

# SARS-CoV-2 VACCINATION for COVID-19 DISEASE SAFETY STUDY (VAC4COVID STUDY)

**First published:** 19/11/2021

**Last updated:** 17/05/2024

Study

Ongoing

## Administrative details

### EU PAS number

EUPAS44124

### Study ID

44381

### DARWIN EU® study

No

### Study countries

☐ United Kingdom

## **Study description**

VAC4COVID is an online study created by MEMO (Medicines Monitoring Unit) Research at the University of Dundee to track all authorised vaccines used to prevent COVID-19 infection. All licensed/authorised vaccines have already been tested extensively for effectiveness and safety in large clinical trials. As with every new medicine, governments and researchers need to keep monitoring how COVID-19 vaccines are working once they are being used in the wider population. This is standard practice and an important part of how medicines and vaccines are regulated. This study will track vaccines to check they are working well. It will be big enough to detect any rare unexpected vaccine side effects. Participation in this study is voluntary and it will not interfere with participants' usual health care. The aim of the study is to monitor the effectiveness and safety of COVID-19 vaccines by gathering information from a large number of participants about their health before and after they are vaccinated. Participants are prompted by email at monthly intervals to state whether they have had any changes in their health and wellbeing. They are also asked if any reported events required emergency care or resulted in hospitalisation. For the four weeks following vaccination, they are asked to provide these updates on their health on a weekly basis, then return to monthly follow-ups for at least 11 months. Participants are asked to provide consent to allow follow-up and investigation of any adverse events they may experience following vaccination against COVID-19.

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## **Study status**

Ongoing

## **Research institutions and networks**

### **Institutions**

# University of Dundee

☐ United Kingdom

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

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Institution

Educational Institution

## Contact details

### Study institution contact

Thomas MacDonald t.m.macdonald@dundee.ac.uk

Study contact

[t.m.macdonald@dundee.ac.uk](mailto:t.m.macdonald@dundee.ac.uk)

### Primary lead investigator

Thomas MacDonald

Primary lead investigator

## Study timelines

### Date when funding contract was signed

Planned: 16/12/2020

Actual: 16/12/2020

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### Study start date

Planned: 28/01/2021

Actual: 02/02/2021

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

Planned: 31/12/2025

## Sources of funding

- Other

## More details on funding

University of Dundee

## 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

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## Other study registration identification numbers and links

ISRCTN -ISRCTN95881792 <https://doi.org/10.1186/ISRCTN95881792>

## Methodological aspects

### Study type

### Study type list

**Study type:**

Non-interventional study

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**Scope of the study:**

Drug utilisation

Safety study (incl. comparative)

**Main study objective:**

To examine the use and safety of licensed/authorised COVID-19 vaccination by collecting event data from participants and healthcare professionals before, during and after the vaccination programme. To monitor, detect and report early pre-specified and emergent safety signals in relation to licensed/authorised COVID-19 vaccinations.

## Study Design

**Non-interventional study design**

Cohort

Other

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**Non-interventional study design, other**

Prescription event monitoring, post-authorisation cohort safety study (PASS)

## Population studied

**Age groups**

Children (2 to < 12 years)

Adolescents (12 to < 18 years)

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**

20000

## Study design details

### **Outcomes**

Recruit consented cohorts of participants to examine the safety of licensed/authorised COVID-19 vaccinations. Outcomes measures: Number of participants offered vaccination Number of participants vaccinated Numbers and rates of SAEs and AESI occurring in vaccinated and non-vaccinated participants, Collect participant self-reported data via questionnaires using electronic data capture. Collect demographics of vaccinated and unvaccinated (unexposed) participants. Collect demographics of vaccinated and unvaccinated (unexposed) participants. Collate and analyse licensed/authorised vaccine utilisation characteristics for the whole vaccinated cohort.

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### **Data analysis plan**

We will conduct frequent interim analyses (at intervals to be determined in collaboration with medicines safety regulators) to detect signals in the data as it accumulates (Phase 1), and further, more detailed analyses when the study is complete (Phase 2). For each primary outcome we will plot cumulative hazards (-log(survival rate), with confidence intervals) for: vaccinated patients, with date of vaccination as time 0, and unvaccinated patients, with date of recruitment as time 0 and censored at vaccination. We will also plot the

cumulative hazards based on an external estimate of population incidence rates (a straight line).

## Documents

### Study, other information

[VAC4COVID Advisory Committee Charter v1 \(e-signed 22-02-21\).pdf](#)(94.33 KB)

## 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](#)

[Drug dispensing/prescription data](#)

[Electronic healthcare records \(EHR\)](#)

[Other](#)

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### Data sources (types), other

Prospective patient-based data collection

## 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