A global comparison of Signature Guides and conventional instrumentation in the Oxford partial knee

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The primary purpose of this study is to compare alignment criteria in the Oxford Partial Knee using conventionalinstrumentation and Signature Custom Guides. Secondly it is questioned whether the Signature technology can reduce the number of surgical...

Ethical review Approved WMO **Status** Recruitment stopped

Health condition type Bone and joint therapeutic procedures

Study type Interventional

Summary

ID

NL-OMON41709

Source

ToetsingOnline

Brief title

GK7 Oxford Signature

Condition

Bone and joint therapeutic procedures

Synonym

joint deterioration, Osteoarthrititis

Research involving

Human

Sponsors and support

Primary sponsor: BioMet

Source(s) of monetary or material Support: Biomet

Intervention

Keyword: Oxford UKP, RCT, Signature patient specific guiding

Outcome measures

Primary outcome

Primary Endpoint: To determine the accuracy and precision of Signature Guides

in the Oxford Knee by the percentage of

knees achieving optimal alignment

Secondary outcome

Secondary Endpoint: Average Number of Instrument Cases Used

In addition, clinical outcomes and cost efficiency data will be collected to

develop economical models.

Study description

Background summary

Recently, the Signature Custom Guide technology was introduced in total knee arthroplasty. Instead of using an x-ray, the

preoperative plan is created by an MRI that is uploaded to a software system so that the surgeon can plan the case

preoperatively while seeing the entire knee and leg (not obscured by soft tissue) on their computer screen prior to surgery.

Using the planning software, a custom guide is created to align pins used to secure traditional cutting blocks while

performing distal femoral and proximal tibial cuts.

By using patient specific guides it is hoped to reduce the number of outliers in placement of the uni knee. Further it is

questioned what is the influence of high versus low volume users with this technique.

Study objective

The primary purpose of this study is to compare alignment criteria in the Oxford Partial Knee using conventional

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instrumentation and Signature Custom Guides.

Secondly it is questioned whether the Signature technology can reduce the number of surgical instruments needed during surgery.

Overall, the intent is to collect the performance and clinical outcomes of the Oxford Partial Knee System using Signature

Custom Guides or Conventional Instrumentation to develop economical models.

Study design

Prospective Multi-Center Randomized Two Armed Trial

Intervention

placement of the unicompartmental knee (Oxford) with Signature (patient specific instrumentation) compared to traditional instruments

Study burden and risks

Because two routine ways to perform Unicompartmental knee replacement with the Oxford are compared and only one additional CT scan is made to objectify the outline of the prosthesis placement is determined, the burden for the patient is limited.

Contacts

Public

Biomet Nederland BV

Toermalijnring 600 Dordrecht 3316LC NL

Scientific

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Trial sites

Listed location countries

Netherlands

Eligibility criteria

Age

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

Inclusion criteria

- -Individuals with osteoarthritis or Avascular necrosis limited to the medial compartment of the knee that is also lacking patellofemoral or lateral compartment disease.
- -Patients 21 and over.

Exclusion criteria

- -Active Infection, sepsis or osteomyelitis
- -Use of prosthesis in lateral compartment of the knee
- -Rheumatoid arthritis or other forms of inflammatory joint disease
- -Revision of failed prosthesis, failed upper tibial osteotomy, or post traumatic arthritis after tibial plateau fracture
- -Insufficiency of the collateral, anterior, or posterior cruciate ligaments which would preclude stability of the device.
- -Disease or damage to the lateral compartment of the knee
- -Patients incapable of following directions regarding rehabilitation
- -Patients who refuse, cannot, or should not receive a CT or MRI.

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): 14-09-2015

Enrollment: 22

Type: Actual

Medical products/devices used

Generic name: Signature

Registration: No

Ethics review

Approved WMO

Date: 09-04-2014

Application type: First submission

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

metc-ldd@lumc.nl

Approved WMO

Date: 12-06-2015

Application type: Amendment

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

metc-ldd@lumc.nl

Approved WMO

Date: 25-03-2016

Application type: Amendment

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

ClinicalTrials.gov NCT01763684 CCMO NL47831.098.14