Rotavirus vaccinatie voor zuigelingen met een medisch risico

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Ethische beoordeling Positief advies **Status** Werving gestart

Type aandoening -

Onderzoekstype Observationeel onderzoek, zonder invasieve metingen

Samenvatting

ID

NL-OMON21569

Bron

NTR

Verkorte titel

RIVAR

Aandoening

RV vaccine coverage rates and timeliness among qualifying high-risk infants RV related hospitalizations among high-risk infants. (severe) RV gastroenteritis among high-risk infants up to 18 months of age.

Ondersteuning

Primaire sponsor: UMC Utrecht

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Overige ondersteuning: ZonMw

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Innovatiefonds Zorgverzekeraars Sparrenheuvel 16 Postbus 304, 3700 AH Zeist The Netherlands

GlaxoSmithKline Huis ter Heideweg 62 3705 LZ Zeist The Netherlands

Onderzoeksproduct en/of interventie

Uitkomstmaten

Primaire uitkomstmaten

- The impact of the RV vaccination program on number of hospitalizations due to RV gastroenteritis and symptomatic nosocomial RV infections among high-risk infants in the first and second year after implementation. Occurrence is expressed as cumulative number of children with at least one hospitalization due to RV gastroenteritis or symptomatic nosocomial RV infection in one of the participating hospitals per number of children at risk.

- Vaccine effectiveness in reducing episodes of severe RV gastroenteritis between 2 and 18 months of age among high-risk infants included in the follow-up study. Occurrence is expressed as number of events per 1000 person-years.

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- Proportion of eligible high-risk children receiving a full 2-dose course of RV vaccination after implementation of RV vaccination program.

Toelichting onderzoek

Achtergrond van het onderzoek

BACKGROUND AND MOTIVATION

Children with underlying medical conditions, prematurity and low birth weight are at increased risk of hospitalizations due to rotavirus (RV) gastroenteritis, RV related complications and mortality. In a recent Dutch study it was demonstrated that high-risk infants had increased risks of requiring hospitalization due to RV (1.7 to 4.4), of ICU

admission (RR: 4.2 to 7.9) and of acquiring nosocomial RV infection (OR: 3.2 to 3.6) compared to healthy infants. High-risk infants also experienced prolonged hospitalization (1.5 to 3.0 excess days) and generated higher healthcare costs (€648 to €1533 per patient). Most importantly, it was estimated that on average 6-7 high-risk infants succumbed due to direct and indirect consequences of RV infection annually in the Netherlands. No mortality was observed among healthy infants. A subsequent cost-effectiveness analysis demonstrated that RV vaccination of high-risk infants would be very cost-effective and potentially cost-saving from the healthcare payer perspective.

The results demonstrate that prevention against RV gastroenteritis is urgently needed for high-risk infants. Yet, there is no RV vaccination program in the Netherlands. Organizing a successful RV vaccination program within the Dutch infant immunization framework generates additional challenges concerning adequate reach-out to high-risk infants because of the strict age-window indicated for RV vaccine administration. The first dose of RV vaccine should be administered between 6 and 14 weeks of age and vaccination completed no later than 24 or 32 weeks of age, depending on the vaccine. Secondary and tertiary pediatric care would provide an excellent environment for RV vaccination in order to reach optimal RV vaccine coverage rates in high-risk infants and to ascertain timely vaccination, because high-risk infants are closely followed here during their first month of life. There is however little experience in the Netherlands with organizing immunization programs for target groups through secondary and tertiary pediatric care.

Furthermore, there is a lack of data on RV vaccine performance among the special populations of high-risk infants, although these patients require protection most. Further data on vaccine effectiveness are needed to improve vaccination guidelines pertaining to high-risk infants.

This project will study the feasibility and impact of implementing a RV vaccination program for high-risk infants organized through secondary and tertiary care.

This study will also determine RV vaccine effectiveness among high-risk infants.

PRIMARY OBIECTIVES

- To evaluate the feasibility of RV vaccination of high-risk infants organized through secondary and tertiary pediatric care as measured by vaccine coverage and timeliness of vaccination.
- To evaluate the impact of RV vaccination of high-risk infants organized through secondary and tertiary pediatric care on rotavirus related hospitalizations among this patientgroup.
- To evaluate the protective effectiveness of at least 1 dose of RV vaccine against severe RV gastroenteritis up to 18 months of life.

