DARWIN EU® Drug Utilisation Study of prescription opioids

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Administrative details

EU PAS number

EUPAS100000479

Study ID

100000479

DARWIN EU® study

Yes

Study countries

Belgium

Denmark

____ Estonia

France

Netherlands

∣Norway

Study description

Prescription opioids, while effective for managing severe pain, have led to a public health crisis due to misuse, addiction, and overdose, particularly in the US.

Recently, concerns have been growing in Europe due to increasing opioid use and related mortality.

Factors such as chronic pain, mental health disorders, and advanced age can exacerbate misuse and the development of dependence.

Given the potential for global spread of this issue, enhanced surveillance and indepth research into opioid utilization patterns are imperative.

A drug utilization study using a Common Data Model (CDM) is a promising approach to supplement European opioid monitoring systems, providing more granular data to inform evidence-based decisions on this complex topic.

Study status

Ongoing

Research institutions and networks

Institutions

Department of Medical Informatics - Health Data Science, Erasmus Medical Center (ErasmusMC)

Netherlands

First published: 03/11/2022



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
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Contact details

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Primary lead investigator Amy Lam

Primary lead investigator

Study timelines

Date when funding contract was signed Planned: 04/09/2024 Actual: 04/09/2024

Study start date Planned: 04/02/2025 Actual: 04/02/2025

Date of final study report Planned: 30/04/2025

Sources of funding

• EMA

Study protocol

DARWIN EU Protocol_P3-C2-002_DUS Opioids_V3.pdf(862.45 KB)

Regulatory

Was the study required by a regulatory body? Yes

Is the study required by a Risk Management Plan (RMP)? Not applicable

Other study registration identification numbers and links

EUPAS105641 EUPAS1000000615

Methodological aspects

Study type

Study type list

Study topic:

Human medicinal product

Study type:

Non-interventional study

Data collection methods:

Secondary use of data

Study design:

A cohort study will be conducted using routinely-collected health data from 8 databases.

Study Design

Non-interventional study design

Cohort

Study drug and medical condition

Name of medicine, other

Opioids (substances listed in ATC classes N01AH, N02A and R05DA),, namely: acetyldihydrocodeine, alfentanil, anileridine, bezitramide, butorphanol, buprenorphine, codeine, dezocine, dimemorfan, dextromethorphan, dextromoramide, dextropropoxyphene, dihydrocodeine, ethylmorphine, fentanyl, hydrocodone, hydromorphone, ketobemidone, meptazinol, meperidine (pethidine), methadone, morphine, nicomorphine, normethadone, nalbuphine, noscapine, oliceridine, opium, oxycodone, oxymorphone, papaveretum, pentazocine, phenazocine, phenoperidine, pholcodine, pirinitramide, propoxyphene, remifentanil, sufentanil, tapentadol, thebacon, tilidine, tramadol; naloxone; buprenorphine/naloxone, oxycodone/naloxone,pentazocine/naloxone, tilidine/naloxone

Study drug International non-proprietary name (INN) or common name **BUTORPHANOL BUPRENORPHINE** CODEINE DEXTROMETHORPHAN DIHYDROCODEINE **ETHYLMORPHINE** FENTANYL **HYDROMORPHONE** METHADONE MORPHINE OXYCODONE PENTAZOCINE PHOLCODINE SUFENTANIL **TAPENTADOL** TILIDINE TRAMADOL

NALOXONE

Anatomical Therapeutic Chemical (ATC) code

(N01AH) Opioid anesthetics Opioid anesthetics (N02A) OPIOIDS OPIOIDS (R05DA) Opium alkaloids and derivatives Opium alkaloids and derivatives

Population studied

Short description of the study population

The study cohort will comprise all individuals present in the database during the study period (2012-2024) and with at least 365 days of data availability before the day they become eligible for study inclusion. Therefore, children aged <1 year will be excluded.

Age groups

All Paediatric Population (< 18 years) 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

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)

Estonian Biobank IQVIA Longitudinal Patient Data - Belgium Integrated Primary Care Information (IPCI) The Information System for Research in Primary Care (SIDIAP) Clinical Data Warehouse of the Bordeaux University Hospital Danish Health Data Registries Institut Municipal d'Assistència Sanitària Information System Norwegian Linked Health registry at University of Oslo

Use of a Common Data Model (CDM)

CDM mapping Yes CDM Mappings CDM name

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