

Tonsillectomy versus outpatient lasertonsillotomy in adult patients with tonsil related disease.

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When conservative treatment fail in patients with tonsil related complaints, a tonsillectomy using the classical dissection technique can be performed. In adults substantial morbidity is reported following classical tonsillectomy under general...

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

Samenvatting

ID

NL-OMON26334

Bron

Nationaal Trial Register

Verkorte titel

'SMOKE-protocol'

Aandoening

tonsillotomy
carbondioxidelaser
lokal anesthesia
adults

tonsillotomie
carbondioxidelaser
lokale anesthesie
volwassenen

Ondersteuning

Primaire sponsor: Maatschap KNO Haga Ziekenhuis / MC Haaglanden

Overige ondersteuning: not applicable

Onderzoeksproduct en/of interventie

Uitkomstmaten

Primaire uitkomstmaten

Presence or absence of the complaints for which the patient has undergone surgery (chronic / recurrent tonsillitis, tonsillolithiasis, halitosis).

Toelichting onderzoek

Achtergrond van het onderzoek

The purpose of this study is to demonstrate the effectiveness of treatment using the CO2-laser in one selected group of patients compared with a classical tonsillectomy. We also compare peri-operative and post-operative morbidity, pain, complications and patient satisfaction.

Doel van het onderzoek

When conservative treatment fail in patients with tonsil related complaints, a tonsillectomy using the classical dissection technique can be performed. In adults substantial morbidity is reported following classical tonsillectomy under general anesthesia. An interesting alternative treatment for a specific selection of adult patients could be the CO2-lasertonsillotomy under local anesthesia in an outpatient clinical setting. Several articles describe good treatment results and a decrease in perioperative and post-operative morbidity. Our hypothesis is that CO2-lasertonsillotomy is effective and there are better secondary outcome results.

Onderzoeksopzet

After an intake interview and randomisation, an appointment will be made for the intervention. The laser treatment is performed in the outpatient operating room, the traditional tonsillectomy in the operating room on an outpatient / short clinical stay. Two weeks after surgery, the patient is seen again to check the wound, then a questionnaire will also be taken. After 6 months, 1 and 2 years there will also be an evaluation and a questionnaire will be administered again. This is sent by email or taken by telephone. We also ask our patients to contact the clinic / researcher when tonsil related complaints exist during the study period.

Onderzoeksproduct en/of interventie

Study treatment:

The CO₂-lasertonsillotomies take place in our outpatient department that met the criteria for performing laser treatments. Patient will be instructed to take 1000 mg Paracetamol one hour prior to the procedure. Both the physician and the patient wear safety laser goggles and outside the room a warning lamp is clearly visible while the laser is in operation.

The patient is half lying. Subsequently, the superior, lateral and anterior parts of the tonsillar pillars will be infiltrated bilaterally with Xylocaine 2% and Adrenaline 1: 80,000. The F125 laser tube by Lumenis will be used with the laser in the continuous wave mode of operation and a beam diameter of 3 mm. Depending on the tonsil size, the power can be raised to 29 watts. With a tongue blade the tonsil will be presented and the tonsil surface is evaporated in a continuous sweeping motion. This act repeated layer by layer until a total cryptolysis occurred. The patient is instructed to hold his breath during activation of the laser and to exhale slowly after deactivation, to avoid inhaling the resulting smoke. On average a patient can hold his breath for 45 seconds (range 8-98 seconds). During the procedure the resulting smoke was continuously aspirated using a smoke evacuator. When a persistent local bleeding emerged, bipolar coagulation was used.

Control treatment:

The classical tonsillectomy will be planned in daycare or short clinical stay which is possible in all participating centra. Before the operation patients get a peripheral infuse. In the operation room the patient receives general anesthesia. The patient is placed in supine position after which the patient is intubated. The mouth is opened using a mouth gag. An Alyss clip will be attached to the superior pole of the tonsil. Then an incision is made through the anterior pillar of the tonsil to view the underlying tonsillar capsule. The incision is made close to the anterior fold and will be extended through the mucosa to the base of the tonsil. The space can be enlarged using scissors if necessary. Using a tonsil pliers the tonsil will be removed. Gauze are used to stop the bleeding. After 5 minutes we remove the gauzes and check whether the wound is dry and, if necessary bleeding can be coagulated. The mouth gag is removed if the wound is dry. After surgery, the patient will be transported to the recovery and then to the day care unit. The anaesthesiologist will decide on post-operative pain medication / anti-emetics if necessary.

Contactpersonen

Publiek

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Wetenschappelijk

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Deelname eisen

Belangrijkste voorwaarden om deel te mogen nemen (Inclusiecriteria)

1. Age > 18 years;
2. Tonsilrelated complaints with an indication for intervention (chronic/rec. tonsillitis, tonsillolithiasis, hallitosis).

Belangrijkste redenen om niet deel te kunnen nemen (Exclusiecriteria)

1. Not cooperative / restless;
2. Unable to open the mouth for a longer period;
3. Presence of a strong gag reflex;
4. History of peritonsillar abcess;
5. Estimated duration of treatment > 30 min (based on tonsilsize and cooperation);
6. Immunocompromised;
7. Hemorrhagic diathesis;
8. Cardiac history.

Onderzoeksopzet

Opzet

Type:	Interventie onderzoek
Onderzoeksmodel:	Anders
Toewijzing:	Gerandomiseerd
Blinding:	Open / niet geblindeerd
Controle:	Actieve controle groep

Deelname

Nederland	
Status:	Werving gestart
(Verwachte) startdatum:	22-08-2011
Aantal proefpersonen:	470
Type:	Verwachte startdatum

Ethische beoordeling

Positief advies	
Datum:	01-08-2011
Soort:	Eerste indiening

Registraties

Opgevolgd door onderstaande (mogelijk meer actuele) registratie

Geen registraties gevonden.

Andere (mogelijk minder actuele) registraties in dit register

Geen registraties gevonden.

In overige registers

Register	ID
NTR-new	NL2867
NTR-old	NTR3010
Ander register	METC Zuidwest Holland : 11-084
ISRCTN	ISRCTN wordt niet meer aangevraagd.

Resultaten

Samenvatting resultaten

N/A