

# DARWIN EU® - Coverage of meningococcal vaccines in the target population in Europe

**First published:** 15/07/2025

**Last updated:** 23/01/2026

Study

Finalised

## Administrative details

### EU PAS number

EUPAS1000000675

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### Study ID

1000000675

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### DARWIN EU® study

Yes

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### Study countries

-  Croatia
  -  Denmark
  -  Finland
  -  Germany
  -  Spain
  -  United Kingdom
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## **Study description**

Vaccine exposure data supports readiness for safety signal management and can inform the feasibility of vaccine effectiveness or safety studies. However, coverage data is often mostly available from MAH sales/distribution data, which does not accurately reflect actual use. Coverage data for meningococcal vaccines, which are recommended to prevent invasive meningococcal disease in children and adolescents, is scarce and of limited granularity.

This population-level drug utilisation study generated evidence on temporal trends in the coverage of meningococcal vaccines, by brand and/or vaccine composition (as available in the data sources), and by age group and dose, in 6 data sources in 5 countries (Croatia, Spain, UK, Denmark, Finland). Exposure included: MenB vaccines (Bexsero® and Trumenba®); MenC and Haemophilus influenzae type b (Hib)/MenC conjugate vaccines; MCV4 vaccines (serogroups A, C, W-135, and Y: Menveo® and Nimenrix®). As expected, countries with universal meningococcal vaccination in childhood (ES, UK) showed higher and more consistent coverage than those without this recommendation (DK, FI, HR), reflecting the different immunisation policies across Europe. Recent decrease in coverage observed in some countries may be related to the impact of the COVID-19 pandemic on vaccine availability and access to vaccination, and increased recent increasing public vaccine hesitancy.

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## **Study status**

Finalised

## **Research institutions and networks**

### **Institutions**

# Department of Medical Informatics - Health Data Science, Erasmus Medical Center (ErasmusMC)

 Netherlands

**First published:** 03/11/2022

**Last updated:** 02/05/2024

**Institution**

**Educational Institution**

**ENCePP partner**

## Networks

### Data Analysis and Real World Interrogation Network (DARWIN EU®)

 Belgium

 Croatia

 Denmark

 Estonia

 Finland

 France

 Germany

 Greece

 Hungary

 Italy

 Netherlands

 Norway

 Portugal

 Spain

 Sweden

 United Kingdom

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

**Last updated:** 30/04/2025

Network

## Contact details

### Study institution contact

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Study contact

[study@darwin-eu.org](mailto:study@darwin-eu.org)

### Primary lead investigator

Ivan Lam

Primary lead investigator

## Study timelines

### Date when funding contract was signed

Planned: 27/03/2025

Actual: 27/03/2025

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### Study start date

Planned: 30/06/2025

Actual: 30/06/2025

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

Planned: 15/11/2025

Actual: 10/12/2025

## Sources of funding

- EMA

## Study protocol

[DARWIN EU\\_Protocol\\_P4-C1-006\\_Utilisation of Meningococcal Vaccines in Europe\\_V3.0.pdf](#) (1.02 MB)

## Regulatory

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

Yes

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

Not applicable

## Methodological aspects

### Study type

### Study type list

**Study topic:**

Human medicinal product

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**Study type:**

Non-interventional study

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**Scope of the study:**

Drug utilisation

**Data collection methods:**

Secondary use of data

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**Study design:**

A drug utilisation cohort study will be conducted, using routinely collected health data from seven nationwide or regional databases in six European countries

**Main study objective:**

1. To examine the coverage of MenB vaccines in children at age one and two years by dose received ( $\geq 1$  dose,  $\geq 2$  doses,  $\geq 3$  doses, =1 dose, =2 doses, =3 doses)
2. To examine the coverage of MenC or Hib/MenC conjugate vaccines in children at age two years ( $\geq 1$  dose, =1 dose)
3. To examine the coverage of MCV4 vaccines in individuals at age 18 years ( $\geq 1$  dose, =1 dose)
4. To estimate the coverage of specific brand of MenB vaccines (Bexsero® and Trumemba®) in individuals aged two years and MCV4 vaccines (Menveo® and Nimenrix®) in individuals aged 18 years =
5. To characterise the age distribution of receipt of MenB, MenC, and MCV4 vaccines

## Study Design

## **Non-interventional study design**

Cohort

# Study drug and medical condition

## **Medicinal product name, other**

Monovalent Meningococcal group B surface protein vaccine, Monovalent Meningococcal group B surface protein vaccine, Quadrivalent Meningococcal conjugate vaccine (groups A, C, W-135, and Y)

# Population studied

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

This study will include all individuals present in their respective database who reach two years of age for examining the coverage of MenB or Hib/MenC vaccines, and 18 years of age for examining the coverage of MCV4 vaccines

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## **Age groups**

- **In utero**
- **Paediatric Population (< 18 years)**
  - Neonate
    - Preterm newborn infants (0 - 27 days)
    - Term newborn infants (0 - 27 days)
  - Infants and toddlers (28 days - 23 months)
  - Children (2 to < 12 years)
  - Adolescents (12 to < 18 years)
- **Adult and elderly population (≥18 years)**
  - Adults (18 to < 65 years)
    - Adults (18 to < 46 years)

- Adults (46 to < 65 years)
- Elderly ( $\geq$  65 years)
  - Adults (65 to < 75 years)
  - Adults (75 to < 85 years)
  - Adults (85 years and over)

## Study design details

### Data analysis plan

The point prevalence of recipients of vaccine coverage at quarterly and yearly time interval were estimated using the IncidencePrevalence package. The age distribution of vaccine recipients was characterised using the CohortCharacteristics package in R. Missing records for a given variable or outcome were assumed to indicate that the information was not present, or that a given individual was unvaccinated. Sensitivity analyses were conducted by 1) defining the study population as the target population for specific meningococcal vaccines without the requirement for complete follow-up and 2) defining the study population for the coverage of MCV4 as all individuals aged between 12 and 18 to test for the robustness of the findings and to examine the coverage of MCV4 in a broader population. A minimum cell count of 5 will be used when reporting results, with any smaller count reported as “<5” and zero counts as “0”.

## Documents

### Abstract of study report

[DARWIN EU\\_Report\\_P4-C1-006\\_Uptake of meningococcal vaccines\\_V3.0.pdf](#)

(2.25 MB)

## 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 source(s)

Croatia National Public Health Information System (Nacionalni javnozdravstveni informacijski sustav)

The Information System for Research in Primary Care (SIDIAP)

Danish Health Data Registries

Hospital District of Helsinki and Uusimaa patient cohort (FinOMOP)

BIFAP - Base de Datos para la Investigación Farmacoepidemiológica en el Ámbito Público (Pharmacoepidemiological Research Database for Public Health Systems)

Clinical Practice Research Datalink (CPRD) GOLD

## Use of a Common Data Model (CDM)

### CDM mapping

Yes

## CDM Mappings

### CDM name

OMOP

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### CDM website

<https://www.ohdsi.org/Data-standardization/>

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### CDM version

<https://ohdsi.github.io/CommonDataModel/index.html>

## 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

Unknown