# DARWIN EU® – Paracetamol prescribing and paracetamol overdose in Europe: a descriptive analysis of trends and patient characteristics

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**Last updated:** 03/03/2025





# Administrative details

EU PAS number
EUPAS1000000329
Study ID
100000329
DARWIN EU® study
Yes
Study countries
Croatia
Denmark
France

Germany	
Portugal	
Spain	
United Kingdom	

### Study description

Paracetamol (acetaminophen) is one of the most widely used medicines worldwide and is available over the counter in the European Union. It is one of the most common causes of drug poisonings and can result in severe hepatic failure. Different regulatory interventions at national level have occurred to reduce the incidence of paracetamol overdose, but it is uncertain how paracetamol is prescribed across Europe and to what extent prescription may be involved in poisonings.

Research question and objectives

The aim of the study is to provide an overview of paracetamol prescribing and paracetamol overdose trends in selected European databases, and to characterise patients presenting with paracetamol overdose.

The specific objectives of the study are:

- 1. To examine the incidence/prevalence of paracetamol prescribing (overall, and by age, sex, formulation and country/database).
- 2. To examine the incidence of paracetamol overdose (overall, and by age, sex, country/database).
- 3. To characterise patients with paracetamol overdose, in terms of comorbidities, co-prescribed medications, prior paracetamol prescription, and incidence of short-term complications and mortality.

### **Study status**

Finalised

# Research institutions and networks

# Institutions

Department of Medical Informatics - Health Data
Science, Erasmus Medical Center (ErasmusMC)
☐ Netherlands
First published: 03/11/2022
Last updated: 02/05/2024
Institution

# **Networks**

Data Analysis and Real World Interrogation Network
(DARWIN EU®)
Belgium
Croatia
Denmark
Estonia
Finland
France
Germany
Greece
Hungary
☐ Italy
☐ Netherlands
Norway

Portugal
Spain
Sweden
United Kingdom
First published: 01/02/2024
Last updated: 30/04/2025
Network

# Contact details

### **Study institution contact**

Ilse Schuemie study@darwin-eu.org

Study contact

study@darwin-eu.org

# **Primary lead investigator**

Berta Raventos

**Primary lead investigator** 

# Study timelines

Date when funding contract was signed

Planned: 24/07/2024

Actual: 24/07/2024

Study start date

Planned: 12/09/2024

Actual: 12/09/2024

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

Planned: 07/02/2025

Actual: 06/02/2025

# Sources of funding

EMA

# Study protocol

DARWIN EU\_Protocol\_P3-C1-007\_Paracetamol\_V2.pdf(724.56 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:

Drug utilisation

### **Data collection methods:**

Secondary use of data

### Study design:

- 1. Population-level drug utilisation study to assess incidence and prevalence of paracetamol prescribing (objective 1) 2. Population-level descriptive epidemiology study to estimate the incidence of paracetamol overdose (objective 2)
- 3. Patient-level characterisation study

### Main study objective:

The aim of the study is to provide an overview of paracetamol prescribing and paracetamol overdose trends in selected European databases, and to characterise patients presenting with paracetamol overdose.

The specific objectives of the study are:

- 1. To examine the incidence/prevalence of paracetamol prescribing (overall, and by age, sex, formulation and country/database).
- 2. To examine the incidence of paracetamol overdose (overall, and by age, sex, country/database).
- 3. To characterise patients with paracetamol overdose, in terms of

comorbidities, co-prescribed medications, prior paracetamol prescription, and incidence of short-term complications and mortality.

# Study Design

### Non-interventional study design

Cohort

# Study drug and medical condition

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

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

(N02BE01) paracetamol paracetamol

# Population studied

### Short description of the study population

For objective 1 and 2, the study population will comprise all individuals present in the database at any time from 1st January 2010 to 31st of December 2023 (or the latest year with complete observation). For objective 3, the study will comprise of individuals with paracetamol overdose for the first time in their patient history during the study period.

For incidence calculations (objective 1 and 2), individuals with a record of the outcome will re-enter the study after a certain amount of time (i.e. washout

period) so further occurrences of the outcome can be captured. This washout period will be defined as 60 days following the end of the prescribed treatment for paracetamol prescribing and 365 days for paracetamol overdose. For objective 3, individuals with a prior history of paracetamol overdose any time prior to index date will be excluded.

A year of observation history prior to index date will be required for all individuals within selected databases except CDWBordeaux. Individuals aged less than 1 year of observation history will be excluded.

# Study design details

### **Setting**

This study will be conducted using routinely collected data from 8 databases in 7 European countries selected from the DARWIN EU® Database Catalogue. All databases were previously mapped to the Observational Medical Outcomes Partnership (OMOP) Common Data Model (CDM).

# **Documents**

### Study report

DARWIN EU\_Report\_P3-C1-007\_Paracetamol\_V2.pdf(2.73 MB)

# Data management

# Data sources

### Data source(s)

Croatia National Public Health Information System (Nacionalni javnozdravstveni informacijski sustav)

BIFAP - Base de Datos para la Investigación Farmacoepidemiológica en el Ámbito Público (Pharmacoepidemiological Research Database for Public Health Systems)

Clinical Practice Research Datalink (CPRD) GOLD

Danish Health Data Registries

IQVIA Disease Analyzer Germany

**UK Biobank** 

Clinical Data Warehouse of the Bordeaux University Hospital

Egas Moniz Database

# 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

# Unknown

# **Check completeness**

**Check conformance** 

Unknown

# **Check stability**

Unknown

# **Check logical consistency**

Unknown

# Data characterisation

### **Data characterisation conducted**

Unknown