Be on the TEAM: Teenagers Against Meningitis

Submission date 12/03/2018	Recruitment status No longer recruiting	[X] Prospectively registered [X] Protocol
Registration date 14/03/2018	Overall study status Completed	 Statistical analysis plan Results
Last Edited 18/06/2025	Condition category Infections and Infestations	Individual participant data[X] Record updated in last year

Plain English summary of protocol

Background and study aims

Teenagers and young children are at increased risk of diseases such as meningitis and blood poisoning due to bacteria called meningococcus. Although these diseases can be serious, the meningococcus bacteria are carried in the back of the throat of 1 in 10 teenagers without causing any symptoms. Most meningococcal disease in teenagers is due to Meningitis B (also known as MenB). The aim of this study is to see whether immunising teenagers with vaccines against MenB can reduce the number of teenagers carrying these bacteria in their throat. This would be important because it could mean that teenage MenB immunisation would not only help protect teenagers against these potentially deadly diseases, but also that babies, children and older adults are less likely to be exposed to the bacteria. In short, immunising teenagers with a MenB vaccine might mean lower rates of meningococcal disease across all ages.

Who can participate?

Students aged 16-18 attending year 12 (or equivalent) at one of the participating 6th form colleges in England, Scotland and Wales

What does the study involve?

Participating schools are randomly allocated to deliver one of two types of MenB vaccine: 4CMenB (also known as Bexsero) and MenB-fHBP (also known as Trumenba). Participants either get two doses of 4CMenB or MenB-fHBP given 6 months apart at their first two study visits, or two doses of 4CMenB 1 to 6 months apart at their last two study visits. These vaccines are approved for use in the UK, but are not routinely given to teenagers in this country. Samples are collected from the participants' throats to compare rates of MenB carriage before and after getting the MenB vaccine. Teenagers have three study visits over 12 to 18 months and all visits take place within schools.

What are the possible benefits and risks of participating? Not provided at time of registration

Where is the study run from? University of Oxford(UK)

When is the study starting and how long is it expected to run for? October 2017 to March 2024 Who is funding the study? 1. Department of Health (UK) 2. Pfizer (UK)

Who is the main contact? Emma Plested

Contact information

Type(s) Scientific

Contact name Mrs Emma Plested

Contact details Oxford Vaccine Group, University of Oxford CCVTM Churchill Hospital Headington Oxford United Kingdom OX3 7LE

Additional identifiers

EudraCT/CTIS number 2017-004609-42

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers 37350

Study information

Scientific Title

Evaluating the effect of immunisation with group B meningococcal vaccines on meningococcal carriage

Study objectives

Teenagers and young children are at increased risk of diseases such as meningitis and blood poisoning due to bacteria called meningococcus. Although these diseases can be serious, the meningococcus bacteria are 'carried' in the back of the throat of 1 in 10 teenagers without causing any symptoms. Most meningococcal disease in teenagers is due to Meningitis B (also known as MenB). The aim of this study is to find out whether immunising teenagers with vaccines against MenB can reduce the number of teenagers carrying these bacteria in their throat. This would be important because it could mean that teenage MenB immunisation would not only help protect teenagers against these potentially deadly diseases, but also that babies, children and older adults are less likely to be exposed to the bacteria. In short, immunising teenagers with a MenB vaccine might mean lower rates of meningococcal disease across all ages.

Ethics approval required

Old ethics approval format

Ethics approval(s) South Central – Berkshire B Research Ethics Committee, 02/03/2018, ref: 18/SC/0055

Study design Non-randomised; Both; Design type: Prevention, Vaccine, Cross-sectional

Primary study design Interventional

Secondary study design

Non randomised study

Study setting(s) School

Study type(s)

Prevention

Participant information sheet

Not available in web format, please use the contact details to request a patient information sheet

Health condition(s) or problem(s) studied

Vaccination against meningitis

Interventions

Vaccines are randomly allocated by the project statistician on a site by site basis. Sites remain assigned to one vaccine group only for the duration of the study. Two types of MenB vaccine are used: 4CMenB (also known as Bexsero) and MenB-fHBP (also known as Trumenba). Participants in this study will either get two doses of 4CMenB or MenB-fHBP given 6 months apart at their first two study visits, or two doses of 4CMenB 1 to 6 months apart at their last two study visits. These vaccines are approved for use in the UK, but are not routinely given to teenagers in this country. The two doses of MenB vaccine will be given by IM injection into the deltoid by trained research nurses/doctors within the school setting. Oropharyngeal samples are collected from teenager's throats to compare rates of MenB 'carriage' in teenagers before and after getting a MenB vaccine. Teenagers have three study visits, over 12 to 18 months and all visits would be held within schools. The follow up is 13 months for group 1 + 2 and up to 18 months for group 3.

