

# Long-term safety study of Rinvoq in RA patients enrolled in the Corrona RA Registry in the United States

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Study

Ongoing

## Administrative details

### EU PAS number

EUPAS39194

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### Study ID

47971

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### DARWIN EU® study

No

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### Study countries

United States

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### Study description

The overall goal of the study is to characterize the safety of Rinvoq in RA patients in the post-approval setting. The primary objective is to compare the incidence of malignancy (excluding non-melanoma skin cancer, NMSC), NMSC, MACE, VTE, SIE, and mortality in adults with RA who receive Rinvoq relative to those who are treated with biologic medications approved for the treatment of RA, in the course of routine clinical care.

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### **Study status**

Ongoing

## Contact details

### **Study institution contact**

Clinical Trial Disclosure AbbVie [CT.Disclosures@abbvie.com](mailto:CT.Disclosures@abbvie.com)

**Study contact**

[CT.Disclosures@abbvie.com](mailto:CT.Disclosures@abbvie.com)

### **Primary lead investigator**

Clinical Trial Disclosure AbbVie

**Primary lead investigator**

## Study timelines

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

Planned: 09/01/2020

Actual: 09/01/2020

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### **Study start date**

Planned: 16/08/2019

Actual: 09/08/2021

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## **Date of final study report**

Planned: 16/02/2030

## Sources of funding

- Pharmaceutical company and other private sector

## More details on funding

AbbVie

## Study protocol

[p19141-protocol abstract-pmos\\_Redacted.pdf](#) (97.25 KB)

[P19-141\\_protocol v3.0\\_abstract\\_Redacted.pdf](#) (154.98 KB)

## 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)

## Other study registration identification numbers and links

P19-141

Methodological aspects

**Study type:**

Non-interventional study

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**Scope of the study:**

Assessment of risk minimisation measure implementation or effectiveness

**Main study objective:**

The main objective is to compare the incidence of malignancy (excluding non-melanoma skin cancer, NMSC), NMSC, MACE, VTE, SIE, and mortality in adults with RA who receive Rinvog relative to those who are treated with biologic medications approved for the treatment of RA, in the course of routine clinical care.

## Study Design

**Non-interventional study design**

Cohort

## Study drug and medical condition

**Medicinal product name**

RINVOQ

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**Medical condition to be studied**

Rheumatoid arthritis

## Population studied

## **Age groups**

- Adults (18 to < 46 years)
  - Adults (46 to < 65 years)
  - Adults (65 to < 75 years)
  - Adults (75 to < 85 years)
  - Adults (85 years and over)
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## **Estimated number of subjects**

7500

# Study design details

## **Outcomes**

To compare the incidence of malignancy (excluding non-melanoma skin cancer, NMSC), NMSC, MACE, VTE, SIE, and mortality in adults with RA who receive Rinvoq relative to those who are treated with biologic medications approved for the treatment of RA, in the course of routine clinical care. To describe the incidence rates of the following adverse events: herpes zoster (HZ), opportunistic infections (OI), active tuberculosis (TB), gastrointestinal (GI) perforations, and evidence of drug-induced liver injury (DILI) in adults with RA who receive Rinvoq relative to those who are treated with biologic medications approved for the treatment of RA, in the course of routine clinical care.

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## **Data analysis plan**

The population of patients in the study will be characterized with respect to demographic, clinical, disease and patient-reported outcomes using descriptive statistics. The main analyses will employ propensity score methods to address confounding by indication (channeling bias) followed by estimation of incident rates as well as multivariable Cox proportional hazards modeling to estimate

HRs and associated 95% CIs.

## 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)

[Disease registry](#)

## 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