# Bleb Dysesthesia Trial

Submission date	Recruitment status	Prospectively registered
30/09/2004	No longer recruiting	Protocol
Registration date	Overall study status	Statistical analysis plan
30/09/2004	Completed	Results
Last Edited	Condition category	Individual participant data
29/05/2014	Surgery	[] Record updated in last year

## Plain English summary of protocol

Not provided at time of registration

## Contact information

Type(s)

Scientific

Contact name

Mr John Salmon

#### Contact details

John Radcliffe Hospital Headley Way Headington Oxford United Kingdom OX3 9DU

## Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

**Secondary identifying numbers** N0203125674; N0199129912; N0176115690

## Study information

Scientific Title

A randomised trial of the effect of bleb site on eye comfort following trabeculectomy

#### **Study objectives**

To investigate long-term post-operative eye discomfort following glaucoma surgery; in particular whether two different wound locations and therefore eye fluid drainage, differ in the level of eye comfort experienced as measured by a validated questionnaire, the 'Glaucoma Symptom Scale'.

On 14/07/2011 the following changes were made to this trial record:

- 1. The end date was changed from 31/12/2008 to 30/06/2010.
- 2. The target number of participants was changed from 40 to 150.
- 3. The trial was originally a multi-centre trial but due to recruitment issues at other centres the Oxford site proceeded single-handedly.

## Ethics approval required

Old ethics approval format

#### Ethics approval(s)

Oxford Research Ethics Committee B, 10/09/2002, ref: C02.200

#### Study design

Randomised controlled trial

#### Primary study design

Interventional

## Secondary study design

Randomised controlled trial

## Study setting(s)

Hospital

## Study type(s)

Other

## Participant information sheet

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

## Health condition(s) or problem(s) studied

Surgery: Trabeculectomy

#### **Interventions**

The study design is a randomised trial designed to compare two accepted variations in surgical techniques of trabeculectomy and its effect on postoperative eye comfort. Patients who are undergoing first trabeculectomy will be approached to participate in the study. Patients will either be recruited in person or by invitation letter. When the patient gives written consent, they are randomised either to superior trabeculectomy or superonasal trabeculectomy. At the preoperative visit they will answer a 10-question self-administered questionnaire, the Glaucoma Symptom Scale. Some pre-operative data will be collected on study forms identified only by hospital number and study number.

During the trabeculectomy, the bleb will be placed below the middle upper lid, or placed below the more inner upper lid.

After the trabeculectomy, usual medical care will take place uninfluenced by their participation in this study. At the 6-month post-operative visit, the patient will be examined for the study and a second Glaucoma Symptom Scale questionnaire will be completed by the patient. The examination will record clinical findings such as visual acuity, eye pressure and eye examination findings. The study ends here and patients continue their usual care.

### Intervention Type

Procedure/Surgery

#### **Phase**

Not Applicable

#### Primary outcome measure

- 1. Eye comfort at 6 months post-surgery as measured by the Glaucoma Symptom Scale
- 2. Eye features associated with eye discomfort (dysesthesia) at 6 months post-surgery

#### Secondary outcome measures

No secondary outcome measures

#### Overall study start date

10/09/2002

### Completion date

30/06/2010

## Eligibility

#### Key inclusion criteria

- 1. Aged 18 years and over, either sex
- 2. Listed for trabeculectomy
- 3. Satisfy the inclusion criteria

#### Participant type(s)

Patient

#### Age group

Adult

#### Lower age limit

18 Years

#### Sex

Both

## Target number of participants

150

## Key exclusion criteria

Does not match inclusion criteria

#### Date of first enrolment

10/09/2002

#### Date of final enrolment

30/06/2010

## Locations

## Countries of recruitment

England

**United Kingdom** 

## Study participating centre John Radcliffe Hospital

Oxford United Kingdom OX3 9DU

## Sponsor information

#### Organisation

Department of Health

#### Sponsor details

Richmond House 79 Whitehall London United Kingdom SW1A 2NL

#### Sponsor type

Government

#### Website

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

## Funder(s)

## Funder type

## Hospital/treatment centre

#### Funder Name

Royal Devon and Exeter NHS Trust (UK)

#### Funder Name

Royal Berkshire and Battle Hospitals NHS Trust (UK)

#### Funder Name

Oxford Radcliffe Hospitals NHS Trust (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