# Medical assessment of adverse health outcomes in Dutch childhood cancer survivors; a nationwide project; SKION LATER Q2008 research: Is pulmonary dysfunction a late effect of cyclophosphamide treatment?

Published: 23-01-2015 Last updated: 27-04-2024

to assess the pulmonary toxicity in patients treated with cyclophosphamide compared to a

control group

**Ethical review** Approved WMO

**Status** Recruitment stopped

**Health condition type** Lower respiratory tract disorders (excl obstruction and infection)

**Study type** Observational invasive

# **Summary**

#### ID

NL-OMON47559

#### Source

ToetsingOnline

#### **Brief title**

SKION LATER Q2008 - pulmonology

## Condition

• Lower respiratory tract disorders (excl obstruction and infection)

#### **Synonym**

pulmonary dysfunction

## Research involving

Human

# **Sponsors and support**

Primary sponsor: Stichting Kinderoncologie Nederland

Source(s) of monetary or material Support: Quality of life gala

## Intervention

Keyword: late effects, lung function, pulmotoxicity

## **Outcome measures**

#### **Primary outcome**

Abnormal PFT, defined as TLC/FVC <75% (restrictive dysfunction) of predicted

for age and sex matched controls of a normal population.

Other PFT parameters, i.e. FEV1%FVC <75% (obstruction) and/or DLCO/TLCO <75%

(diffusion).

# **Secondary outcome**

not applicable

# **Study description**

#### **Background summary**

Advances in diagnosis and treatment of childhood cancer over the last decades have dramatically increased long-term survival. As a result, the numbers of childhood cancer survivors (CCS) are growing and it has become increasingly clear that the former disease and its treatment can significantly impair long-term health. The need for long-term follow-up is uniformly recognized. Research focusing on identification and characterization of high-risk populations is an essential foundation on which to build evidence-based recommendations for long-term follow-up. Furthermore, research focusing on more accurate screening tests and effective interventions is needed to reduce excess morbidity and mortality in CCS. The SKION LATER Q2008 - pulmonology study phocuses on late pulmonary toxicity in CCS

## Study objective

to assess the pulmonary toxicity in patients treated with cyclophosphamide

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# Study design

In 2 groups of childhood cancer survivors (CCS, each 260 patients) pulmonary late effects will be assessed by a pulmonary function test:

- CCS treated with cyclophosphamide but no other potentially pulmotoxic treatment modalities (chemo and/or radiotherapy)
- CCS treated with neither cyclophosphamide nor any other potentially pulmotoxic treatment modalities (chemo and/or radiotherapy)

## Study burden and risks

no risks associated with the pulmonary function test. The test is not painful, patients wil have to blow in a tube

# **Contacts**

#### **Public**

Stichting Kinderoncologie Nederland

Heidelberglaan 25 Utrecht 3584CS

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#### Scientific

Stichting Kinderoncologie Nederland

Heidelberglaan 25 Utrecht 3584CS NL

# **Trial sites**

## **Listed location countries**

**Netherlands** 

# **Eligibility criteria**

## Age

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Adolescents (12-15 years) Adolescents (16-17 years) Adults (18-64 years) Elderly (65 years and older)

## Inclusion criteria

childhood cancer survivors (CCS) who either received cyclophosphamide (Cy) treatment but no other potentially pulmotoxic treatment or who received neither Cy nor any other potentially pulmotoxic treatment

## **Exclusion criteria**

childhood cancer survivors who received potentially pulmotoxic treatment

# Study design

# **Design**

Study type: Observational invasive

Intervention model: Other

Allocation: Non-randomized controlled trial

Masking: Open (masking not used)

Control: Active

Primary purpose: Basic science

## Recruitment

NL

Recruitment status: Recruitment stopped

Start date (anticipated): 04-05-2016

Enrollment: 520

Type: Actual

# **Ethics review**

## Approved WMO

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Date: 23-01-2015

Application type: First submission

Review commission: METC Amsterdam UMC

Approved WMO

Date: 05-01-2016

Application type: Amendment

Review commission: METC Amsterdam UMC

Approved WMO

Date: 20-06-2017

Application type: Amendment

Review commission: METC Amsterdam UMC

Approved WMO

Date: 23-05-2018

Application type: Amendment

Review commission: METC Amsterdam UMC

Approved WMO

Date: 22-08-2018

Application type: Amendment

Review commission: METC Amsterdam UMC

# **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

CCMO NL35002.018.11