# Monitoring of pregnancy outcomes in women treated with inclisiran: a noninterventional study

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# Administrative details

#### **EU PAS number**

EUPAS42905

#### **Study ID**

50706

#### DARWIN EU® study

No

#### **Study countries**

Switzerland

## **Study description**

This post-authorisation safety study is a worldwide, single-arm descriptive noninterventional study that collects prospective and retrospective data in women exposed to inclisiran during pregnancy. It uses an enhanced pharmacovigilance data collection and processing system via a set of targeted checklists, structured follow-up, rigorous process of data entry and data quality control, and programmed aggregate analysis. The overall objective of the study is to collect data on pregnancy outcomes in patients treated with inclisiran during or prior to pregnancy. Although pharmacovigilance data may be collected from any country in the world where the product is approved, the anonymized patient level data will be analyzed at a global level in Switzerland.

## **Study status**

Ongoing

# Research institutions and networks

# Institutions

# **Novartis Pharmaceuticals**

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Institution

# Contact details

**Study institution contact** 

# Novartis Clinical Disclosure Officer Trialandresults.registries@novartis.com

Study contact

Trialandresults.registries@novartis.com

Primary lead investigator Novartis Clinical Disclosure Officer

Primary lead investigator

# Study timelines

Date when funding contract was signed Planned: 09/12/2020 Actual: 09/12/2020

**Study start date** Planned: 31/12/2021 Actual: 31/12/2021

Data analysis start date Planned: 31/01/2031

**Date of final study report** Planned: 31/07/2031

# Sources of funding

• Pharmaceutical company and other private sector

# More details on funding

Novartis Pharma AG

# Study protocol

CKJX839A12011-v00--protocol\_Redacted.pdf(726.72 KB)

CKJX839A12011\_Protocol V2\_Redacted.pdf(593.52 KB)

# Regulatory

Was the study required by a regulatory body? Yes

Is the study required by a Risk Management Plan (RMP)? EU RMP category 3 (required)

# Other study registration identification numbers and links

CKJX839A12011

Methodological aspects

Study type

Study type list

## Study type:

Non-interventional study

## Scope of the study:

Safety study (incl. comparative)

## Main study objective:

To estimate the proportion of major congenital malformations among pregnancies exposed to inclisiran during pregnancy prospectively reported to Novartis amongst (i) live births and (ii) live births plus still births plus termination of pregnancy for fetal anomaly (TOPFA).

# Study Design

**Non-interventional study design** Cohort Other

## Non-interventional study design, other

Intensive monitoring schemes

# Study drug and medical condition

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

## Anatomical Therapeutic Chemical (ATC) code (C10AX16) inclisiran inclisiran

# **Population studied**

#### Age groups

Adolescents (12 to < 18 years) Adults (18 to < 46 years) Adults (46 to < 65 years)

## Special population of interest

Pregnant women

## **Estimated number of subjects**

500

# Study design details

#### Outcomes

Major congenital malformations (MCM) among pregnancies exposed to inclisiran during pregnancy, • To estimate the proportion of MCM among pregnancies exposed to inclisiran prior to the last menstrual period (LMP) • To compare the frequency of MCM among pregnancies exposed to inclisiran with the background frequencies • To estimate the proportion of other secondary outcomes in pregnancies exposed to inclisiran prior to LMP and during pregnancy

#### Data analysis plan

Data analysis will focus on the prospectively reported pregnancy cases. Retrospective pregnancy cases will also be analyzed and presented separately from the prospective cases. Data analysis will include the estimation of proportion (and 95% confidence interval) of malformations (major, minor, and overall), and of specific pregnancy outcomes such as live births, stillbirths, spontaneous abortions and elective terminations. Proportions will be estimated by timing of drug exposure in pregnancy (during pregnancy and prior to LMP, as well as by trimester if sample size allows). Descriptive analysis will be performed for all prospective pregnancy cases including case disposition (outcome known, pending, and lost to follow-up) and maternal characteristics (i.e. age, ethnicity, region) by providing the number and percentage of pregnancies in each category. Further stratified analyses may be undertaken if sample size allows.

# 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

Spontaneous reports of suspected adverse drug reactions

#### Data sources (types), other

targeted follow-up questionnaires for reported pregnancy cases

# 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

Not applicable