# A GLOBAL PROSPECTIVE OBSERVATIONAL REGISTRY OF PATIENTS WITH POMPE DISEASE

**First published:** 09/11/2023

**Last updated:** 29/09/2025





# Administrative details

EU PAS number	
EUPAS107299	
Study ID	
107300	
DARWIN EU® study	
No	
Study countries	
\[ \lambda rachtine	
Argentina  Australia	

Bosnia and Herzegovina
Brazil
Canada
Chile
Czechia
Denmark
Estonia
France
Germany
Greece
Hungary
Israel
Italy
Japan
Korea, Republic of
Netherlands
New Zealand
Poland
Portugal
Serbia
Slovakia
Slovenia
Spain
Sweden
Taiwan
Thailand
United Kingdom
United States

# Study description

The goal of this registry is to assess clinical outcomes in patients with Pompe disease, including patients with late-onset Pompe disease (LOPD) or infantile-onset Pompe disease (IOPD), regardless of current or previous therapy.

#### **Study status**

Planned

# Research institutions and networks

#### **Institutions**

# **Amicus Therapeutics**

First published: 01/02/2024

**Last updated:** 01/02/2024

Institution

# Contact details

#### **Study institution contact**

Guiliano Joseph jgiuliano@amicusrx.com

Study contact

jgiuliano@amicusrx.com

## Primary lead investigator

Roberts Mark

Primary lead investigator

# Study timelines

#### Date when funding contract was signed

Planned: 01/10/2023

#### Study start date

Planned: 01/10/2023

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

Planned: 30/12/2033

# Sources of funding

• Pharmaceutical company and other private sector

# More details on funding

Amicus Therapeutics Inc

# 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)

# Other study registration identification numbers and links

# Methodological aspects

# Study type

# Study type list

#### **Study topic:**

Disease /health condition

#### Study type:

Non-interventional study

#### Scope of the study:

Disease epidemiology

Effectiveness study (incl. comparative)

Safety study (incl. comparative)

#### Main study objective:

- To evaluate the long-term safety of Pompe disease treatments through collection of data that describe the frequency of AEs/SAEs occurring in Pompe disease patients - To evaluate the long-term real-world effectiveness - To evaluate the long-term real-world impact of Pompe disease treatments on QOL and PROs - To describe the natural history of untreated Pompe disease

# Study Design

#### Non-interventional study design

Cohort

# Study drug and medical condition

#### **Medicinal product name**

**POMBILITI** 

**OPFOLDA** 

# Population studied

#### Age groups

- Preterm newborn infants (0 27 days)
- Term newborn infants (0 27 days)
- Infants and toddlers (28 days 23 months)
- Children (2 to < 12 years)
- Adolescents (12 to < 18 years)
- 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**

500

# Study design details

#### Data analysis plan

No formal hypotheses will be tested in this registry. Demographics and medical history (including but not limited to disease stage, time since diagnosis, prior and concomitant medication use, prior treatment with ERTs, and co-morbidities) will be summarized using descriptive statistics, with number and percent for categorical variables, and n, mean, SD, SE of the mean, median, minimum, and maximum for continuous variables. Quantitative analyses may include incidence rates of events, reasons for dropout and discontinuation, time-to-event profiles, and descriptions of clinical outcomes such as impact on physical function, HRQOL, and the occurrence of AEs, including but not limited to IARs, hypersensitivity reactions (including anaphylaxis), immune complex related reactions, and medication errors in the home infusion setting. Demographics and medical history will also be summarized using descriptive statistics on different subgroups.

# 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 sources (types)

Disease registry

# Use of a Common Data Model (CDM)

CDM	man	nnna
CDII	map	פיייקי

No

# Data quality specifications

#### **Check conformance**

Unknown

#### **Check completeness**

Unknown

#### **Check stability**

Unknown

#### **Check logical consistency**

Unknown

# Data characterisation

#### **Data characterisation conducted**

No