Nasopharyngeale pneumokokken en meningokokken kolonisatie

Gepubliceerd: 19-09-2018 Laatst bijgewerkt: 15-05-2024

Current study is part of our long-term surveillance intended to monitor changes in serotype specific pneumococcal nasopharyngeal carriage since introduction of pneumococcal vaccination in the NIP in vaccinated children and their unvaccinated parents...

Ethische beoordeling Positief advies **Status** Werving gestopt

Type aandoening

Onderzoekstype Interventie onderzoek

Samenvatting

ID

NL-OMON28461

Bron

Nationaal Trial Register

Verkorte titel

OKIDOKI-5

Aandoening

Pneumococcal carraige and meningococcal carriage

Ondersteuning

Primaire sponsor: National Institute for Public Health and the Environment (RIVM)

Overige ondersteuning: Ministery of Health and Social Affairs

ZonMW

Onderzoeksproduct en/of interventie

Uitkomstmaten

Primaire uitkomstmaten

to determine vaccine- and non-vaccine serotype-specific pneumococcal carriage by culture.

Toelichting onderzoek

Achtergrond van het onderzoek

Current study is part of our long-term surveillance intended to monitor changes in serotype specific pneumococcal nasopharyngeal carriage since introduction of pneumococcal vaccination in the NIP in vaccinated children and their unvaccinated parents. In response to the recent increase of invasive meningococcal disease incidence due to meningococcal W, the Minister of Health decided to replace the monovalent meningococcal conjugate vaccine against MenC by the 4-valent conjugate vaccine MenACWY vaccination in spring 2018. The impact of MenACWY vaccination on the vacant niche is currently unknown. Pneumococcal and meningococcal carriage in young children will be monitored in the OKIDOKI-5 study. The study population consists of 330 24-month-old children that have been vaccinated according to the NIP, siblings of the participating children (24-month-old - <6 years of age) and one of the parents/legal guardians of each child. The children only received MenC vaccination not MenACWY. During a home visit saliva and a throat swab are collected of all participants. Of the 24-month old children also a nose swab is taken. The objective is to monitor changes in pneumococcal carriage after pneumococcal vaccination and to determine the best method for detection of meningococcal carriage. Results of the current meningococcal carriage can serve as a baseline for future studies.

Doel van het onderzoek

Current study is part of our long-term surveillance intended to monitor changes in serotype specific pneumococcal nasopharyngeal carriage since introduction of pneumococcal vaccination in the NIP in vaccinated children and their unvaccinated parents. Streptococcus pneumoniae (pneumococcus) is the leading cause of invasive pneumococcal disease (IPD) like

meningitis, sepsis and bacteremic pneumonia as well as of respiratory infections like community

acquired pneumonia and otitis media. The highest disease incidence is observed in children below two years of age and in elderly > 65 years of age. Disease caused by Streptococcus pneumoniae is preceded by asymptomatic nasopharyngeal acquisition and colonization. Neisseria meningitidis, the bacterium that causes meningococcal disease, is an obligate commensal of humans, which transiently colonizes the mucosa of the upper respiratory tract but only occasionally causing invasive meningococcal disease (IMD), depending on the clonal type and low population immunity to the particular strain. IMD is relatively rare (at present 100-150 cases/year in the Netherlands) but with severe symptoms and with mortality up to 20%, the burden of IMD remains high. Meningococci can be classified into 13 serogroups based on the capsular polysaccharides Chemistry and immunogenicity. Of these 13 serogroups, six are responsible for the vast majority of disease cases with strains of serogroup B, W and Y dominating in IMD in the Netherlands. The incidence is highest in children under 5 years of age followed by adolescents, young adults and elderly of over 75 years of age. Although cases of IMD are generally rare, the severity of

disease and the occurrence of outbreaks strongly support a role for prevention by

