A Post-Authorization Safety Study to Evaluate the Safety of Abrocitinib Exposure During Pregnancy in United States Healthcare Databases

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## Administrative details

### **EU PAS number**

EUPAS100000096

#### **Study ID**

100000096

DARWIN EU® study

No

### **Study countries**

United States

## **Study description**

This US population-based, non-interventional, cohort study will evaluate the risk of adverse pregnancy and infant outcomes among women with moderate-tosevere AD who are exposed to abrocitinib during pregnancy compared to those unexposed during pregnancy. Pregnancies with start dates occurring during the period 14 January 2022 to Q4 2025, and infants born to the women, will be included. The main outcome of interest will be MCMs. Additional outcomes of interest include other adverse infant outcomes (preterm birth, SGA) and pregnancy outcomes (stillbirth, spontaneous abortion).

Study status

Ongoing

# Research institutions and networks

## Institutions

## Pfizer

First published: 01/02/2024

Last updated: 01/02/2024

Institution

## Harvard Pilgrim Health Care Institute

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# Contact details

## Study institution contact

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Study contact

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Primary lead investigator Jenny Sun Primary lead investigator

# Study timelines

### Date when funding contract was signed

Planned: 07/09/2022

Actual: 07/09/2022

## Study start date

Planned: 31/12/2022 Actual: 31/12/2022

Data analysis start date Planned: 01/03/2024 Actual: 17/04/2024

Date of final study report Planned: 30/06/2028

## Sources of funding

• Pharmaceutical company and other private sector

## More details on funding

Pfizer

# Study protocol

B7451098\_ABROCITINIB UPDATED PROTOCOL\_18NOV2022\_Redacted.pdf(2.88 MB)

# Regulatory

## Was the study required by a regulatory body?

Yes

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:

Safety study (incl. comparative)

## Data collection methods:

Secondary use of data

### Study design:

This US population-based, non-interventional, cohort study will evaluate the risk of adverse pregnancy and infant outcomes among women with moderate-tosevere AD who are exposed to abrocitinib during pregnancy compared to those unexposed during pregnancy.

## Main study objective:

To describe and compare the adverse pregnancy and infant outcomes among women with moderate-to-severe AD who are exposed to abrocitinib during pregnancy and women with moderate-to-severe AD who are not exposed to abrocitinib.

# Study Design

## Non-interventional study design

Cohort

# Study drug and medical condition

## Name of medicine CIBINQO

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

## Anatomical Therapeutic Chemical (ATC) code

(D11AH08) abrocitinib abrocitinib

### Additional medical condition(s)

Moderate-to-severe atopic dermatitis (AD)

# **Population studied**

### Short description of the study population

The study population will include singleton pregnancies among women 12 to 49 years of age diagnosed with moderate-to-severe AD who are members of participating Sentinel Data Partners (DPs).

#### Age groups

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

### **Special population of interest**

Pregnant women

# Study design details

### Setting

The study will be conducted using data provided by US Data Partners in the FDA's Sentinel System. The study population will include singleton pregnancies among women 12 to 49 years of age diagnosed with moderate-to-severe AD who are members of participating Sentinel Data Partners (DPs).

### Outcomes

The main outcome of interest will be MCMs. Additional outcomes of interest include other adverse infant outcomes (preterm birth, SGA) and pregnancy outcomes (stillbirth, spontaneous abortion).

## Data management

## Data sources

## Data source(s), other

Sentinel Data Partners

### Data sources (types)

Administrative healthcare records (e.g., claims)

# Use of a Common Data Model (CDM)

## **CDM mapping**

Yes

**CDM Mappings** 

#### **CDM name**

Sentinel

### **CDM** website

https://www.sentinelinitiative.org/methods-Data-tools/sentinel-common-Datamodel

## Data quality specifications

#### **Check conformance**

Unknown

### **Check completeness**

Unknown

## **Check stability**

Unknown

### **Check logical consistency**

Unknown

## Data characterisation

### **Data characterisation conducted**

Unknown