# Surveillance of Efficacy and Tolerability in Von Willebrand's disease

Submission date	Recruitment status	Prospectively registered
18/11/2008	No longer recruiting	☐ Protocol
Registration date	Overall study status	Statistical analysis plan
19/11/2008	Completed	Results
Last Edited	Condition category	Individual participant data
07/04/2011	Haematological Disorders	<ul><li>Record updated in last year</li></ul>

# Plain English summary of protocol

Not provided at time of registration

# Contact information

# Type(s)

Scientific

#### Contact name

Ms Jennifer Feddern

#### Contact details

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# Additional identifiers

Protocol serial number

**WIL-15** 

# Study information

Scientific Title

# Acronym

Wilate-SET

## Study objectives

The efficacy of Wilate® in clinical practice is comparable to the efficacy in clinical trial patients.

As of 07/05/2011 the anticipated end date for this trial has been extended from 31/07/2009 to 31/07/2012

## Ethics approval required

Old ethics approval format

## Ethics approval(s)

As the procedures during this observational study do not interfere with the patient's usual treatment and monitoring of treatment, this study is not regarded as a clinical study as defined by EU Directive 2001/20/EC. Therefore approval by an Independent Ethical Committee as an Institutional Review Board is not required.

## Study design

Non-interventional, observational, open, prospective, multi-centre study

# Primary study design

Observational

# Study type(s)

Treatment

# Health condition(s) or problem(s) studied

Von Willebrand's disease

#### **Interventions**

Documentation of details of diagnosis and bleeding frequency at entry, details of bleeding episodes and surgical procedures during 2-year individual observation period. Documentation of details of each injection including assessment of efficacy and tolerability at a 4-point verbal rating scale. Documentation of relevant laboratory assessments if performed.

#### Intervention Type

Drug

#### Phase

**Not Specified** 

## Drug/device/biological/vaccine name(s)

Wilate®

#### Primary outcome(s)

Percentage of efficacy assessments with rating "excellent/good".

## Key secondary outcome(s))

Percentage of tolerability assessments with rating "excellent/good".

#### Completion date

31/07/2012

# **Eligibility**

# Key inclusion criteria

Patients (any age and gender) suffering from hereditary or acquired Von Willebrand's disease requiring von Willebrand factor (VWF)/coagulation factor eight (FVIII) concentrate.

# Participant type(s)

**Patient** 

# Healthy volunteers allowed

No

#### Age group

Other

#### Sex

All

#### Key exclusion criteria

Does not comply with the above inclusion criteria

#### Date of first enrolment

01/02/2005

## Date of final enrolment

31/07/2012

# Locations

#### Countries of recruitment

Germany

# Study participating centre Octapharma GmbH

Langenfeld Germany 40764

# Sponsor information

#### Organisation

Octapharma GmbH (Germany)

#### **ROR**

https://ror.org/002k5fe57

# Funder(s)

# Funder type

Industry

#### Funder Name

Octapharma GmbH (Germany)

# **Results and Publications**

Individual participant data (IPD) sharing plan

# IPD sharing plan summary

Not provided at time of registration

# **Study outputs**

Output type Details Date created Date added Peer reviewed? Patient-facing?

Participant information sheet Participant information sheet 11/11/2025 11/11/2025 No Yes