

# XYREM EU-RMP: Effectiveness Assessment Protocol of Educational Materials

**First published:** 05/09/2016

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

Study

Finalised

## Administrative details

### EU PAS number

EUPAS15024

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### Study ID

29341

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### DARWIN EU® study

No

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### Study countries

☐ Belgium

☐ Germany

☐ Italy

☐ Spain

☐ Sweden

☐ United Kingdom

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## Study description

The overall research question is to evaluate the effectiveness of the educational material risk minimization measures (RMMs) being implemented in the EU to mitigate the important identified risks of respiratory and central nervous system (CNS) depression, depression and suicidality, abuse and misuse of XYREM, diversion and criminal use, overdose, dependency/withdrawal, and interactions of XYREM with alcohol, the important potential risks associated with sodium overload, in patients prescribed XYREM.

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## Study status

Finalised

## Contact details

### Study institution contact

Clinical Trial Registries and Results Personal identifiable data of lead investigator are not published here, as consent is not available. [clinicaltrials@ucb.com](mailto:clinicaltrials@ucb.com)

**Study contact**

[clinicaltrials@ucb.com](mailto:clinicaltrials@ucb.com)

### Primary lead investigator

Clinical Trial Registries and Results Personal identifiable data of lead investigator are not published here, as consent is not available.

**Primary lead investigator**

## Study timelines

**Date when funding contract was signed**

Actual: 07/12/2015

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**Study start date**

Actual: 01/01/2016

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**Data analysis start date**

Planned: 29/12/2017

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**Date of final study report**

Planned: 14/04/2019

Actual: 09/04/2019

## Sources of funding

- Pharmaceutical company and other private sector

## More details on funding

UCB BioPharma SPRL

## Study protocol

[NA0001 PASS Amendment 4 Protocol-final-redacted.pdf](#)(856.91 KB)

## Regulatory

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

Yes

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## 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

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**Study type:**

Non-interventional study

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**Scope of the study:**

Assessment of risk minimisation measure implementation or effectiveness

**Data collection methods:**

Primary data collection

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**Main study objective:**

Evaluate the effectiveness of the educational material RMMs implemented in the EU to mitigate the important identified risks of respiratory and CNS depression, depression/suicidality, abuse/misuse of XYREM, diversion/criminal use, overdose, dependency/withdrawal, and interactions of XYREM with alcohol, the important potential risks associated with sodium overload in patients prescribed XYREM.

## Study Design

## **Non-interventional study design**

Other

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## **Non-interventional study design, other**

Survey

# Study drug and medical condition

## **Name of medicine**

XYREM

# Population studied

## **Short description of the study population**

XYREM prescribers from Belgium, Germany, Italy, Spain, Sweden, and the UK to whom the XYREM educational materials were sent.

Physicians were required to meet the following inclusion criterion:

1. Must have been included on the XYREM RMP educational materials mailing list
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## **Age groups**

Adults (18 to < 46 years)

Adults (46 to < 65 years)

Adults (65 to < 75 years)

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## **Estimated number of subjects**

200

# Study design details

## Outcomes

Proportion of respondents providing a correct response for each individual question consistent with the XYREM summary of product characteristics (SmPC).

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## Data analysis plan

The primary outcome of the survey is the proportion of prescribers providing a correct response for each individual question in the key risk messages consistent with the XYREM EU SmPC. This is computed as the proportion of prescribers who correctly respond to the individual survey questions concerning the appropriate dosing and important identified risks of XYREM as described in the XYREM SmPC. The proportion responding correctly will be tabulated separately for each item in the survey instrument. Point estimates for the proportion with correct responses, and associated 95% confidence intervals, will be calculated for each question. In the case of multiple choice questions, the number and proportion of prescribers reporting each response will also be provided. Information obtained from the survey will be reported as descriptive statistics for the survey administration, survey population, and the survey questions.

## Documents

### Study results

[CSR\\_UCB\\_Xyrem Online Survey NA0001 PASS Study-final-redacted.pdf](#)(779.89 KB)

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## Data management

## Data sources

## **Data sources (types)**

Other

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### **Data sources (types), other**

Surveys will be used to assess the effectiveness of the risk minimisation methods available to prescribers

## Use of a Common Data Model (CDM)

### **CDM mapping**

No

## Data quality specifications

### **Check conformance**

Unknown

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### **Check completeness**

Unknown

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### **Check stability**

Unknown

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### **Check logical consistency**

Unknown

## Data characterisation

## **Data characterisation conducted**

Unknown