RV VACCINATION PROGRAM

Dutch Hospitals with Neonatal Intensive Units and associated post IC/HC hospitals will be approached for participation in an implementation project of a hospital-based RV vaccination program for high-risk infants. The RV vaccination program consists of offering a 2-dose

course of the oral monovalent RV vaccine free of charge to all high-risk infants receiving care in one of the participating hospitals at discharge or during routine clinic visits and will be accompanied by a RV active surveillance program within the participating hospitals.

STUDY DESIGN

An observational study accompanying the program will assess occurrence of gastroenteritis among high-risk infants pre- and post-implementation, RV epidemiology within hospitals, vaccine coverage rates and timeliness of vaccination among RV vaccine eligible high-risk infants. A step-wedged design is used to account for inter-season variability of RV epidemics. Occurrence of (RV) gastroenteritis among high-risk infants will be studied by recruiting high-risk infants, both pre- and post-implementation, for an individual follow-up study until 18-months of age. Follow-up includes parental reporting of gastroenteritis, symptom and severity scoring and collection of stool samples.

STUDY POPULATION

Hospitals and eligible high-risk infants receiving care at these hospitals.

PRIMARY ENDPOINTS

- RV vaccine coverage rates, timeliness of vaccination and impact of the RV vaccination program in reducing RV related hospitalizations among high-risk infants.
- Vaccine effectiveness against severe RV gastroenteritis among high-risk infants up to 18 months of age.

Doel van het onderzoek

Children with underlying medical conditions, prematurity and low birth weight are at increased risk of hospitalizations due to rotavirus (RV) gastroenteritis, RV related complications and mortality. In a recent Dutch study it was demonstrated that high-risk infants had increased risks of requiring hospitalization due to RV, of ICU admission and of acquiring nosocomial RV infection compared to healthy infants. High-risk infants also experienced prolonged hospitalization and generated higher healthcare costs.

The results demonstrate that prevention against RV gastroenteritis is urgently needed for high-risk infants. Yet there is no RV vaccination program in the Netherlands.

Furthermore, there is lack of data on RV vaccine performance among the special populations of high-risk infants. Further data on vaccine effectiveness are needed to improve vaccination guidelines pertaining to high-risk infants.

This project will 1)

study the feasibility and impact of implementing a RV vaccination program for high-risk infants organized through secondary and tertiary care, and

2) determine RV vaccine effectiveness among high-risk infants.

Onderzoeksopzet

Program impact is measured 1 and 2 years post-implementation. Vaccine effectiveness is measured based on follow-up of each enrolled infant up to 18 months of age.

Onderzoeksproduct en/of interventie

No randomized interventions. This is a step-wedged implementation project of a rotavirus vaccination program for high-risk infants in participating hospitals combined with and observational before-after cohort study measuring rotavirus gastroenteritis occurrence in a pre- and post-implementation cohort of high-risk infants.

Contactpersonen

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Deelname eisen

Belangrijkste voorwaarden om deel te mogen nemen (Inclusiecriteria)

- 1. Infants 6 weeks 0 days to 13 weeks 6 days of age who receive medical care in hospital or through outpatient clinics at the time of the first vaccine dose administration in one of the participating hospitals and
- 2. Diagnosed with at least one high-risk condition:
- Gestational age less than 36 weeks and 0 days
- Birth weight less than 2500 grams
- A qualifying diagnoses of severe congenital malformation or perinatal morbidity

Belangrijkste redenen om niet deel te kunnen nemen (Exclusiecriteria)

- Known hypersensitivity to any of the vaccine components.
- Previous intussusception or an uncorrected congenital condition predisposing to intussusception (such as Meckel's diverticle).
- A diagnosis of severe (congenital) immunodeficiency syndrome.

Onderzoeksopzet

Opzet

Type: Observationeel onderzoek, zonder invasieve metingen

Onderzoeksmodel: Anders

Toewijzing: N.v.t. / één studie arm

Blindering: Open / niet geblindeerd

Controle: N.v.t. / onbekend

Deelname

Nederland

Status: Werving gestart

(Verwachte) startdatum: 01-12-2014

Aantal proefpersonen: 2000

Type: Verwachte startdatum

Ethische beoordeling

Positief advies

Datum: 13-08-2015

Soort: Eerste indiening

Registraties

Opgevolgd door onderstaande (mogelijk meer actuele) registratie

Geen registraties gevonden.

Andere (mogelijk minder actuele) registraties in dit register

Geen registraties gevonden.

In overige registers

Register ID

NTR-new NL5213 NTR-old NTR5361

Ander register 80-83600-98-20129 : ZonMW

Resultaten