# Dissemination of additional risk minimisation measures for patients and healthcare professionals in EU/EEA countries

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# Administrative details

EU PAS number
EUPAS100000524
Study ID
100000524
DARWIN EU® study
No
Study countries
Finland
Hungary
Italy

Lithuania
☐ Netherlands
Romania
Study status
Ongoing
Research institutions and networks
Institutions
University of Naples Federico II
First published: 01/02/2024
Last updated: 01/02/2024
Institution
Netherlands Institute for Health Services Research
(Nivel)
☐ Netherlands
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Institution Not-for-profit

# Contact details

## **Study institution contact**

Anne Brabers a.brabers@nivel.nl

Study contact

a.brabers@nivel.nl

## **Primary lead investigator**

Liset Van Dijk

**Primary lead investigator** 

# Study timelines

## Date when funding contract was signed

Actual: 28/02/2025

#### Study start date

Actual: 28/02/2025

## Date of interim report, if expected

Planned: 28/04/2026

## **Date of final study report**

Planned: 29/06/2026

# Sources of funding

EMA

# Study protocol

# 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:

Assessment of risk minimisation measure implementation or effectiveness

#### **Data collection methods:**

Primary data collection

#### Study design:

Focus groups, interviews, and surveys among the different populations, desk research

#### Main study objective:

- 1. To describe and analyse aRMM dissemination: Process & frequency; key stakeholders involved; roles & responsibilities by type of aRMM, by dissemination method, by medicinal product and by country
- 2. To describe and analyse how access is ensured by type of aRMM, by medicinal product, by key stakeholder involved, and by country
- 3. To identify key challenges of disseminating healthcare professional- and patient-targeted aRMMs by type of aRMM, by dissemination method, by key stakeholder involved and by country
- 4. To identify and describe preferences for aRMM tools by type of aRMM, by dissemination method, by key stakeholder involved and by country
- 5. To provide recommendations by identifying feasible and concrete steps at each step of implementation pathway

# Study Design

## Non-interventional study design

Cross-sectional

# Study drug and medical condition

#### Name of medicine

**XELJANZ** 

**AUBAGIO** 

#### Name of medicine, other

Valproate containing medicinal products

Retinoids containing medicinal products

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

TOFACITINIB CITRATE

**TERIFLUNOMIDE** 

**ALEMTUZUMAB** 

**AFLIBERCEPT** 

**EDOXABAN** 

**ACITRETIN** 

**ADAPALENE** 

**ALITRETINOIN** 

**BEXAROTENE** 

**ISOTRETINOIN** 

**TRETINOIN** 

**TAZAROTENE** 

# Population studied

#### Short description of the study population

Patients using these drugs, HCPs prescribing these drugs, Competent Authorities, pharmacists, Marketing AuthorizationHolders (MAH)

## Data management

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**

No