Multicenter Observational study to evaluate the effectiveness of a biosimilar Etanercept (ErelziTM) in patients with established Rheumatic disease previously treated with reference etanercept. (BRONZE)

First published: 03/04/2019 Last updated: 20/01/2025



# Administrative details

#### **EU PAS number**

EUPAS28777

#### **Study ID**

43383

#### **DARWIN EU® study**

No

#### **Study countries**

France

### Study description

Study design: National, Multicenter, Prospective, Post-Authorization, non-PASS, Real Life, Observational Study. Primary objective: To evaluate effectiveness of ErelziTM in patients with RA, AS or PsA with, stable clinical response to Enbrel®. Secondary objectives:

• To assess safety in a real-life situation of switching from Enbrel® to ErelziTM.

• To characterize the patient populations and drug utilization patterns of RA, AS and PsA patients who are switched to ErelziTM from stable treatment with Enbrel.

• To assess patient satisfaction at baseline and after switching to ErelziTM.

• To explore what are the factors related (clinical, diagnostic factors, comorbidities, patient therapy history, experience/expertise with biologics, patient-oriented program, clinic location, extent of physician's knowledge of biosimilars) to the physician decision to switch.

#### **Study status**

Finalised

# Research institutions and networks

## Institutions

## Sandoz

First published: 01/02/2024

Last updated: 01/02/2024

Institution

# Contact details

### **Study institution contact**

placeholder placeholder placeholder@placeholder.com

Study contact

placeholder@placeholder.com

Primary lead investigator placeholder placeholder

Primary lead investigator

# Study timelines

Date when funding contract was signed Planned: 30/06/2017

Actual: 14/09/2017

Study start date Planned: 15/02/2019 Actual: 08/02/2019

Data analysis start date Actual: 31/05/2024

Date of final study report Planned: 31/08/2023 Actual: 13/12/2024

# Sources of funding

• Pharmaceutical company and other private sector

## More details on funding

SANDOZ FR

# Regulatory

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

No

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

Not applicable

# Methodological aspects

# Study type

# Study type list

### Study type:

Non-interventional study

### Scope of the study:

Effectiveness study (incl. comparative)

#### Main study objective:

To evaluate effectiveness of ErelziTM in patients with RA, AS or PsA with, stable clinical response to Enbrel®.

# Study Design

#### Non-interventional study design

Other

#### Non-interventional study design, other

National, Multicenter, Prospective, Post-Authorization, non-PASS, Real Life, Observational Study.

# Study drug and medical condition

### Name of medicine ERELZI

## Population studied

#### Age groups

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)

#### Estimated number of subjects

650

# Study design details

#### Data analysis plan

No minimum number of patients in each disease indication will be required.

## Data management

## 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**

Unknown

## Data characterisation

#### **Data characterisation conducted**

No