# Post Marketing Surveillance Study for Mircera

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## Administrative details

<b>EU PAS number</b> EUPAS4683		
Study ID		
20705		
DARWIN EU® study		
No		
Study countries  Korea, Republic of		

#### **Study description**

This study (ML22560) is a post-marketing prospective surveillance study (conducted in Korea from 29 Aug 2008 to 28 Aug 2012) to meet local regulatory requirements.

#### **Study status**

**Finalised** 

## Research institutions and networks

## **Institutions**

## F. Hoffmann-La Roche

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Multiple centres: 26 centres are involved in the

study

## Contact details

## Study institution contact

Jenny Shin jenny.shin@roche.com

Study contact

jenny.shin@roche.com

### **Primary lead investigator**

Petersen Jenny

**Primary lead investigator** 

## Study timelines

#### Date when funding contract was signed

Actual: 02/01/2008

#### Study start date

Actual: 29/08/2008

#### Data analysis start date

Actual: 28/08/2012

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

Actual: 28/11/2012

# Sources of funding

• Pharmaceutical company and other private sector

## More details on funding

F. Hoffmann-La Roche

## Regulatory

Was the study required by a regulatory body?

Yes

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

Non-EU RMP only

# Other study registration identification numbers and links

ML22560

## Methodological aspects

## Study type

# Study type list

#### **Study topic:**

Disease /health condition

Human medicinal product

#### Study type:

Non-interventional study

#### Scope of the study:

Assessment of risk minimisation measure implementation or effectiveness Effectiveness study (incl. comparative)

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

Primary data collection

#### Main study objective:

The main objectives are to evaluate the following items under routine practice:1) Serious adverse event(AE)/adverse drug reaction (ADR) 2)
Unexpected ADR 3) Expected ADR 4) Non-serious AE 5) AE occurred by misuse,

abuse and drug interaction 6) Any factors influencing safety and efficacy parameters (influencing lab. data, etc)

# Study Design

#### Non-interventional study design

Other

#### Non-interventional study design, other

Prescription event monitoring

# Study drug and medical condition

#### **Medicinal product name**

**MIRCERA** 

#### Medical condition to be studied

Anaemia

## Population studied

#### Short description of the study population

Patients who were prescribed Mircera® by their physician according to the local Korean Mircera® label, for the treatment of anemia associated with chronic kidney disease who require Erythropoiesis-Stimulating Agents (ESA) therapy.

#### Age groups

- Adults (18 to < 46 years)</li>
- Adults (46 to < 65 years)</li>
- 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**

600

# Study design details

#### **Data analysis plan**

According to the data, t-test or chi-square test was performed. Adverse events were tabulated in summary tables as followings: Summary table of incidence status of AEs/ADRs by System Organ Class Summary table of drug - adverse events relationship Summary table of Intensity of Adverse events/Adverse Drug Reactions Summary table of Incidence rate Unexpected AEs/ADRs by System Organ Class

# 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 sources (types)** 

Other

#### Data sources (types), other

Prospective patient-based data collection

## 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**

# Data characterisation

#### **Data characterisation conducted**

Unknown