# Arthroplasty Pain Experience (APEX) Study

Submission date	<b>Recruitment status</b>
29/04/2010	No longer recruiting
<b>Registration date</b> 29/04/2010	<b>Overall study status</b> Completed
Last Edited	<b>Condition category</b>
04/01/2016	Musculoskeletal Diseases

[] Prospectively registered

[] Protocol

- [] Statistical analysis plan
- [X] Results
- [] Individual participant data

#### Plain English summary of protocol

Not provided at time of registration

## **Contact information**

**Type(s)** Scientific

**Contact name** Miss Vicky Wylde

#### **Contact details**

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## Additional identifiers

EudraCT/CTIS number

**IRAS number** 

ClinicalTrials.gov number

Secondary identifying numbers 7664

## Study information

Scientific Title

A randomised controlled trial to determine if local wound infiltration reduces chronic pain after lower limb arthroplasty

Acronym

APEX study

#### Study objectives

Joint replacement is one of the most common elective surgical procedures performed in the NHS. However, previous research indicates that between 10 - 30% of patients experience chronic pain after hip and knee replacement. Therefore, more research is required to explore methods of minimising chronic pain after joint replacement. The aim of this study is to determine whether an injection of local anaesthethic into the hip or knee during joint replacement surgery, in addition to standard anaesthetic, reduces long-term pain.

To do this, 300 patients listed for knee replacement and 300 patients listed for hip replacement at the Avon Orthopaedic Centre will be recruited into a randomised controlled trial. Participants will complete questionnaires to assess their joint pain and function, psychological status, and use of healthcare resources. Participants will complete questionnaires before their operation, and then at regular intervals after their operation, while they are an in-patient.

Following discharge from hospital, participants will be asked to complete postal questionnaires at 3-months and 6-months after surgery. At 12-months after surgery, participants will be asked to complete a final questionnaire and undergo a joint assessment with a research nurse. Also a small number of participants and health care professionals will be interviewed about their participation in the study, in order to find out how participation in the trial affects them.

#### Ethics approval required

Old ethics approval format

**Ethics approval(s)** Southampton and South West Hampshire REC (B), 27/08/2009, ref: 09/H0504/94

**Study design** Randomised interventional treatment 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

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

Topic: Musculoskeletal; Subtopic: Musculoskeletal (all Subtopics); Disease: Musculoskeletal

#### Interventions

Participants randomised to the interventional arm of the trial will receive a local wound infiltration, in addition to the standard anaesthetic regimen during surgery. The local anaesthetic mixture will consist of 60 ml of 0.25% bupivicaine with 1 in 200,000 adrenaline.

#### Intervention Type

Other

**Phase** Not Applicable

#### Primary outcome measure

The WOMAC Pain score at 12-months post-operative.

#### Secondary outcome measures

1. Length of hospital stay

2. Daily 100mm Visual Analogue Scale (VAS) ratings while an in-patient (starting the day after surgery until discharge or day 5 after surgery)

3. Western Ontario and McMaster Universities Arthritis Index (WOMAC) Function and Stiffness Scale

4. Intermittent and Constant Osteoarthritis Pain (ICOAP) pain measure

5. Pressure pain thresholds of the volar forearm will be measured using pressure algometry preoperatively, at discharge from hospital and 12-month and post-operative

6. Resource use: Health service resource use including staff time and other resources used in the intervention, inpatient stays, outpatient visits and general practitioner visits, will be collected using hospital records and participant self-completed questionnaires (including the EQ-5D). These questionnaires will be administered at 3, 6 and 12 months post-operatively and will also be used to measure the time and travel of the patient and the carer if applicable.

#### Overall study start date

25/11/2009

**Completion date** 

01/09/2010

## Eligibility

#### Key inclusion criteria

1. Patients undergoing primary total hip replacement or primary total knee replacement for osteoarthritis at the Avon Orthopaedic Centre

Are willing and able to provide fully informed consent and complete the study questionnaires
Male and female, lower age limit of 18 years

**Participant type(s)** Patient

Age group

#### Adult

#### Lower age limit

18 Years

#### Sex

Both

#### Target number of participants

Planned Sample Size: 600

#### Key exclusion criteria

- 1. Patients undergoing revision joint replacement
- 2. Patients undergoing joint replacement for a diagnosis other than osteoarthritis
- 3. Patients under the age of 18 years

4. Patients with any medical comorbidity that precludes spinal anaesthetic, regional blocks or the use of strong analgesics postoperatively

5. Diagnosis of severe dementia or psychiatric illness such that they are unable to complete the questionnaires or provide informed consent

6. Patients undergoing stimulatenous bilateral joint replacement

7. Patients who have been in the trial for a previous joint replacement

8. Patients who are unable to understand English will be exclude because not all the validated questionnaires have been translated into languages other than English

### Date of first enrolment

25/11/2009

#### Date of final enrolment

01/09/2010

### Locations

#### **Countries of recruitment** England

United Kingdom

#### Study participating centre Southmead Hospital

Bristol United Kingdom BS10 5NB

### Sponsor information

**Organisation** Southmead Hospital (UK)

Sponsor details Southmead Road Westbury-On-Trym Bristol England United Kingdom BS10 5NB helen.lewis@nbt.nhs.uk

**Sponsor type** Hospital/treatment centre

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

ROR https://ror.org/05d576879

## Funder(s)

**Funder type** Government

**Funder Name** National Institute for Health Research (NIHR) (UK) - Central Commissioning Facility (CCF)

## **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

#### Study outputs Output type

<u>Results article</u>	results	26/02/2011		Yes	No
Results article	results	26/06/2015		Yes	No
HRA research summary			28/06/2023	No	No