# Evaluation of potential off-label use of dabigatran etexilate in Europe

First published: 08/10/2014

**Last updated:** 02/07/2024





# Administrative details

EU PAS number	
EUPAS7591	
Study ID	
25630	
DARWIN EU® study	
No	
Study countries	
Denmark	
France	
United Kingdom	

#### **Study description**

This is a descriptive, observational, multi-country European cross-sectional study of new users of dabigatran etexilate that aims to characterise on and off-label status and other medical characteristics at the time of the first captured prescription of dabigatran etexilate in each database. The study will be conducted using Cegedim Strategic Database (CSD, France), Danish National Databases (Denmark) and Clinical Practice Research Datalink (CPRD, UK).

## **Study status**

Finalised

# Research institutions and networks

# Institutions

RTI Health Solutions (RTI-HS)
France
Spain
Sweden
United Kingdom
United Kingdom (Northern Ireland)
United States
First published: 21/04/2010
Last updated: 13/03/2025
Institution Not-for-profit ENCePP partner

CSD Medical Research (CSDMR)

France
First published: 18/07/2012
Last updated: 20/08/2024
Institution Other
Clinical Practice Research Datalink (CPRD)
United Kingdom
First published: 15/03/2010
Last updated: 17/01/2025
Institution (Laboratory/Research/Testing facility) (ENCePP partner)
Pharmacoepi center, University of Southern
Denmark
Denmark
First published: 22/04/2010
Last updated: 27/07/2023
Institution

# Contact details

**Study institution contact** 

# Manel Pladevall-Vila mpladevall@rti.org

Study contact

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## **Primary lead investigator**

Manel Pladevall-Vila

**Primary lead investigator** 

# Study timelines

## Date when funding contract was signed

Actual: 27/11/2013

#### Study start date

Planned: 03/11/2014

Actual: 28/11/2014

#### Date of final study report

Planned: 30/11/2016

Actual: 15/12/2016

# Sources of funding

• Pharmaceutical company and other private sector

# More details on funding

Boehringer Ingelheim GmbH

# Study protocol

1160-144--protocol-revision-04 redacted.pdf(1.66 MB)

# Regulatory

Was the study required by a regulatory body?

Yes

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

EU RMP category 3 (required)

# 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:**

#### Main study objective:

Estimate the proportion of off-label use in new users of dabigatran according to the recorded clinical indication or generated proxies as available in each DB.Describe the characteristics of new users of dabigatran including dose, demographics, clinical indication, morbidity and use of other medications prior to the first captured prescription, stratified by usage sub-group—on- or off-label use.

# Study Design

### Non-interventional study design

Cross-sectional

# Study drug and medical condition

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

# Population studied

## Short description of the study population

Patients who received a new prescription of dabigatran etexilate in the study period and have at least 1 year of enrolment in the electronic database and have not been prescribed dabigatran etexilate during the 1-year period prior to the index date.

#### Age groups

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)

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**

Renal impaired

#### **Estimated number of subjects**

15000

# Study design details

#### **Outcomes**

The main outcome of this study is the proportion of off-label use estimated among new users of dabigatran etexilate, new users will be characterised.

#### **Data analysis plan**

The analyses will be descriptive at baseline. The main analysis will be to estimate (with 95% confidence intervals) the prevalence proportion of off-label use among new users of dabigatran etexilate during the overall study period in each of the study populations. All the results will be presented for each country-specific database. A weighted, pooled prevalence of off-label use among new

users of dabigatran etexilate for the entire study population (study populations of the CPRD, Cegedim, and Danish national databases combined) will be estimated (with 95% confidence intervals) when the individual results of all three databases are available.

## **Documents**

## **Study results**

nis-existing-data-report-text-part-0303644-bi-dabigatran\_redacted.pdf(182.25 KB)

### **Study publications**

Cainzos-Achirica M, Varas-Lorenzo C, Pottegård A, Asmar J, Plana E, Rasmussen L...

Cainzos-Achirica M, Varas-Lorenzo C, Pottegård A, Asmar J, Plana E, Rasmussen L...

# 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)

Clinical Practice Research Datalink

#### Data source(s), other

Cegedim Strategic Data - Longitudinal Patient Database France

## **Data sources (types)**

Administrative healthcare records (e.g., claims)

Drug dispensing/prescription data

Electronic healthcare records (EHR)

# 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**

Unknown