Levonorgestrel spiraal versus progesteron oraal voor de behandeling van hyperplasie zonder atypie van het endometrium.

Published: 07-01-2009 Last updated: 06-05-2024

Treatment of endometrial hyperplasia by an IUD that contains progesteron is more effective and has less side-effects than the treatment with progesteron oral.

Ethical review Approved WMO **Status** Recruitment stopped

Health condition type Reproductive neoplasms female benign

Study type Interventional

Summary

ID

NL-OMON36959

Source

ToetsingOnline

Brief title

HYPROLEV-study

Condition

Reproductive neoplasms female benign

Synonym

abnormal endometrial tissue, precancer leasion of woomb

Research involving

Human

Sponsors and support

Primary sponsor: obstetrie en gynaecologie

Source(s) of monetary or material Support: Ministerie van OC&W

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Intervention

Keyword: endometrium, hyperplasia, levonorgestrel IUD, progesteron

Outcome measures

Primary outcome

endometrial atrofy after treatment

Secondary outcome

less side-effects in the study-group than the control-group

Study description

Background summary

Endometrial hyperplasia without atypie is treated by oral progestron. This treatment has some side-effects. Another way of treating this condiion is by an IUD that contains progesteron. This method is already been investigated and is also effective. We think that this second option (the IUD which contains progesteron) is more effective and is also has less side-effects than the oral treatment. That is why we want to do this ranomized controlled trial. For this study we need to include 90 patients.

Study objective

Treatment of endometrial hyperplasia by an IUD that contains progesteron is more effective and has less side-effects than the treatment with progesteron oral.

Study design

prospective randomized clinical trial

Intervention

One group will be treated bij oral progesteron and the oter group will get a levonorgestrel-IUD.

Study burden and risks

Patients are four times asked to fill in a form which will take about 10-15 minutes each time.

There are no risks for the patient in this study.

Contacts

Public

Selecteer

Debyelaan 25 6202 AZ Maastricht Nederland **Scientific** Selecteer

Debyelaan 25 6202 AZ Maastricht Nederland

Trial sites

Listed location countries

Netherlands

Eligibility criteria

Age

Adults (18-64 years) Elderly (65 years and older)

Inclusion criteria

endometrial hyperplasia (simple and complex), Dutch language

Exclusion criteria

endometrial cancer, hyperplasia with atypie, history of hyperplasia, contra-indication for treatment with progesteron, biopt not possible for any reason

Study design

Design

Study type: Interventional

Intervention model: Parallel

Allocation: Randomized controlled trial

Masking: Open (masking not used)

Control: Active

Primary purpose: Treatment

Recruitment

NL

Recruitment status: Recruitment stopped

Start date (anticipated): 11-04-2011

Enrollment: 90

Type: Actual

Medical products/devices used

Product type: Medicine

Brand name: Mirena

Generic name: Levonorgestrel IUD

Registration: Yes - NL outside intended use

Product type: Medicine

Brand name: provera

Generic name: medroxyprogesteron

Registration: Yes - NL intended use

Ethics review

Approved WMO

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Date: 07-01-2009

Application type: First submission

Review commission: METC academisch ziekenhuis Maastricht/Universiteit

Maastricht, METC azM/UM (Maastricht)

Approved WMO

Date: 21-07-2009

Application type: First submission

Review commission: METC academisch ziekenhuis Maastricht/Universiteit

Maastricht, METC azM/UM (Maastricht)

Approved WMO

Date: 23-06-2010

Application type: Amendment

Review commission: METC academisch ziekenhuis Maastricht/Universiteit

Maastricht, METC azM/UM (Maastricht)

Approved WMO

Date: 09-02-2011
Application type: Amendment

Review commission: METC academisch ziekenhuis Maastricht/Universiteit

Maastricht, METC azM/UM (Maastricht)

Approved WMO

Date: 21-03-2011

Application type: Amendment

Review commission: METC academisch ziekenhuis Maastricht/Universiteit

Maastricht, METC azM/UM (Maastricht)

Approved WMO

Date: 26-04-2011
Application type: Amendment

Review commission: METC academisch ziekenhuis Maastricht/Universiteit

Maastricht, METC azM/UM (Maastricht)

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

EudraCT EUCTR2008-006715-21-NL

CCMO NL24978.068.08