Immunological response after early extra and regular MMR immunization: 10 years follow-up

Submission date 11/07/2023	Recruitment status No longer recruiting	[X] Prospectively registered [_] Protocol
Registration date 05/10/2023	Overall study status Completed	 Statistical analysis plan Results
Last Edited 19/12/2024	Condition category Infections and Infestations	 Individual participant data [X] Record updated in last year

Plain English summary of protocol

Background and study aims

From May 2013 until March 2014, a measles epidemic occurred in the Netherlands. During this epidemic, the Dutch Ministry of Health decided to offer infants between 6 and 12 months of age, living in the measles outbreak area, an early extra MMR (MMR-0) immunization during this outbreak, to protect children below 1 year of age in areas with lower immunization coverage (less than 90%) and thus with higher risk of exposure to measles virus. Researchers previously investigated the immunological response to early vaccination in a group of these children and a control group. Blood samples were taken at 14 months, 15.5 months, 2 years, 4 years and 6-7 years of age. About 12% of the children who received MMR-0 between 6 and 8 months of age had no detectable antibodies at 14 months of age before MMR-1. All other children had MMR-0induced antibody levels above the protective threshold and were regarded as protected against measles during the epidemic. After the regular MMR-1 vaccination at 14 months of age, all children had developed protective measles antibody levels. However, 3 years later, measles antibody levels had dropped below the protective threshold in 11% of children, particularly children who received MMR-0 between 6 and 8 months of age. At 6-7 years of age, this percentage had increased to 68%. Among the children who received MMR-0 between 9 and 12 months of age and the children of the control groups who only received their regular MMR-1 at 14 months of age, 21% and 11% had dropped below the threshold of protection, respectively. The aim of this study is to analyse the levels of measles-specific antibodies in children who participated in the EMI study or BMR-nul-study at about 11 years of age, which is 2 years (+/- 2 months) after having received the regular MMR-2 vaccination. The proportion of children who have gained protective antibody levels through MMR-2 vaccination compared to the ones who haven't will be assessed. Finally, the levels and the quality of antibodies will be compared between children who had received an MMR-0 at 6-12 months of age and the children of the control group who only received the regular MMR-1 vaccination at 14 months of age.

Who can participate?

Children aged 10 to 12 years who previously participated in the EMI clinical study on the immunological response after MMR-0 immunization or the epidemiological questionnaire study BMR-nul.

What does the study involve? One blood sample is obtained by fingerstick and one short questionnaire is completed.

What are the possible benefits and risks of participating? The children have no direct benefit from participating in the study. Blood collection will be done using a finger stick, which poses no risk.

Where is the study run from? Dutch National Institute of Health and the Environment (RIVM) (Netherlands)

When is the study starting and how long is it expected to run for? June 2023 to November 2024

Who is funding the study? The Dutch Ministry of Health, Welfare and Sport (Netherlands)

Who is the main contact? bmr-nul-emi@rivm.nl

Study website https://www.rivm.nl/mazelen/onderzoek#BMR-0

Contact information

Type(s) Principal Investigator

Contact name Dr I.M. Slits

Contact details 9 Antonie van Leeuwenhoeklaan Bilthoven Netherlands 3721MA +31 (0)886897312 bmr-nul-emi@rivm.nl

Additional identifiers

EudraCT/CTIS number Nil known

IRAS number

ClinicalTrials.gov number Nil known

Secondary identifying numbers IIV-628

Study information

Scientific Title

Observational study on immunological response 2 years after MMR-2 vaccination in children that received early extra or regular MMR immunization as infants to assess long-term effect of early extra MMR immunization

Acronym

EMI-3

Study objectives

Previously the immunological response to early vaccination was investigated in a cohort of children with the last blood sample obtained at 6-7 years of age, which is 6 years after the MMR-1 vaccination (NL45616.094.13/IIV-273 and NL69434.100.19/IIV-411). A large part of the early extra MMR-0 vaccinated children had protective measles levels at the age of 14 months. However, the decline in antibody levels was steeper than in children who received the regular MMR-1 vaccination and for some children antibody levels dropped below seroprotective levels at 4 and 6-7 years of age. This study will monitor the effect of MMR-2 on these children.

