual: 16/07/2020

# Sources of funding

• Pharmaceutical company and other private sector

# More details on funding

Amgen

# Study protocol

01.02.06 Public Redacted Protocol Ver 1.0 2019-11-19 English.pdf (297.16 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 type	
Study type list	
Study topic: Human medicinal product Disease /health condition	
Study type: Non-interventional study	
Scope of the study: Drug utilisation Effectiveness study (incl. comparative)	
Data collection methods: Secondary use of data	
Main study objective:	

Among patients who received romiplostim in routine care in one of seven countries (Austria, Belgium, Czech Republic, France, Greece, Portugal and Sweden), describe the following overall and within strata of duration of ITP: patient profile at romiplostim initiation, patterns of romiplostim use and overall platelet count trends, effectiveness of romiplostim via platelet-based and other endpoints.

## Study Design

#### Non-interventional study design

Other

#### Non-interventional study design, other

Secondary analysis of data collected through a European, multi-center, observational cohort study (Amgen Study 20070225)

# Study drug and medical condition

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

#### Medical condition to be studied

Immune thrombocytopenia

# Population studied

Short description of the study population

The previously conducted cohort study (Amgen Study 20070225) enrolled 340 eligible participants. Patients were ≥ 18 years old, diagnosed with primary ITP, and had received at least one dose of romiplostim. Exclusion criteria included receipt (or planned receipt) of platelet related products (eg, recombinant human thrombopoietin [rHuTPO], thrombopoietin receptor agonists, pegylated recombinant human megakaryocyte growth and development factor [PEG-rHuMGDF]), participation in any interventional clinical study, or initiation of romiplostim prior to commercial availability of the product

#### 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

Immune thrombocytopenic purpura patients

## **Estimated number of subjects**

340

# Study design details

#### **Outcomes**

Durable platelet response, median overall platelet count, overall platelet response, time to first platelet response, Discontinuation of concurrent medications, bleeding, splenectomy, adverse drug reaction, thrombotic events, bone marrow fibrosis

#### Data analysis plan

Descriptive statistics of the clinical and demographic characteristics of the cohort will be provided, as will patterns of romiplostim use and platelet count trends. Estimates of medians, proportions, probabilities, and rates will be calculated for the endpoints of interest and presented with appropriate measures of variability. No hypotheses will be tested.

## **Documents**

#### Study results

20190407\_Observational Research Study Report Published Report Redacted.pdf (149.37 KB)

## 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

ta sources (types), other	
mpleted retrospective chart review in European clinical pract	ices
se of a Common Data Model (CDM)	
M mapping	
ata quality specifications	
eck conformance	
known	
eck completeness	
known	
eck stability	
known	

# Data characterisation

## **Data characterisation conducted**

No