# Cerliponase alfa Observational Study

**First published:** 26/03/2019

**Last updated:** 10/07/2024





# Administrative details

EU PAS number	
EUPAS29031	
Study ID	
46783	
DARWIN EU® study	
No	
Study countries	
Denmark	
France	
Germany	
Italy	
Netherlands	
Romania	
Sweden	

United	Kingdom
--------	---------

### **Study description**

This is a voluntary, multicenter, multinational, cerliponase alfa observational study for patients with a confirmed diagnosis of neuronal ceriod lipofuscinosis type 2 (CLN2 disease), also known as TPP1 deficiency, who intend to be or are currently being treated with cerliponase alfa.

#### **Study status**

Ongoing

### Research institutions and networks

### **Institutions**

ICON Commercialisation & Outcomes
Germany
Ireland
First published: 19/03/2010
<b>Last updated:</b> 05/07/2024
Institution Non-Pharmaceutical company ENCePP partner

### Contact details

### Study institution contact

Program 190-504 Director medinfo@bmrn.com

Study contact

medinfo@bmrn.com

### **Primary lead investigator**

Program 190-504 Director

**Primary lead investigator** 

## Study timelines

#### Date when funding contract was signed

Planned: 14/03/2019

Actual: 19/03/2019

### Study start date

Planned: 28/10/2019

Actual: 10/10/2019

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

Planned: 27/04/2030

## Sources of funding

• Pharmaceutical company and other private sector

### More details on funding

BioMarin Pharmaceutical

## Regulatory

Was the study	required b	y a regulatory	y body?
---------------	------------	----------------	---------

Yes

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

EU RMP category 2 (specific obligation of marketing authorisation)

## Methodological aspects

# Study type

## Study type list

#### Study type:

Non-interventional study

### Scope of the study:

Safety study (incl. comparative)

### Main study objective:

To evaluate long-term safety of cerliponase alfa in patients with neuronal ceroid lipofuscinosis Type 2 (CLN2 disease).

## Study Design

### Non-interventional study design

Other

### Non-interventional study design, other

## Study drug and medical condition

#### Name of medicine

**BRINEURA** 

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

**CERLIPONASE ALFA** 

#### **Anatomical Therapeutic Chemical (ATC) code**

(A16AB17) cerliponase alfa cerliponase alfa

#### Medical condition to be studied

Neuronal ceroid lipofuscinosis

#### Additional medical condition(s)

Neuronal ceroid lipofuscinosis type 2 (CLN2)

## Population studied

#### Age groups

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)

#### Special population of interest

Hepatic impaired

Immunocompromised

Renal impaired

### **Estimated number of subjects**

45

## Study design details

### Data analysis plan

All analysis will be descriptive. Assessments are collected based on the country, institution, and individual patient standard of care.

### 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

Other	(types)				
Data sources	(types), othe	r			
Prospective pa	ient-based dat	a collectio	n		
Use of a (	Common	Data N	Model (	CDM)	
CDM mapping					
No					
Data qua	ity spacit	fication	2.5		
Data qua	ity specii	icatioi	15		
Check confor		icatioi	15		
•		icatioi	15		
Check confor	nance	icatioi	15		
Check confor	nance	icatioi	15		
Check conford Unknown Check comple	nance teness	icatioi	15		

# Data characterisation

### **Data characterisation conducted**

No