# Pharmacokinetic evaluation of long-term Xolair-therapy in severe asthma patients

No registrations found.

**Ethical review** Positive opinion **Status** Recruiting

Health condition type -

**Study type** Observational non invasive

# **Summary**

### ID

NL-OMON25198

**Source** 

Nationaal Trial Register

Brief title PHLOX

**Health condition** 

Allergic Asthma

# **Sponsors and support**

**Primary sponsor:** Medical Centre Leeuwarden

Source(s) of monetary or material Support: MCL Academy Science-committee

#### Intervention

#### **Outcome measures**

## **Primary outcome**

The percentage of patients with a ratio omalizumab trough-level to baseline IgE larger than 15:1

## **Secondary outcome**

Association of the ratio omalizumab trough-level to baseline IgE and patients yearn for the next gift

Association of the ratio omalizumab trough-level to baseline IgE and asthma-related outcomes (ACQ, AQLQ, FEV1, NO)

# **Study description**

## **Background summary**

Omalizumab has been used in the treatment of severe allergic asthma for several years. Early trials found that adequate suppression of IgE was achieved when omalizumab was in excess over IgE 15:1. It remains unknown what this ratio is in patients treated for more than a year. Furthermore, some patients are able to extend their dose-interval, while some patients indicate that they yearn for their next omalizumab gift. In this trial the ratio between trough omalizumab-level and baseline IgE will be examined. The patients yearn for the next gift and the correlation with the ratio between trough omalizumab-level and baseline IgE will be examined.

## Study objective

Our hypothesis is that the ratio omalizumab trough-level to baseline IgE differs from the 15:1 ratio, which was found to induce adequate asthma-suppression in early trials. The patients yearn for the next gift may be associated with an inadequate ratio.

## Study design

Data will be collected during the 6-month evaluation

## Intervention

N/A

# **Contacts**

#### **Public**

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#### Scientific

Medisch Centrum Leeuwarden

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# **Eligibility criteria**

## **Inclusion criteria**

Severe asthma patients, receiving more than 12 months of omalizumab therapy, >18 months old, able to sign informed consent.

## **Exclusion criteria**

None

# Study design

## **Design**

Study type: Observational non invasive

Intervention model: Other

Allocation: Non controlled trial

Masking: Open (masking not used)

Control: N/A, unknown

## Recruitment

NL

Recruitment status: Recruiting
Start date (anticipated): 01-06-2019

Enrollment: 30

Type: Anticipated

## **IPD** sharing statement

Plan to share IPD: No

## Plan description

N/A

# **Ethics review**

Positive opinion

Date: 06-06-2019

Application 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 ID

NTR-new NL7794

Other RTPO Leeuwarden: nWMO 375

# **Study results**

## **Summary results**

N/A