Is preventing hypothermia in patients undergoing a primairy Total Knee Prosthesis or Total Hip prosthesis with the Barrier Easywarm as effective as with Forced-Air Warming.

Published: 03-04-2017 Last updated: 11-04-2024

If there is comparable Efficacy in preventing hypothermia we can start using the barrier easywarm instead of the BairHugger. With the Bair hugger there is a disruption in airflow in the operation theatre which might bring a higher risk of infection...

Ethical review Approved WMO **Status** Recruitment stopped

Health condition type Bone and joint therapeutic procedures

Study type Interventional

Summary

ID

NL-OMON45308

Source

ToetsingOnline

Brief title

Comparable Efficacy in Orthopedic Patients?

Condition

Bone and joint therapeutic procedures

Synonym

Hypothermia Patientheating

Research involving

Human

Sponsors and support

Primary sponsor: HagaZiekenhuis

Source(s) of monetary or material Support: Ziekenhuis

Intervention

Keyword: Bair Hugger, Barrier easywarm, Forced air warming, Hypothermia

Outcome measures

Primary outcome

Temperature during operation.

Secondary outcome

NVT

Study description

Background summary

Prevention of hypothermia in patients undergoing a primairy Total Knee Prosthesis or Total Hip prosthesis. We want to investigate if the Barrier Easywarm is as effective as with the Forced-Air Warming.

Study objective

If there is comparable Efficacy in preventing hypothermia we can start using the barrier easywarm instead of the BairHugger.

With the Bair hugger there is a disruption in airflow in the operatieon theatre which might bring a higher risk of infection.

Study design

Randomized Controlled Trial, non inferiority study

Intervention

40 patients will be randomized in the barrier Easywarm group, the other 40 in the Bairhugger group.

During the operatioon the core temperature will be measured by the 3M SpotOn

monitoring system

Study burden and risks

Both warming systems are already used in operating theatres.

Contacts

Public

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Trial sites

Listed location countries

Netherlands

Eligibility criteria

Age

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

Inclusion criteria

Patients undergoing total hip or knee prosthesis

Exclusion criteria

severe arterial disease

Study design

Design

Study type: Interventional

Intervention model: Parallel

Allocation: Randomized controlled trial

Masking: Open (masking not used)

Control: Active

Primary purpose: Prevention

Recruitment

NL

Recruitment status: Recruitment stopped

Start date (anticipated): 29-05-2017

Enrollment: 80

Type: Actual

Ethics review

Approved WMO

Date: 03-04-2017

Application type: First submission

Review commission: METC Leiden-Den Haag-Delft (Leiden)

metc-ldd@lumc.nl

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 NL60416.098.16