# 205514 (V72\_380B) - Post-licensure observational effectiveness study of meningococcal B vaccine 4CMenB (Bexsero®) vaccination

**First published:** 28/07/2015

**Last updated:** 02/04/2024





## Administrative details

EU PAS number
EUPAS10416
Study ID
33821
DARWIN EU® study
No
Charles countries
Study countries
United Kingdom

#### Study description

The purpose of this study is to investigate the effectiveness of 4CMenB vaccination during routine clinical care in the UK national immunisation programme (NIP).

## **Study status**

Finalised

## Research institutions and networks

## Institutions

Public Health England (PHE)

## Contact details

## **Study institution contact**

Call Center EU Clinical Trials
Vx.publicdisclosureglobal@gsk.com

Study contact

Vx.publicdisclosureglobal@gsk.com

## Primary lead investigator

Call Center EU Clinical Trials

**Primary lead investigator** 

# Study timelines

#### Date when funding contract was signed

Actual: 02/11/2011

#### Study start date

Actual: 01/09/2015

#### **Data analysis start date**

Actual: 01/10/2015

#### Date of final study report

Actual: 24/05/2019

## Sources of funding

Other

# More details on funding

This surveillance was conducted by Public Health England. Post-marketing surveillance reports were provided to GSK to comply with their Risk Management Strategy.GSK provided funding for purchasing the reports and provided the MATS kits.See section 19

# Study protocol

V72\_38OB-04 Trial Registration Form-ENCePP Registration Redacted Protocol-2015-07-03.pdf(622.26 KB)

gsk-205514-protocol-redact.pdf(597.14 KB)

# Regulatory

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

Yes

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

EU RMP category 3 (required)

# Methodological aspects

# Study type

# Study type list

#### **Study topic:**

Human medicinal product

## **Study type:**

Non-interventional study

## Scope of the study:

Disease epidemiology

Effectiveness study (incl. comparative)

#### **Data collection methods:**

Primary data collection

## Main study objective:

The objective of this post-marketing observational study is to assess the impact on MenB and effectiveness of 4CMenB vaccination against MenB disease, after

# Study Design

#### Non-interventional study design

Other

#### Non-interventional study design, other

Descriptive study, Vaccine effectiveness (VE) will be assessed by the screening method, or by a case-control method if the screening method cannot be used (for example, if appropriate coverage data cannot be determined)

## Study drug and medical condition

#### Name of medicine

**BEXSERO** 

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

RECOMBINANT NEISSERIA MENINGITIDIS GROUP B NHBA FUSION PROTEIN
RECOMBINANT NEISSERIA MENINGITIDIS GROUP B NADA PROTEIN
RECOMBINANT NEISSERIA MENINGITIDIS GROUP B FHBP FUSION PROTEIN
PRODUCED IN E. COLI CELLS BY RECOMBINANT DNA TECHNOLOGY ADSORBED
ON ALUMINIUM HYDROXIDE

OUTER MEMBRANE VESICLES FROM NEISSERIA MENINGITIDIS GROUP B (STRAIN NZ 98/254)

**NEISSERIA MENINGITIDIS** 

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

(J07AH09) meningococcus B, multicomponent vaccine

# Population studied

#### Short description of the study population

General population in England.

Individuals were included in the cohorts targeted for vaccination in England.

## **Age groups**

Infants and toddlers (28 days - 23 months)

#### **Estimated number of subjects**

1

# Study design details

#### **Outcomes**

The primary outcome is a capsular group B confirmed case by culture and/or PCR from a normally sterile site (case definition A), regardless of MATS, The secondary outcome is a confirmed or probable case of capsular group B 4CMenB-vaccine-type where protection would have been expected based on the vaccine antigens (case definition B)

#### **Data analysis plan**

 $VE = 1 - (PCV \times (1-PPV)) / ((1-PCV) \times PPV)VE$ : Vaccine EffectivenessPCV:

Proportion Cases Vaccinated PPV: Proportion Population Vaccinated

## **Documents**

## Study, other information

Sources of funding information.pdf(70.55 KB)

## **Study publications**

Ladhani SN, Andrews N, Parikh SR, Campbell H, White J, Edelstein M, Bai X, Luci...

# Data management

## Data sources

#### **Data sources (types)**

Other

## 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

## **Check completeness**

Unknown

## **Check stability**

Unknown

## **Check logical consistency**

Unknown

# Data characterisation

## **Data characterisation conducted**

Unknown