

Comparison of femoral nerve block with posterior capsule infiltration versus anterior and posterior capsule infiltration after total knee replacement

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This prospective randomized controlled trial designed to compare the quality of analgesia offered by SFNB and LiA and their effects on morphine consumption, mobilization and pain control postoperatively. Our first hypothesis concerning opioid use is...

Ethische beoordeling	Positief advies
Status	Werving gestart
Type aandoening	-
Onderzoekstype	Interventie onderzoek

Samenvatting

ID

NL-OMON22055

Bron

Nationaal Trial Register

Verkorte titel

LiFeAnKeR

Aandoening

Total knee arthroplasty femoral nerve block Local infiltration analgesia

Ondersteuning

Primaire sponsor: S. van Kralingen

M. Vogel

OLVG

Amsterdam

Overige ondersteuning: none

Onderzoeksproduct en/of interventie

Uitkomstmaten

Primaire uitkomstmaten

- Opioid consumption measured by mg of morphine i.v. used daily (every morning at am by painconsulent and/or researcher)

- Mobilization measured by Modified Iowa Levels of Assistance Scale (MILAS)

Toelichting onderzoek

Achtergrond van het onderzoek

This prospective randomized controlled trial compares the SFNB group with the LiA group. The SFNB group gets a femoral nerve block with ropivacaine 0.375% and 75 microgram clonidine, and furthermore receives 50 cc ropivacaine 0.2% with epinephrine 1:100.000 in the posterior part of the capsule. The LiA group receives local infiltration with 50 cc ropivacaine 0.2% in the anterior capsule and 50 cc ropivacaine 0.2% in the posterior capsule. And the third injection in the subcutis with 50 cc ropivacaine 0.2% without epinephrine to prevent tissue necrosis. Patients meeting the inclusion criteria will receive information about the study and informed consent letters at the anesthesiology outpatients department. The eligible patients receive a call a week after visiting the outpatients department to obtain permission. Informed consent documents will be signed before surgery by the patient and the investigator/anesthesiologist. Patients will be randomized just before the surgery to the SFNB or LiA group.

All patients will receive the same pain medication operatively: acetaminophen 1000 mg Q?D and naproxen BID and a single dose of intravenous metamizol 1000 mg (dipyrone). Moreover all patients receive a patient controlled analgesia pump with morphine. Every morning at 10.00 am the painconsulent and/or the researcher will note how much morphine is consumed. At the post-anesthesia care unit (PACU) the patients were instructed that no pain and worst possible pain equals to 0 and 10 respectively, on the visual analog scale (VAS). This VAS score is measured at the PACU until day 3 after surgery, and is noted by the patient in their patient journal during rest, flexion exercises and during walking. Side effects like nausea, vomiting, drowsiness, pruritus, paresthesia (day 1 to 3) and obstipation (at discharge) are registered by the patient in their patient journal as well as by the clinician in the medical file in EPIC.

A total of 90 patients will be included by interviewing them at the time of intake and asking informed consent. Fewer calculations showed that 80 patients are needed to show a difference in opioid consumption

Doel van het onderzoek

This prospective randomized controlled trial designed to compare the quality of analgesia offered by SFNB and LiA and their effects on morphine consumption, mobilization and pain control postoperatively. Our first hypothesis concerning opioid use is a significantly reduced opioid use in the SFNB group. Secondly, we hypothesize that with respect to the mobilization there is no significant difference between SFNB en LIA group

Onderzoeksproduct en/of interventie

Patients in the SFNB group will get a femoral nerve block with ropivacaine 0,375% and 75 microgram clonidine, and furthermore will receive 50cc ropivacaine 0,2% with epinephrine 1:100.000 in the posterior part of the capsule. The LiA group will receive 3 local infiltrations: 50cc ropivacaine 0,2% in the anterior capsule and 50cc ropivacaine 0,2% in the posterior capsule, and the third injection in the subcutis with 50cc ropivacaine 0,2% without epinephrine to prevent tissue necrosis. Ropivacaine is a local anestheticum which is favorable considering its low cardiovascular and neurologic toxicity comparing to bupivacaine. Moreover it has less motor blokkade compared to bupivacaine.

Contactpersonen

Publiek

OLVG West

Marlou Vogel
Postbus 9243

Amsterdam 1006 AE
The Netherlands

Wetenschappelijk

OLVG West

Marlou Vogel
Postbus 9243

Amsterdam 1006 AE
The Netherlands

Deelname eisen

Belangrijkste voorwaarden om deel te mogen nemen (Inclusiecriteria)

Patients 18 years and older who are scheduled for a TKA with spinal anesthesia.

- age above 18 years
- mentally competent
- eligible for TKA

Belangrijkste redenen om niet deel te kunnen nemen (Exclusiecriteria)

- contra indication for spinal anesthesia(severe aortic stenosis, severely compromised cardiac function
- infection at interspace voor spinal injection
- allergy to used medication
- repeat TKP
- all previous surgeries concerning arthrotomy
- participation in other research protocol

Onderzoeksopzet

Opzet

Type:	Interventie onderzoek
Onderzoeksmodel:	Parallel
Toewijzing:	Gerandomiseerd
Blinding:	Open / niet geblindeerd
Controle:	N.v.t. / onbekend

Deelname

Nederland
Status: Werving gestart
(Verwachte) startdatum: 01-06-2015
Aantal proefpersonen: 90
Type: Verwachte startdatum

Ethische beoordeling

Positief advies
Datum: 21-04-2016
Soort: Eerste indiening

Registraties

Opgevolgd door onderstaande (mogelijk meer actuele) registratie

ID: 42184
Bron: ToetsingOnline
Titel:

Andere (mogelijk minder actuele) registraties in dit register

Geen registraties gevonden.

In overige registers

Register	ID
NTR-new	NL5736
NTR-old	NTR5881
CCMO	NL51548.100.15
OMON	NL-OMON42184

Resultaten