# The Benralizumab Pregnancy Exposure Study: A VAMPSS Post-Marketing Surveillance Study

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## Administrative details

EU PAS number
EUPAS26461
Study ID
45903
DARWIN EU® study
No
Study countries
Canada
United States

## **Study status**

**Finalised** 

Research institutions and networks

## Institutions

# Organization of Teratology Information Specialists (OTIS)

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# University California SD

## **Networks**

# Organization of Teratology Information Specialists (OTIS) Network

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## Contact details

## **Study institution contact**

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**Study contact** 

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### Primary lead investigator

## Christina Chambers

**Primary lead investigator** 

# Study timelines

### Date when funding contract was signed

Planned: 20/12/2017

Actual: 20/12/2017

#### Study start date

Planned: 01/09/2018

Actual: 20/03/2019

#### Data analysis start date

Planned: 01/11/2025

Actual: 26/12/2023

### Date of interim report, if expected

Planned: 15/12/2019

Actual: 12/12/2018

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

Planned: 30/03/2027

Actual: 23/08/2024

# Sources of funding

Pharmaceutical company and other private sector

## More details on funding

AstraZeneca AB

# Study protocol

d3250r00026-csp-v2\_Redacted\_15Nov2018.pdf(1.26 MB)

d3250r00026-csp-v3 Redacted 27Jul2020(final version).pdf(491.03 KB)

# Regulatory

Was the study required by a regulatory body?

No

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

EU RMP category 3 (required)

# Other study registration identification numbers and links

D3250R00026

# Methodological aspects

Study type

Study type list

#### Study type:

Non-interventional study

#### Scope of the study:

Assessment of risk minimisation measure implementation or effectiveness

#### Main study objective:

To monitor planned and unplanned pregnancies exposed to benralizumab and evaluate the potential teratogenic effect of this medication relative to the primary pregnancy outcome of major structural birth defects and the secondary pregnancy outcomes of preterm delivery, small for gestational age infants, spontaneous abortion, stillbirth, elective termination and postnatal growth to one year of age.

## Study Design

#### Non-interventional study design

Other

#### Non-interventional study design, other

This is a prospective, observational, exposure cohort study of pregnancy outcomes in pregnant women with asthma exposed to benralizumab and comparison women

# Study drug and medical condition

#### Name of medicine, other

FasenraTM

#### Medical condition to be studied

Asthma

# Population studied

#### Age groups

Preterm newborn infants (0 - 27 days)

Term newborn infants (0 – 27 days)

Infants and toddlers (28 days - 23 months)

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**

800

## Study design details

#### **Outcomes**

Major structural birth defects, - Spontaneous abortion/miscarriage - Stillbirth - Elective termination/abortion - Preterm delivery - Small for gestational age infants - Small for age postnatal growth of live born children to 1 year of age

#### **Data analysis plan**

Descriptive tables will be prepared for characteristics of each of the cohorts in each interim and final report displaying n, means, standard deviations, minimums and maximums or proportions and percentages. For the primary endpoint and for the secondary endpoints un-adjusted relative risk estimates will be presented together with exact two-sided 95% and 80% confidence intervals. For the secondary endpoints survival methods will be used (Kaplan Meier) to estimate crude rates and confidence intervals accounting for gestational timing of enrollment in the study. Adjusted analyses producing relative risks and 95% and 80% confidence intervals or hazard ratios and 95% and 80% confidence intervals, where numbers permit, will be conducted. Confounders will be considered for each adjusted analysis separately, using several methods including the method of change in estimate of the effect of exposure to benralizumab by 10% or more.

## **Documents**

#### Study results

OTIS Benralizumab Final Analysis Report\_Synopsis\_Redacted.pdf(354.46 KB)

#### **Study report**

2019 FINAL OTIS BENRALIZUMAB ANNUAL INTERIM REPORT\_03Dec2019.pdf (384.32 KB)

2020 OTIS Benralizumab Pregnancy Registry Annual Report\_10Nov2020.pdf (450.17 KB)

2021 FINAL OTIS Fasenra Annual Interim Report\_08Feb2022.pdf(725.07 KB)

#### Study, other information

2020 OTIS Benralizumab Pregnancy Registry Annual Report\_10Nov2020.pdf (450.17 KB)

2021 FINAL OTIS Fasenra Annual Interim Report\_08Feb2022.pdf(725.07 KB)

## Data management

## Data sources

Other	
Data sources (types), other	
Prospective patient-based data collection, Medical records	
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