Ethics approval required

Ethics approval required

Ethics approval(s)

Approved 14/09/2023, Dutch Medical Ethics Committee MEC-U (Postbus 2500, Nieuwegein, 3430 EM, Netherlands; +31 (0)88 320 8784; info@mec-u.nl), ref: NL84855.100.23 (R23.050)

Study design Observational case-control study

Primary study design Observational

Secondary study design Case-control study

Study setting(s) Home, Internet/virtual

Study type(s) Prevention

Participant information sheet

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

Health condition(s) or problem(s) studied Measles immunization in children

Interventions

In this follow-up study, a single small blood sample will be collected by finger-stick of children who previously were immunized with an MMR-0 vaccination between 6-12 months of age and children of a control group who only received the regular MMR-1 at 14 months of age. Blood sampling through a finger-stick will be performed at 2 years (+/- 2 months) post MMR-2 vaccination. Parents can perform the finger-stick on their child themselves and mail the blood sample to the RIVM. If parents do not feel comfortable performing the finger-stick themselves, an alternative can be arranged. Furthermore, a digital questionnaire is filled out with questions about the health of the child and other vaccinations.

Intervention Type

Other

Primary outcome measure

Measles-specific serum IgG measured by Luminex multiplex-based serology and virusneutralizing antibodies by Plaque Reduction Neutralization Test (PRNT) 2 years (+/- 2 months) after MMR-2 vaccination

Secondary outcome measures

Serum IgG antibody concentrations against mumps and rubella measured by Luminex multiplexbased serology 2 years (+/- 2 months) after MMR-2 vaccination

Overall study start date

01/06/2023

Completion date 26/11/2024

Eligibility

Key inclusion criteria

Current inclusion criteria as of 30/01/2024:

1. Participation in the study on the immunological effects of early extra measles vaccination, as described in a separate study protocol (NL45616.094.13/IIV-273) or participation in the BMR-nulstudy on epidemiological effects of early extra measles vaccination as described in a separate study protocol

2. The parents/legal representatives accept participation in the trial according to the described procedures

- 3. Presence of an informed consent signed by both parents/legal representatives
- 4. Children must have received the MMR-2 vaccination

Previous inclusion criteria:

1. Participation in the study on the immunological effects of early extra measles vaccination, as described in a separate study protocol (NL45616.094.13/IIV-273)

2. The parents/legal representatives accept participation in the trial according to the described procedures

- 3. Presence of an informed consent signed by both parents/legal representatives
- 4. Children must have received the MMR-2 vaccination

Participant type(s)

Healthy volunteer

Age group Child

Lower age limit 10 Years

Upper age limit

12 Years

Sex Both

Target number of participants 80

Total final enrolment 88

Key exclusion criteria

1. Presence of a serious disease that requires medical care that can interfere with the results of the study

2. Known or suspected immunological disorder

3. Bleeding disorders

Date of first enrolment 09/10/2023

Date of final enrolment 18/07/2024

Locations

Countries of recruitment Netherlands

Study participating centre

Rijksinstituut voor Volksgezondheid en Milieu Antonie van Leeuwenhoeklaan 9 Bilthoven Netherlands 3721 MA

Sponsor information

Organisation National Institute for Public Health and the Environment

Sponsor details 9 Antonie van Leeuwenhoeklaan Bilthoven Netherlands 3721MA +31 (0)886894154 mensgebonden-onderzoek@rivm.nl

Sponsor type Research organisation

Website http://www.rivm.nl/en/

ROR https://ror.org/01cesdt21

Funder(s)

Funder type Government

Funder Name Ministerie van Volksgezondheid, Welzijn en Sport

Alternative Name(s) Dutch Ministry of Health, Welfare and Sport, VWS

Funding Body Type Government organisation

Funding Body Subtype National government

Location Netherlands

Results and Publications

Publication and dissemination plan Planned publication in a high-impact peer-reviewed journal

Intention to publish date

01/09/2025

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