# DARWIN EU® - Clozapine and the incidence of agranulocytosis over time

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# Administrative details

### **EU PAS number**

EUPAS100000549

### **Study ID**

100000549

#### **DARWIN EU® study**

Yes

#### **Study countries**

Croatia

🗌 Denmark

Finland

Germany

Spain

### **Study description**

Clozapine is an effective treatment for treatment-resistant schizophrenia and Parkinson's disease psychosis, but it carries a risk of severe hematological complications, including neutropenia and agranulocytosis.

Emerging evidence suggests that the risk is highest in the initial months of treatment, yet stringent hematological monitoring requirements remain in place throughout long-term use.

These requirements may hinder clinical practice, leading to underuse, early treatment discontinuation, or reluctance to initiate therapy.

This study aims to provide epidemiological evidence on the incidence and timing of clozapine-associated neutropenia and agranulocytosis across Europe.

### Study status

Ongoing

# Research institutions and networks

### Institutions



Netherlands

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### Networks

Data Analysis and	Real	World	Interrogation	Network
(DARWIN EU®)				

Belgium

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Croatia

Denmark

- Estonia
- Finland
- France

Germany

Greece

Hungary

Italy

Netherlands

Norway

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Sweden

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# Contact details

Study institution contact

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Study contact

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Primary lead investigator Dina Vojinovic

Primary lead investigator

# Study timelines

### Date when funding contract was signed Planned: 17/02/2025 Actual: 17/02/2025

**Study start date** Planned: 31/03/2025 Actual: 31/03/2025

Date of final study report Planned: 30/05/2025

### Sources of funding

• EMA

### Study protocol

DARWIN EU\_Protocol\_P4-C1-001\_Clozapine\_agranulocytosis\_V2.pdf(760.15 KB)

# Regulatory

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

Yes

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

Not applicable

# Methodological aspects

# Study type

# Study type list

### Study topic:

Human medicinal product

### Study type:

Non-interventional study

### Scope of the study:

Disease epidemiology

### Data collection methods:

Secondary use of data

### Study design:

A cohort study will be conducted using routinely collected health data from 5 data sources. The study will comprise three consecutive parts:

• Population-level cohort study (Objective 1, Population-level descriptive

epidemiology of agranulocytosis and neutropenia in new users of clozapine).

Cohort

### Main study objective:

1. To estimate the incidence rates of agranulocytosis and neutropenia in consecutive weekly and monthly intervals following the initiation of clozapine treatment, overall and stratified by age and sex.

2. To characterise the timing of agranulocytosis and neutropenia events during clozapine treatment using Kaplan-Meier curves, overall and stratified by age and sex.

3. To characterise individuals initiating clozapine treatment in terms of demographics and pre-specified conditions related to the indication for clozapine use.

4. To determine the treatment duration for clozapine use.

# Study Design

### Non-interventional study design

Cohort

# Study drug and medical condition

### Name of medicine, other

Clozapine

### Anatomical Therapeutic Chemical (ATC) code

(N05AH02) clozapine clozapine

### Medical condition to be studied

Agranulocytosis

# Population studied

### Short description of the study population

The study population will include all new users of clozapine registered in the respective data sources between 1st of January 2010 and 31st of December 2024 (or latest date available).

"New use" refers to a first prescription of clozapine within the study period with no clozapine prescription in the medical history.

Eligibility Criteria:

At least 1 year of data visibility before starting clozapine treatment. Additional eligibility criteria:

To ensure sufficient follow-up, only individuals who initiated clozapine treatment at least one year before the end of the available data in the respective data source will be included.

### Age groups

All In utero Paediatric Population (< 18 years) Neonate 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) Adult and elderly population ( $\geq$ 18 years) Adults (18 to < 65 years) Adults (18 to < 46 years) Adults (46 to < 65 years) Elderly ( $\geq$  65 years) Adults (65 to < 75 years) Adults (75 to < 85 years) Adults (85 years and over)

### Data management

### Data sources

### Data source(s)

Danish Health Data Registries Croatia National Public Health Information System (Nacionalni javnozdravstveni informacijski sustav) IQVIA Disease Analyzer Germany The Information System for Research in Primary Care (SIDIAP)

Data source(s), other

FinOMOP-HILMO

# Use of a Common Data Model (CDM)

### **CDM mapping**

Yes

**CDM Mappings** 

#### **CDM** name

OMOP

#### **CDM** website

https://www.ohdsi.org/Data-standardization/

#### **CDM version**

https://ohdsi.github.io/CommonDataModel/index.html

# Data quality specifications

#### **Check conformance**

Unknown

#### **Check completeness**

Unknown

### **Check stability**

Unknown

#### **Check logical consistency**

Unknown

### Data characterisation

#### **Data characterisation conducted**

Unknown