# Mercurius 5 years follow-up study

No registrations found.

Ethical review	Positive opinion
Status	Pending
Health condition type	-
Study type	Observational non invasive

## **Summary**

## ID

NL-OMON28614

Source NTR

**Brief title** Mercurius 5 year FU

#### **Health condition**

Healthy infants

## **Sponsors and support**

**Primary sponsor:** Nutricia Research **Source(s) of monetary or material Support:** Nutricia Research

### Intervention

### **Outcome measures**

#### **Primary outcome**

Key outcome: Body Mass Index (BMI) at 3, 4, and 5 years of age

#### Secondary outcome

- Prevalence of overweight and obesity at 3, 4, and 5 years of age
- Sum of skin fold thicknesses at 3, 4, and 5 years of age

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- Percentage total body fat (derived from skinfolds)
- Head- and waist- circumference at 3, 4, and 5 years of age

## **Study description**

#### **Background summary**

The present study is a follow up study of the original Mercurius study (NTR3683) at the age of 3-5 years. The main objective of this study is to gain insight into the possible long-term effects, up to 5 years of age, on growth and body composition development in subjects who have received the test product compared to subjects who have received the control product in the first 4 months of age and in comparison to the breastfed reference group. Study visits will take place at 3, 4 and 5 years of age. A phone call will take place at 3.5 and 4.5 years of age.

#### **Study objective**

The infant formula containing a new fat blend, consumed in the first 4 months of life, will have a positive effect on the growth trajectory in toddlerhood up to 5 years of age.

#### Study design

Visit at 3, 4 and 5 years of age. Phone call at 3.5 and 4.5 years of age

#### Intervention

N/A

## Contacts

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## **Eligibility criteria**

## **Inclusion criteria**

- Subjects that participated in the original Mercurius study (NTR3683) up to visit 5 (approx. 4 months of age) are eligible for this follow up study.

- Written Informed Consent

## **Exclusion criteria**

- Investigator's uncertainty about the willingness or ability of the child and parents to comply with the protocol requirements.

- Subjects that were recruited by investigators who decided not to participate in the 5 year follow up study.

## Study design

## Design

Study type:	Observational non invasive
Intervention model:	Other
Masking:	Single blinded (masking used)
Control:	N/A , unknown

### Recruitment

#### NL

Recruitment status:	Pending
Start date (anticipated):	01-01-2016
Enrollment:	230
Туре:	Anticipated

## **Ethics review**

Positive opinionDate:21-12-2015Application type:First submission

## **Study registrations**

### Followed up by the following (possibly more current) registration

No registrations found.

## Other (possibly less up-to-date) registrations in this register

No registrations found.

### In other registers

**Register** NTR-new NTR-old Other ID NL5421 NTR5538 : EPI1.C.C Nutricia Research

## **Study results**