Analysis of patients with arginase 1 deficiency treated with Loargys in standard clinical care enrolled in a European, noninterventional, multicentre registry

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# Administrative details

#### **EU PAS number**

EUPAS100000495

#### **Study ID**

100000495

#### **DARWIN EU® study**

No

## Study countries

Austria

France

Netherlands

### **Study description**

This is a post-authorisation efficacy study (PAES) conducted in collaboration with the European Registry and Network for Intoxication Type Metabolic Diseases (E-IMD) that will analyse the ARG1-D population of the pre-existing and active registry held by the E-IMD.

The E-IMD international, multicentric, observational registry collects health data generated during the routine care of affected patients with urea cycle disorders, including ARG1-D.

#### **Study status**

Planned

# Research institutions and networks

## Institutions

Immedica Pharma AB
Sweden
First published: 30/06/2025
Last updated: 30/06/2025
Institution Pharmaceutical company

European registry and network for Intoxication type Metabolic Diseases (E-IMD)

First published: 01/02/2024

Last updated: 01/02/2024

# Heidelberg University Hospital

First published: 01/02/2024

Last updated: 01/02/2024

Institution

# Contact details

## Study institution contact Mattias Rudebeck clinical@immedica.com

Study contact

clinical@immedica.com

**Primary lead investigator** Mattias Rudebeck

Primary lead investigator

# Study timelines

Date when funding contract was signed Planned: 30/07/2025

Study start date

Planned: 03/11/2025

Data analysis start date

Planned: 15/12/2025

**Date of interim report, if expected** Planned: 02/02/2026

**Date of final study report** Planned: 04/02/2036

## Sources of funding

• Pharmaceutical company and other private sector

## Study protocol

IMM-PEG-003 Study Protocol v1.0 24Jun2025\_Redacted.pdf(444.91 KB)

# Regulatory

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

Yes

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

EU RMP category 2 (specific obligation of marketing authorisation)

**Regulatory procedure number** EMA/PAM/0000264956

## Methodological aspects

## Study type

### **Study topic:**

Human medicinal product

### Study type:

Non-interventional study

### Scope of the study:

Effectiveness study (incl. comparative)

#### Data collection methods:

Primary data collection

### Study design:

This PAES is conducted in collaboration with E-IMD that will analyse the ARG1-D population of the pre-existing and active registry held by the E-IMD.The E-IMD international, multicentric, observational registry collects health data generated during the routine care of patients with UCD, incl. ARG1-D

#### Main study objective:

The objective of the study is to evaluate the long-term effectiveness of Loargys treatment in patients with ARG1-D in standard clinical care

# Study Design

#### Non-interventional study design

Cohort

# Study drug and medical condition

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

### Anatomical Therapeutic Chemical (ATC) code

(A16AB24) pegzilarginase pegzilarginase

### Medical condition to be studied

Arginase deficiency

# **Population studied**

### Short description of the study population

Adult and paediatric patients enrolled in the E-IMD registry with a confirmed diagnosis of ARG1-D

### Age groups

Paediatric Population (< 18 years) Infants and toddlers (28 days – 23 months) Children (2 to < 12 years) Adolescents (12 to < 18 years) Adult and elderly population (≥18 years)

### Estimated number of subjects

15

# Study design details

#### Setting

European E-IMD centres treating patients with ARG1-D.

### 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)

European registry and network for intoxication type metabolic diseases

#### Data sources (types)

**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