

# Active post-marketing surveillance of Levonorgestrel IUS insertion related difficulties: a non-interventional post-authorisation safety study

**First published:** 12/11/2014

**Last updated:** 23/04/2024

Study

Finalised

## Administrative details

### EU PAS number

EUPAS7857

### Study ID

26348

### DARWIN EU® study

No

### Study countries

☐ Bulgaria

☐ Czechia

☐ Denmark

- ☐ Hungary
  - ☐ Lithuania
  - ☐ Norway
  - ☐ Poland
  - ☐ United Kingdom
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### Study description

The PASS STUDY is an observational study with primary objective to characterize, under routine practice, the ease of insertion and the safety profile of Levonorgestrel IUS during insertion in a study population that is representative of the actual users of the IUS (either new user and patients switching from Mirena). And with secondary objective to characterize the utilization pattern for Levonorgestrel IUS. The Data Clarification Form (DCF) will be included in the Levonorgestrel IUS package. The doctors will be asked to complete and send it to the sponsor on spontaneous basis.

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### Study status

Finalised

## Research institutions and networks

### Institutions

**Gedeon Richter**

**First published:** 01/02/2024

**Last updated:** 01/02/2024

**Institution**

Multiple centres: 200 centres are involved in the study

## Study timelines

### **Date when funding contract was signed**

Planned: 15/10/2014

Actual: 15/10/2014

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### **Study start date**

Planned: 16/10/2014

Actual: 16/10/2014

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

Planned: 21/12/2020

Actual: 28/08/2020

## Sources of funding

- Pharmaceutical company and other private sector

## More details on funding

Gedeon Richter Plc

## Study protocol

[Levosert Pass Study Amend 02 140618.pdf](#)(602.49 KB)

[Levosert PASS Protocol Amendment\\_5 2.pdf](#)(3.66 MB)

## 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

Disease /health condition

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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

Safety study (incl. comparative)

**Data collection methods:**

Primary data collection

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**Main study objective:**

The primary objective of this PASS is to characterize, under routine practice, the ease of insertion and the safety profile of Levonorgestrel IUS during insertion in a study population that is representative of the actual users of the IUS (either new users and patients switching from Mirena).

## Study Design

**Non-interventional study design**

Other

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**Non-interventional study design, other**

Multinational, multi-centre, PASS study

## Study drug and medical condition

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

LEVONORGESTREL

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**Medical condition to be studied**

Abnormal uterine bleeding

## Population studied

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

The study focused on women who had prescribed with levonorgestrel intrauterine system (IUS) for any indication as a part of routine care.

Inclusion criteria:

1. Willing and able to provide written informed consent
2. Prescribed levonorgestrel IUS as part of routine clinical care prior to enrolment.

Exclusion criteria:

1. There are no exclusion criteria for this study.
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## **Age groups**

Adults (18 to < 46 years)

Adults (46 to < 65 years)

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## **Estimated number of subjects**

1000

# Study design details

## **Outcomes**

IUS insertion related adverse events, The indication of use of levonorgestrel IUS

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## **Data analysis plan**

DCF information will be entered in the Clinical and, partially, in the Pharmacovigilance databases. Details of treatment initiation and prescribing reasons as reported on the questionnaire will be provided using descriptive statistics. Prescribing pattern will be assessed. The reported insertion problems will be provided in a listing which will also be part of the Final Clinical Study Report. General comments about the readability of the instructions for use and

handling will be recorded for potential future improvements.

## Documents

### Study publications

[Velev R.; A multiple center, randomised, parallel group, single-blind clinical ...](#)  
[Trinh XB, Talma WA, Makar AP, Buytaert G, Weyler J, van Dam PA, Use of the levo...](#)

[Mawet M, Nollevaux F, Nizet D, Wijzen F, Gordenne V, Tasev N, Segedi D, Marines...](#)

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## 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](#)

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### 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

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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

No