vaccination. Disease onset is often a-specific but, within hours IMD progresses towards meningitis, sepsis and/or septic shock. Vaccination with a 7-valent pneumococcal vaccine (Prevenar-7, PCV-7) was introduced in the Dutch National Immunization Program (NIP) for children in 2006 and replaced in 2011, by a 10-valent vaccine (PCV-10). In response to the recent increase in MenW IMD incidence, the Minister of Health decided to replace the monovalent meningococcal conjugate vaccine against MenC by the 4-valent conjugate vaccine MenACWY vaccination in spring 2018. In addition, MenACWY conjugate vaccination will be implemented at the age of 14 years in September 2018. PCV-10 and possibly

also MenACWY vaccine reduce the acquisition and density of vaccine serotypes in the nasopharynx of vaccinated children and subsequent reduces transmission to others leading to an indirect protection of the community (herd effects) thus enhancing the public health effect and costeffectiveness

of this approach. After PCV vaccination the vacant niche in the nasopharynx of vaccinated children is immediately filled by non-vaccine pneumococci and possibly other potential

pathogens that may be involved in respiratory or invasive disease. The impact of MenACWY vaccination on the vacant niche is currently unknown. Pneumococcal and meningococcal carriage in young children will be monitored in the OKIDOKI-5 study.

Onderzoeksopzet

One home visit per child/parent.

Onderzoeksproduct en/of interventie

A transnasal and transoral naopharyngeal swab and two saliva samples will be taken from the participants (parents and older siblings; no transnasal swab).

Contactpersonen

Publiek

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Wetenschappelijk

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Deelname eisen

Belangrijkste voorwaarden om deel te mogen nemen (Inclusiecriteria)

- •The children have to be of normal health (same health criteria apply as used in well-baby clinics when a child receives a vaccination, e.g. also children with small increases in temperature or cold are seen as children with normal health, fever >38.5°C in the last two days is not considered as normal health)
- •The parents/legal guardians have to be willing and able to participate in the trial according to procedure
- •The child is 24-months-old (\pm 8 weeks) or for older siblings; the child is between 24 month (-4 weeks) and <6 years of age)
- •The child has been vaccinated according to the Dutch NIP (including MenC vaccination, not MenACWY vaccination)
- Presence of a signed informed consent (the parents/legal guardians have given written informed consent after receiving oral and written information)

Parents/legal guardians of 24-month-old children are included when the child fulfils the inclusion criteria

Belangrijkste redenen om niet deel te kunnen nemen (Exclusiecriteria)

A potential participating child who meets any of the following criteria will be excluded from participation in this study:

- •Previous vaccinations with PCV using a vaccine and schedule that differs from the Dutch NIP of that age group
- Previous vaccinations with MenACWY vaccine
- Medical conditions that will severely affect immunological responses to vaccinations or nasopharyngeal carriage rates (certain chromosomal abnormalities or craniofacial abnormalities (like Trisomy 21 or schisis), known or suspected immunodeficiency disease or other medical conditions)

A parent/legal guardian who meets any of the following criteria will be excluded from participation in this study:

• Medical conditions that will severely affect immunological responses to vaccinations or nasopharyngeal carriage rates (certain chromosomal abnormalities or craniofacial abnormalities (like Trisomy 21 or schisis), known or suspected immunodeficiency disease or other medical conditions)

Onderzoeksopzet

Opzet

Type: Interventie onderzoek

Onderzoeksmodel: Parallel

Toewijzing: N.v.t. / één studie arm

Blindering: Open / niet geblindeerd

Controle: N.v.t. / onbekend

Deelname

Nederland

Status: Werving gestopt

(Verwachte) startdatum: 15-08-2018

Aantal proefpersonen: 760

Type: Werkelijke startdatum

Voornemen beschikbaar stellen Individuele Patiënten Data (IPD)

Wordt de data na het onderzoek gedeeld: Nog niet bepaald

Ethische beoordeling

Positief advies

Datum: 19-09-2018

Soort: Eerste indiening

Registraties

Opgevolgd door onderstaande (mogelijk meer actuele) registratie

ID: 46096

Bron: ToetsingOnline

Titel:

Andere (mogelijk minder actuele) registraties in dit register

Geen registraties gevonden.

In overige registers

Register ID

NTR-new NL7263 NTR-old NTR7485

CCMO NL65919.100.18 OMON NL-OMON46096

Resultaten