

The effects of patellar tape on neural correlates during knee joint proprioception tests using fMRI: a pilot study

Submission date

29/09/2006

Recruitment status

No longer recruiting

Registration date

29/09/2006

Overall study status

Completed

Last Edited

09/08/2021

Condition category

Musculoskeletal Diseases

☐ Prospectively registered

☐ Protocol

☐ Statistical analysis plan

☒ Results

☐ Individual participant data

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s)

Scientific

Contact name

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Additional identifiers

Protocol serial number

N0453162803

Study information

Scientific Title

The effects of patellar tape on neural correlates during knee joint proprioception tests using fMRI: a pilot study

Study objectives

The principle objective is to discover if there is any increase in brain activity when subjects perform a simple proprioceptive test with and without a piece of tape across the knee cap.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Added June 2008: approved by Salford and Trafford LREC, ref 05/Q1404/17.

Study design

Pilot RCT

Primary study design

Interventional

Study type(s)

Not Specified

Health condition(s) or problem(s) studied

Musculoskeletal Diseases: Knee

Interventions

The subjects will lie in the scanner wearing shorts. A plastic block will support the thigh and knee at an angle of 45 degrees of knee inflexion. A strap will be placed over the hips to further limit head motion as a consequence of lower limb motion. Every effort will be made to control head movement by using a foam pad velcro strap and a bite bar. To ensure minimum ankle, foot and toe movements during the test, these joints will be placed in a neutral position and held by a temporary plaster cast and velcro straps.

MR scanning using the BOLD technique will be performed using a 1.5 tesla machine. The scanning protocol will be based on a functional time series. Each time series will consist of 4 blocks of 4 conditions: knee joint extension to 0deg, extension to 20deg with or without patellar tape. Each condition will last 5 minutes consisting of 30 sec on (ie the task) and 30 sec off (no task) and will be triggered by an auditory command given by using headphones customised for fMRI experiments. A metronome will pace the movements using sound in order to impose a constant timing and equal no of cycles across conditions.

Patellar taping will then be applied. The order of tape/no tape will be randomised

The acquired images will be assessed carefully for head movement and in order to correct the confounding effects induced by head movement realignment parameters. To process the data,

statistical parametric mapping will be used to display the significance of activation and to analyse functional and anatomical image. SPM is the most prevalent approach to characterising activity related changes (Friston et al 2000).

Intervention Type

Other

Phase

Not Specified

Primary outcome(s)

Neural processing in the brain as a result of application of patellar tape detected by the BOLD technique using fMRI.

Key secondary outcome(s))

No secondary outcome measures

Completion date

31/01/2007

Eligibility**Key inclusion criteria**

8 healthy volunteers between the ages of 20-40 years will have refrained from any physical exercise for 3 days prior to the testing. They will sign a consent form after reading the study information sheet and after having had the opportunity to ask questions about any aspect of the study and their role.

There is no control group. All subjects will act as their own internal controls.

Participant type(s)

Healthy volunteer

Healthy volunteers allowed

No

Age group

Adult

Sex

Not Specified

Total final enrolment

8

Key exclusion criteria

Added June 2008:

1. Histories of neurological or cardiovascular disease
2. Cochlear implants or any metal objects in the body
3. Cardiac or neural pacemakers
4. Histories of serious musculoskeletal injury in both lower limbs

Date of first enrolment

26/04/2005

Date of final enrolment

31/01/2007

Locations

Countries of recruitment

United Kingdom

England

Study participating centre

Clinical Specialist & Research Associate

Manchester

United Kingdom

M13 9PT

Sponsor information

Organisation

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

Funder(s)

Funder type

Government

Funder Name

Central Manchester and Manchester Children's University Hospitals NHS Trust (UK)

Funder Name

NHS R&D Support Funding

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Not provided at time of registration

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article		26/01/2012	09/08/2021	Yes	No