# Radio surgery versus 80% phenol for toe nail matrix ablation: a randomised comparison study

Submission date	Recruitment status	Prospectively registered
29/09/2006	No longer recruiting	Protocol
Registration date	Overall study status	Statistical analysis plan
29/09/2006	Completed	Results
Last Edited	Condition category	[] Individual participant data
19/08/2015	Surgery	<ul><li>Record updated in last year</li></ul>

#### **Plain English Summary**

Not provided at time of registration

# Contact information

# Type(s)

Scientific

#### Contact name

Mr Ian Robinson

#### Contact details

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Dean Clarke House
Southernhay East
Exeter
United Kingdom
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# Additional identifiers

**EudraCT/CTIS** number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers N0620171683

# Study information

#### Scientific Title

Radio surgery versus 80% phenol for toe nail matrix ablation: a randomised comparison study

#### Study hypothesis

Does the wound heal faster when 80% phenol is used to destroy the nail-producing cells or is healing faster when radio surgery is used to destroy the nail-producing cells?

#### Ethics approval required

Old ethics approval format

#### Ethics approval(s)

Not provided at time of registration

#### Study design

Single-centre blinded randomised comparative trial

#### Primary study design

Interventional

## Secondary study design

Randomised controlled trial

#### Study setting(s)

Hospital

## Study type(s)

Treatment

#### Participant information sheet

Not available in web format, please use the contact details below to request a patient information sheet

#### Condition

Surgery: Toenail matrix ablation

#### **Interventions**

60 participants (patients who have been referred for nail surgery) will be allocated an ID number. 30 randomised to 80% phenol and 30 will receive radio surgery technique for nail matrix ablation using a random number code.

#### Intervention Type

Drug

#### Phase

Not Applicable

## Drug/device/biological/vaccine name(s)

Phenol

#### Primary outcome measure

Healing time of the wound (to the nearest week). Pain experienced at each return visit following surgery measured using a visual scale, post-operative infection incidence measured by clinical signs and symptoms, nail regrowth incidence measured by clinical signs and symptoms, time to nail regrowth following surgery.

## Secondary outcome measures

Not provided at time of registration

#### Overall study start date

01/09/2005

#### Overall study end date

06/10/2007

# **Eligibility**

#### Participant inclusion criteria

- 1. Subjects will be males and females
- 2. Aged 18-80 years
- 3. They will be individuals referred to the podiatry department at the Royal Devon and Exeter Hospital (Heavitree) for nail surgery under local anaesthesia
- 4. They will present with a condition indicating the need for nail surgery including the following conditions:
- 4.1 Ingrowing toenail
- 4.2 Involuted nail
- 4.3 Fungal infection of the nail
- 4.4 Thickening of the nail
- 4.5 Severe nail hypertrophy
- 5. Each subject must be able to attend for follow-up visits including a 6-month visit and be able to provide informed consent
- 6. Participants will be happy to have their toe photographed and will be happy to be contacted by telephone after 1 year

# Participant type(s)

Patient

# Age group

Adult

# Lower age limit

18 Years

# Upper age limit

80 Years

#### Sex

Both

# Target number of participants

2 groups of 30

#### Participant exclusion criteria

- 1. Fitted with a pacemaker, artificial heart valve, artificial joints or any other type of implant because these are contra-indications to radiosurgery
- 2. Subungual exostosis because treatment other than nail surgery is required for this condition
- 3. Contraindication to anaesthesia or the procedure according to standard guidelines
- 4. Pregnancy or breastfeeding
- 5. Unable to give informed consent
- 6. Inadequate blood supply to the foot or toe

#### Recruitment start date

01/09/2005

#### Recruitment end date

06/10/2007

# Locations

#### Countries of recruitment

England

**United Kingdom** 

# Study participating centre

Podiatry

Exeter United Kingdom EX1 1PQ

# Sponsor information

#### Organisation

Record Provided by the NHSTCT Register - 2006 Update - Department of Health

#### Sponsor details

The Department of Health Richmond House 79 Whitehall London United Kingdom SW1A 2NL +44 (0)20 7307 2622 dhmail@doh.gsi.org.uk

#### Sponsor type

Government

#### Website

http://www.dh.gov.uk/Home/fs/en

# Funder(s)

# Funder type

Government

#### **Funder Name**

Exeter Primary Care Trust (UK), NHS R&D Support Funding

# **Results and Publications**

## Publication and dissemination plan

Not provided at time of registration

Intention to publish date

Individual participant data (IPD) sharing plan

## IPD sharing plan summary

Not provided at time of registration