

# id.DRIVE study of brand-specific respiratory syncytial virus (RSV) vaccine effectiveness in Europe

**First published:** 01/03/2024

**Last updated:** 01/10/2025

Study

Ongoing

## Administrative details

### EU PAS number

EUPAS1000000035

### Study ID

1000000035

### DARWIN EU® study

No

### Study countries

- France
- Germany
- Italy
- Spain

United Kingdom

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### Study description

This is a European, non-interventional study to estimate the effectiveness of vaccines against respiratory syncytial virus (RSV). Of interest is the effectiveness against hospitalisation with severe acute respiratory infection (SARI) caused by RSV.

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### Study status

Ongoing

## Research institutions and networks

### Institutions

#### P95 Clinical and Epidemiology Services

- Belgium
- Colombia
- Netherlands
- South Africa
- Thailand
- United States

**First published:** 07/11/2022

**Last updated:** 21/02/2025

**Institution**

**Laboratory/Research/Testing facility**

**Non-Pharmaceutical company**

**ENCePP partner**

# The Foundation for the Promotion of Health and Biomedical Research of Valencia Region (FISABIO)

Spain

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

**Last updated:** 31/10/2025

**Institution**

## University Hospital Vall d'Hebron (HUVH)

Spain

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

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

**Institution**

**Educational Institution**

**Hospital/Clinic/Other health care facility**

## Imperial College London

United Kingdom

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

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

**Institution**

**Educational Institution**

Valencia Hospital Network for the Study of  
Influenza and Other Respiratory Viruses (VAHNSI)  
Germans Trias i Pujol University Hospital  
Hospital Clínico Universitario de Santiago de  
Compostela  
Hospital Universitario La Paz

Centro Interuniversitario per la Ricerca  
sull’Influenza e le altre Infezioni (CIRI-IT)

Universitätsklinikum Ulm  
Charité - Universitätsmedizin Berlin

Hôpitaux Universitaires Henri-Mondor  
Centre Hospitalier Annecy Genevois

Manchester University NHS Foundation Trust  
(MFT)  
NHS Lothian

# Networks

## [id.DRIVE \(former COVIDRIVE\)](#)

- Austria
- Belgium
- Croatia
- Czechia
- Denmark
- France
- Germany
- Iceland
- Italy
- Poland
- Romania
- Spain

**First published:** 06/09/2021

**Last updated:** 29/02/2024

**Network**

**ENCePP partner**

# Contact details

## **Study institution contact**

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**Study contact**

[kaatje.bollaerts@p-95.com](mailto:kaatje.bollaerts@p-95.com)

## Primary lead investigator

Kaatje Bollaerts

**Primary lead investigator**

## Study timelines

### Date when funding contract was signed

Planned: 15/03/2024

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### Study start date

Planned: 01/06/2024

Actual: 29/08/2024

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### Date of final study report

Planned: 01/12/2026

## Sources of funding

- Pharmaceutical company and other private sector

## More details on funding

Pfizer

## Study protocol

[RSV\\_Master Protocol V2.0\\_07 Jun 2024.pdf \(994.21 KB\)](#)

[RSV Vaccine Effectiveness protocol v3.0\\_21 Feb 2025.pdf \(856.05 KB\)](#)

## Regulatory

**Was the study required by a regulatory body?**

No

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

Not applicable

**Other study registration identification numbers and links**

<https://iddrive.eu>

## Methodological aspects

**Study type**

**Study type list**

**Study topic:**

Disease /health condition

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**Study type:**

Non-interventional study

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**Scope of the study:**

Effectiveness study (incl. comparative)

**Data collection methods:**

Combined primary data collection and secondary use of data

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### **Study design:**

This study is a multi-country, multi-centre, hospital-based case-control study with test-negative controls (TNCC design). A combination of primary and secondary data collection will be used to obtain the relevant data.

### **Main study objective:**

To estimate brand-specific RSV vaccine effectiveness (VE) against hospitalisation due to laboratory-confirmed RSV infection in severe acute respiratory infection (modified SARI) older adult patients.

## Study Design

### **Non-interventional study design**

Case-control

## Study drug and medical condition

### **Medicinal product name**

ABRYSVO

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### **Medicinal product name, other**

Respiratory syncytial virus vaccine (bivalent, recombinant)

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### **Anatomical Therapeutic Chemical (ATC) code**

(J07BX05) respiratory syncytial virus vaccines  
respiratory syncytial virus vaccines

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## **Medical condition to be studied**

Severe acute respiratory syndrome

## **Population studied**

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

The study population consists of hospitalised "modified severe acute respiratory infection (SARI)"\*  $\geq 60$  year-old patients.

A "modified SARI" patient is a person hospitalised primarily for a suspicion of a respiratory infection with at least one of the following symptoms:

- cough,
- shortness of breath,
- fever ( $\geq 38$  C°)

with symptom onset within the last 10 days prior to hospital admission.

\*This SARI definition is modified from the latest European Centre for Disease Prevention and Control (ECDC) case definition [EUROPEAN CENTRE FOR DISEASE PREVENTION AND CONTROL (2021). ECDC Technical report: Core protocol for ECDC studies of COVID-19 vaccine effectiveness against hospitalisation with Severe Acute Respiratory Infection laboratory-confirmed with SARS-CoV-2, version 1.0, 2021].

## **Study design details**

### **Setting**

Hospitals

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### **Outcomes**

The outcome of interest for the primary analysis will be RSV detection in patients hospitalised with symptoms consistent with the modified ECDC SARI definition.

RSV infection must be laboratory-confirmed by reverse transcription polymerase chain reaction (RT-PCR) or another ribonucleic acid (RNA) amplification system with at least the same sensitivity as RT-PCR (e.g., transcription-mediated amplification [TMA]). RT-PCR assays must be used for the detection of at least RSV, SARS-CoV-2, and influenza.

The secondary outcomes in patients hospitalised with modified SARI symptoms include:

- detection of RSV subtypes A and B
  - detection of RSV genotypes
  - modified SARI severity level of study participants (both cases and controls), including respiratory support severity level
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## **Data analysis plan**

A SAP is developed prior to the conduct of the analysis. The SAP specifies all statistical analyses conducted, and includes tables shells and mock figures.

## Documents

[Link to id.DRIVE.eu](#)

## 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.

## Use of a Common Data Model (CDM)

### **CDM mapping**

No

## Data quality specifications

### **Check conformance**

Yes

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### **Check completeness**

Yes

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### **Check stability**

Yes

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### **Check logical consistency**

Yes

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## Data characterisation

### **Data characterisation conducted**

Yes

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### **Data characterisation moment**

after data extraction