

Disposable-Syringe Jet Injector for Measles-Mumps-Rubella vaccine (DSJI MMR) study

Submission date 08/07/2010	Recruitment status No longer recruiting	☐ Prospectively registered
		☐ Protocol
Registration date 29/07/2010	Overall study status Completed	☐ Statistical analysis plan
		☒ Results
Last Edited 10/07/2019	Condition category Infections and Infestations	☐ Individual participant data

Plain English summary of protocol
Not provided at time of registration

Contact information

Type(s)
Scientific

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Additional identifiers

Protocol serial number
Asclin/002/2009

Study information

Scientific Title
A randomised, non-inferiority study comparing safety and immunogenicity of a disposable-syringe jet injector to needle and syringe for the administration of measles-mumps-rubella combination vaccine to healthy Brazilian infants aged 12 to 18 months

Acronym

DSJI/MMR

Study objectives

Immunogenicity of measles-mumps-rubella (MMR) vaccine administered by needle free device is non inferior to that induced by needle and syringe.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Approved on 4th April 2010 by:

1. PATH REC (ref: 499)
2. Brazilian CONEP National Ethics Committee (ref: 15810)

Primary study design

Interventional

Study design

Randomised controlled non-inferiority study

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Measles, mumps and rubella vaccination

Interventions

MMR vaccine, against measles, mumps and rubella (Schwarz, RIT 4385 and Wistar RA 27/3 strains, respectively), to be administered as a single 0.5 mL dose, subcutaneously, delivered by a needle-free device (experimental group) or by the conventional needle and syringe method (control group), in a 2:1 proportion, that is, 388 volunteers allocated to the experimental group and 194 to the control group. Volunteers will be followed-up for 42 days after vaccination. Blood collections for immunogenicity before vaccination and 42 days after vaccination. Adverse events will be registered in diary cards by parents/guardians during this period.

Intervention Type

Drug

Phase

Not Applicable

Drug/device/biological/vaccine name(s)

Measles-mumps-rubella vaccine

Primary outcome(s)

Antibody titres 42 days after vaccination: cut offs for enzyme-linked immunosorbent assay (ELISA) 231 units/mL (mumps) and 4 IU/mL for rubella. Plaque reduction neutralisation test (PRNT) for measles, cut off 0.20 IU/mL.

Key secondary outcome(s)

Local and systemic adverse events during 42 days after vaccination

Completion date

26/07/2011

Eligibility

Key inclusion criteria

1. Healthy children from 12 to 18 months of age, either sex
2. Up-to-date with their immunisation schedule according to age
3. Not enrolled in other clinical studies

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Child

Lower age limit

12 Months

Upper age limit

18 Months

Sex

All

Total final enrolment

582

Key exclusion criteria

1. Parents/guardians unable or unwilling to give consent for the study
2. Unable to follow the study procedures

Date of first enrolment

26/07/2010

Date of final enrolment

26/07/2011

Locations

Countries of recruitment

Brazil

Study participating centre
Av. Brasil 4365
Rio de Janeiro
Brazil
21040-360

Sponsor information

Organisation
Bio-Manguinhos/Fiocruz (Brazil)

ROR
<https://ror.org/05gj5j117>

Funder(s)

Funder type
Charity

Funder Name
Bill and Melinda Gates Foundation (USA) - PATH DSJI project

Funder Name
Bio-Manguinhos/Fiocruz (Brazil)

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary
Not provided at time of registration

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	results	01/03/2015	10/07/2019	Yes	No