Treatment of intertrigo with zinc oxide in ketoconazolecream with or without substitution of hydrocortisone-acetaat

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Clinical trial to investigate a difference in effectiveness for the treatment of intertrigo in wich the combination of zinc oxide 10% in ketoconazole will to be compared with zinc oxide 10% and hydrocortisone-acetaat 1% in ketoconazole.

Ethical review Approved WMO

Status Recruitment stopped

Health condition type Skin and subcutaneous tissue disorders NEC

Study type Interventional

Summary

ID

NL-OMON30558

Source

ToetsingOnline

Brief title

Treatment of intertrigo with or without hydrocortisone?

Condition

Skin and subcutaneous tissue disorders NEC

Synonym

intertigo, moist lesion

Research involving

Human

Sponsors and support

Primary sponsor: Medisch Centrum Leeuwarden

Source(s) of monetary or material Support: De kosten vallen geheel binnen de kosten

van de reguliere gezondheidszorg

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Intervention

Keyword: Hydrocortisone, intertrigo, Randomized Controled Trial, Therapy

Outcome measures

Primary outcome

Reduce in percentage of the area of the intertigo compared to the other side that has not been treated with the hydrocortisone-acetaat.

Secondary outcome

Local complainments as pruritis, pain, redness and scaly. Also the results of the cultured samples are evaluate whereby we will look at the percentage of positive results and the reaction or the different treatments.

Study description

Background summary

Intertrigo is a skin disease especially seen in hospitals and other care-centers. This disease starts in the large skin folds as the groin, inframammary and in the axillae. There is no consensus for the treatment at this moment but the main point is to dry the skin. Of all the topical treatments, only the anti-myotics have an controled investigation. In the Medical Center of Leeuwarden intertrigo is treatened with the combination of zinc oxide in ketoconazol or zinc oxide and hydrocortisone-acetaat in ketoconazole. Zinc oxide is to dry-in the skin, ketoconazole for the dermatofyt or yeast and hydrocortisone-acetaat for the redness and irritation of the skin.

Study objective

Clinical trial to investigate a difference in effectiveness for the treatment of intertrigo in wich the combination of zinc oxide 10% in ketoconazole will to be compared with zinc oxide 10% and hydrocortisone-acetaat 1% in ketoconazole.

Study design

It is a dubble-bline, randomized, standardtherapy-controled phase III study,

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with a duration of aproximately 6 months. There is an minimum of 48 inclusions for the statistical calculations so there will be approximately 65 participants.

Intervention

All patients will have both the combination with substitution with hydrocortisone-acetaat as the cream combination without it.

Study burden and risks

There will be no extra burdening when a patient participate in our study because the study is therapeuticaly. Physical there may be a burdening because of the location of the intertrigo, wich are on the whole the intimate areas. For the study it is necessary to have a good look at the intertrigo and trace it onto foil. This has to be done twice.

A sample has to be taken once, but this is a part of the normal care for the patients.

No invasive treatment is necessary in the interest of the study.

Contacts

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Trial sites

Listed location countries

Netherlands

Eligibility criteria

Age

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

Inclusion criteria

- Two-sided intertrigo in the folds of the groin, axillairy or submammary
- The intertrigo may not been treated yet beside treatment with zincoxide.
- There has to be a control after 5-7 days.

Exclusion criteria

- Patients may not have a hypersensibility for one of the componements of the different creams
- Patients may not have a (skin)disease that might can influence the aspect or the threatment for this study neither have a contraindication to start the treatment.

Study design

Design

Study phase: 3

Study type: Interventional

Intervention model: Parallel

Allocation: Randomized controlled trial

Masking: Double blinded (masking used)

Control: Active

Primary purpose: Treatment

Recruitment

NI

Recruitment status: Recruitment stopped

Start date (anticipated): 01-04-2007

Enrollment: 65

Type: Actual

Medical products/devices used

Product type: Medicine

Brand name: -

Generic name: hydrocortisone-acetaat

Registration: Yes - NL outside intended use

Product type: Medicine

Brand name: -

Generic name: ketoconazole

Registration: Yes - NL intended use

Product type: Medicine

Brand name: -

Generic name: Zinc oxide

Registration: Yes - NL intended use

Ethics review

Approved WMO

Date: 12-03-2007

Application type: First submission

Review commission: RTPO, Regionale Toetsingscie Patientgebonden Onderzoek

(Leeuwarden)

Approved WMO

Date: 26-03-2007

Application type: First submission

Review commission: RTPO, Regionale Toetsingscie Patientgebonden Onderzoek

(Leeuwarden)

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 EUCTR2006-006234-18-NL

CCMO NL15277.099.06