Intervention Type

Biological/Vaccine

Phase

Phase IV

Drug/device/biological/vaccine name(s)

Bexsero, Trumenba

Primary outcome measure

Rates of carriage prevalence of any of meningococci genogroup B, C, W, X and Y before and after immunisation in both immunisation cohorts, compared with unimmunised controls; Timepoint(s): End of the study

Secondary outcome measures

Rates of carriage prevalence of particular Neisseria before and after immunisation in both immunisation cohorts, compared with controls, specifically:

- 1. Serogroup B meningococci
- 2. Hyper-invasive meningococcal strains
- 3. All meningococcal strains
- 4. Other Neisseria species
- 5. Meningococci of other non B serogroups and capsule null meningococci

6. Meningococci expressing antigens contained in 4CMenB and MenB-fHBP

The difference in acquisition of carriage of all N. meningitidis over a 12-month period in both immunised cohorts compared to unvaccinated participants

Overall study start date

01/10/2017

Completion date

31/03/2024

Eligibility

Key inclusion criteria

1. Male or female, aged 16-18 years attending year 12 (or equivalent) at one of the participating 6th form colleges in England, Scotland and Wales

2. Participant is willing and able to give informed consent for participation

3. In the Investigator's opinion, is able and willing to comply with all trial requirements.

4. Willing to have bacterial isolates from throat swabs stored for future research in ethically approved studies

5. Willing to allow his or her General Practitioner to be contacted to confirm vaccination status if necessary

Participant type(s)

All

Age group Child

Lower age limit 16 Years

Upper age limit 18 Years **Sex** Both

Target number of participants Planned Sample Size: 24000; UK Sample Size: 24000

Total final enrolment 24047

Key exclusion criteria

1. Evidence of a course of either 4CMenB or MenB-fHBP in the past (documentation or self-report)

2. History of anaphylaxis to any component of 4CMenB or MenB-fHBP

3. Any other significant disease or disorder which, in the opinion of the investigator, may either put the participants at risk because of participation in the trial, or may influence the result of the trial, or the participants ability to participate

4. Participant is known to be pregnant

Date of first enrolment

19/03/2018

Date of final enrolment

22/12/2019

Locations

Countries of recruitment England

Scotland

United Kingdom

Wales

Study participating centre Royal Alexandra Children's Hospital BSUHT Eastern Road East Sussex Brighton United Kingdom BN2 5BE

Study participating centre Bristol Children's Vaccine Centre Level 6 Education and Research Centre Upper Maudlin Street Bristol United Kingdom BS2 8AE

Study participating centre Public Health Wales

4th Floor, Number 2 Capital Quarter Tyndall Street Cardiff United Kingdom CF10 4BZ

Study participating centre

Health Protection Scotland 4th Floor, Meridian Court 5 Cadogan Street Glasgow United Kingdom G2 6QE

Study participating centre St Mary's Hospital Praed St

London United Kingdom W2 1NY

Study participating centre Research and Development Department

Above Breast Care Centre – First floor Maidstone Hospital Hermitage Lane Maidstone United Kingdom ME16 9QQ

Study participating centre Paediatric Research Team 5th Floor Royal Manchester Children's Hospital Oxford Road Manchester United Kingdom M13 9WL

Study participating centre University of Nottingham Health Service University Park Derby Rd Nottingham United Kingdom NG7 2QW

Study participating centre Oxford Vaccine Group CCVTM Churchill Hospital Oxford United Kingdom OX3 7LE

Study participating centre Research & Development

The Lantern centre Vicarage Lane Fulwood Preston United Kingdom PR2 8DW

Study participating centre University Hospital Southampton NHS Foundation Trust Southampton General Hospital Tremona Road Southampton United Kingdom SO16 6YD

Study participating centre

St George's, University of London

Cranmer Terrace London United Kingdom SW17 0RE

Study participating centre Research and Innovation

Room F08, Pinewood House Stockport NHS Foundation Trust Stepping Hill Hospital Stockport United Kingdom SK2 7JE

Study participating centre Clinical Trials Unit Wrightington Hospital Hall Lane Appley Bridge United Kingdom WN6 9EP

Sponsor information

Organisation University of Oxford

Sponsor details

Clinical Trials and Research Governance (CTRG) Joint Research Office, Block 60 Churchill Hospital Oxford England United Kingdom OX3 7LJ

Sponsor type Hospital/treatment centre

ROR

https://ror.org/052gg0110

Funder(s)

Funder type Government

Funder Name Department of Health; Grant Codes: PR-R18-0117-21001

Funder Name Pfizer UK

Alternative Name(s) Pfizer Ltd, Pfizer Limited

Funding Body Type Private sector organisation

Funding Body Subtype For-profit companies (industry)

Location United Kingdom

Results and Publications

Publication and dissemination plan

Planned publication of the results in a high impact peer reviewed journal as soon after the completion of the project as possible.

Intention to publish date

31/12/2025

Individual participant data (IPD) sharing plan

The data sharing plans for the current study are unknown and will be made available at a later date.

IPD sharing plan summary

Data sharing statement to be made available at a later date

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient- facing?
<u>Protocol article</u>	protocol	22/10/2020	26/10 /2020	Yes	No

<u>Protocol file</u>	version 7.0	28/02/2022 09/05 /2022	No	No
<u>HRA research</u> <u>summary</u>		28/06 /2023	No	No
<u>Protocol file</u>	version 8.1	08/01 15/12/2023 08/01 /2024	No	No
Other publications	Observational study of the vacccination programme	13/07/2022 <mark>18/06</mark> /2025	Yes	No