A double-blind randomised controlled trial comparing porcine dermal collagen patch saphenoplasty with conventional saphenofermoral ligation to prevent recurrent saphenofemoral incompetence

Submission date 21/10/2010	Recruitment status No longer recruiting	Prospectively registeredProtocol
Registration date 21/10/2010	Overall study status Completed	 Statistical analysis plan Results
Last Edited 18/04/2017	Condition category Surgery	 Individual participant data Record updated in last year

Plain English summary of protocol Not provided at time of registration

Contact information

Type(s) Scientific

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Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Study information

Scientific Title

A double-blind randomised controlled trial comparing porcine dermal collagen patch saphenoplasty with conventional saphenofermoral ligation to prevent recurrent saphenofemoral incompetence

Study objectives

The study aims to compare the use of porcine dermal collagen patch saphenoplasty with standard saphenofemoral flush ligation in primary and recurrent varicose vein surgery. Post-operative subjective and objective measures of presence and severity of varicosity recurrence shall be compared. This shall be the first study to establish the efficacy of Permacol™ in this setting, and shall use the following endpoints to test the given null hypothesis.

Ethics approval required

Old ethics approval format

Ethics approval(s) MREC approved, ref: 06/Q1105/27

Study design Multicentre non-randomised observational case-control study

Primary study design Observational

Secondary study design Case-control study

Study setting(s) GP practice

Study type(s) Diagnostic

Participant information sheet

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

Health condition(s) or problem(s) studied

Topic: Generic Health Relevance and Cross Cutting Themes; Subtopic: Generic Health Relevance (all Subtopics); Disease: Surgery

Interventions

1. Imaging investigations (non-radiation): Patients need to undergo a handheld Doppler ultrasound examination of the saphenofemoral junction to document the presence of radiologically apparent recurrence.

2. Other examinations: Patients shall be clinically examined for evidence of recurrent varicose veins

3. Questionnaire: A research nurse assessor, blinded to the technique used, shall administer a questionnaire at 1 and 5 years follow-up

Intervention Type

Procedure/Surgery

Phase Not Applicable

Primary outcome measure

50% reduction in the clinical presence of recurrence at 5 years.

Secondary outcome measures

- 1. Annual clinical recurrence rates
- 2. Local complication rate fistula, haematoma, groin wound infection
- 3. Radiographically demonstrable recurrence rates at 1 year and 5 years
- 4. Serious adverse event rates
- 5. 36-item short form health survey (SF-36) and EUROQOL questionnaire scores
- 6. Visual analogue pain scores

Overall study start date 10/07/2006

10/07/2006

Completion date 09/07/2011

Eligibility

Key inclusion criteria Not provided at time of registration

Participant type(s) Patient

Age group Not Specified

Sex Not Specified

Target number of participants Planned sample size: 100

Key exclusion criteria Not provided at time of registration

Date of first enrolment 10/07/2006

Date of final enrolment 09/07/2011

Locations

Countries of recruitment England

United Kingdom

Study participating centre Diana, Princess of Wales Hospital Grimsby United Kingdom DN33 2BA

Sponsor information

Organisation Northern Lincolnshire and Goole Hospitals NHS Foundation Trust (UK)

Sponsor details

Research and Development Department Scunthorpe General Hospital Cliff Gardens Scunthorpe England United Kingdom DN15 7BH

Sponsor type Hospital/treatment centre

Website http://www.nlg.nhs.uk

ROR https://ror.org/01ep18d71

Funder(s)

Funder type Industry **Funder Name** Tissue Science Laboratories PLC (UK)

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