# Injury, inflammatory markers & the exacerbation of confusion: ASCRIBED

Submission date	<b>Recruitment status</b> No longer recruiting	[X] Prospectively registered		
21/11/2016		[X] Protocol		
Registration date	Overall study status Completed Condition category	Statistical analysis plan		
11/05/2017		<ul><li>☐ Results</li><li>☐ Individual participant data</li></ul>		
Last Edited				
03/10/2023	Mental and Behavioural Disorders	☐ Record updated in last year		

#### Plain English summary of protocol

Background and study aims

Dementia is an umbrella term used to refer to a wide range of symptoms linked to with a reduction in memory and/or other thinking skills which reduce a person's ability to perform everyday activities. Inflammation is generally a beneficial response to tissue damage or infection. However, when inflammation is extensive or prolonged this can damage healthy tissues and disrupt normal cellular function. Research suggests that acute illnesses or injury causing inflammation can accelerate dementia. However there are few studies which examine underlying mechanisms of how this happens in humans. This study aims to address this gap. The study will compare markers of inflammation and injury found in the blood and cerebrospinal fluid (fluid which bathes the spinal cord (CSF)) of people with and without confirmed dementia who fracture their fracture. A hip fracture is a common example of an acute injury causing an inflammatory response. People who fracture their hip will undergo an operation to repair it. A common procedure during this operation is the giving of spinal aesthetic. This involves inserting a needle into the patient's spinal space and injecting anaesthesia into CSF. This means CSF can be collected before operation via the same needle.

#### Who can participate?

Patients due to have a hip fracture operation via spinal anaesthesia with all levels of preoperative confusion.

#### What does the study involve?

Patients are allocated to one of three groups after their operation: those with confirmed dementia (as obtained from patient's GP notes/hospital records and/or carer insight), non-dementia (no evidence of dementia found in patent's GP notes/medical records) groupings; and pre-operatively confused but without confirmed dementia. Consent is gained for the storage of surplus samples in a bio-bank to help future studies. Pre-operative blood and CSF samples, post-operative blood samples and a short cognitive questionnaire is administered to all patients.

What are the possible benefits and risks of participating?

For patients and their families there are no direct benefits taking part. However it is hoped that

the research may help similar patient groups in the future. Postdural-puncture headaches (PDPH) are a common side-effect of spinal anaesthesia. The collection of CSF may slightly increase the chance of PDPH.

Where is the study run from?

This study is being run by University of East Anglia (UK) and takes place in hospitals in the UK.

When is the study starting and how long is it expected to run for? November 2016 to December 2022

Who is funding the study? Alzheimer's Research UK (UK)

Who is the main contact? Dr Simon Hammond s.hammond@uea.ac.uk

#### Contact information

#### Type(s)

Public

#### Contact name

Dr Simon Hammond

#### **ORCID ID**

http://orcid.org/0000-0002-0473-3610

#### Contact details

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#### Additional identifiers

**EudraCT/CTIS** number

**IRAS** number

ClinicalTrials.gov number

Secondary identifying numbers

1

#### Study information

#### Scientific Title

ASCRIBED: The impact of Acute SystematiC inflammation upon cerebRospinal fluId and blood BiomarkErs of brain inflammation and injury in Dementia: a study in acute hip fracture patients

#### Acronym

**ASCRIBED** 

#### Study objectives

The aim of this study is to evaluate whether hip fracture patients with dementia show elevated markers of systemic inflammation and of brain inflammation in comparison to stable patients with dementia and hip fracture patients without dementia, as measured by biomarkers in cerebrospinal fluid (CSF) and blood.

#### Ethics approval required

Old ethics approval format

#### Ethics approval(s)

NRES Committee North East - Newcastle & North Tyneside 1, 24/03/2017, ref: ISRCTN43803769

#### Study design

Observational case-control study

#### Primary study design

Observational

#### Secondary study design

Case-control study

#### 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

Dementia patients admitted to acute hospital settings with a hip fracture.

#### Interventions

Participants due to have a hip fracture operation via spinal anaesthesia with all levels of preoperative confusion (as assessed by clinical screening procedures, AMTS (England) and 4AT (Scotland)) will be approached and recruited. Capacity will be assessed and consent or consultee agreement gained. During pre-operative procedures a blood sample will be taken and cerebrospinal fluid (CSF) collected.

48 hours post-operatively patients will be approached to undertake a short cognitive test and have a second blood sample collected. A suitable informant will also be sought to complete a

proxy measure about the patient's memory and thinking. At 1 month post-operatively patient's GP and medical records will be accessed to search for evidence of dementia. Patients also taking part in ASCRIBED's sister study PERFECTED (ISRCTN99336264) will be asked to give blood samples at 1, 3 and 6 months post-operatively.

#### Intervention Type

Other

#### Primary outcome measure

CSF inflammation and injury is measured by TNF- $\alpha$ , IL-1RA, IL-1 $\beta$ , IL-6 and Neurogranin, tTau, Synaptotagmin, SNAP-25 at baseline.

#### Secondary outcome measures

Magnitude of the brain inflammation is measured by brain injury markers (phospho-Tau, NFL, neurogranin, synaptotagmin, SNAP-25) in CSF at baseline.

#### Exploratory:

1. Higher levels of inflammatory and brain injury markers in CSF and blood are associated with worsening cognitive and functional decline measured by MMSE at 6 months post-operatively 2. Cytokines ratio CSF:blood pre-op time will show higher ratios in dementia than non-dementia patients measures in blood and CSF at baseline

#### Overall study start date

01/11/2016

#### Completion date

31/12/2022

#### **Eligibility**

#### Key inclusion criteria

Group 1: Pre-operative acute hip fracture patients with confusion as defined by the Abbreviated Mental Test (England) or 4AT (Scotland):

Inclusion Criteria:

- 1. Patient must have had a confirmed proximal hip fracture requiring an operation and be aged 60 or older at the time of operation
- 2. Patient has a pre-operative Abbreviated Mental Test score of 8 or below or 4AT score of 1 or above
- 3. Patient must be undergoing spinal anaesthesia

Group 2: Pre-operative acute hip fracture patients without confusion as defined by the Abbreviated Mental Test (England) or 4AT (Scotland):

**Inclusion Criteria:** 

- 1. Patient must have had a confirmed proximal hip fracture requiring an operation and be aged 60 or older at the time of operation
- 2. Pre-operative Abbreviated Mental Test score of 9 or above or 4AT score of 0
- 3. Patient must be undergoing spinal anaesthesia

#### Participant type(s)

Mixed

#### Age group

Adult

#### Sex

Both

#### Target number of participants

400

#### Total final enrolment

469

#### Key exclusion criteria

Group 1 and 2:

- 1. Decision taken not to have hip surgery
- 2. Patient has head trauma with bleeding as indicated by a CT scan
- 3. Patient has confirmed diagnosis of Parkinson's disease
- 4. Patient not expected to survive beyond 4 weeks

#### Date of first enrolment

01/06/2017

#### Date of final enrolment

31/08/2019

#### Locations

#### Countries of recruitment

England

Scotland

United Kingdom

#### Study participating centre Royal Infirmary of Edinburgh

51 Little France Crescent Edinburgh United Kingdom EH16 4SA

#### Study participating centre Leicester Royal Infirmary

Infirmary Square Leicester United Kingdom LE1 5WW

#### Study participating centre Royal Sussex County Hospital

Eastern Road Brighton United Kingdom BN2 5BE

#### Study participating centre Princess Royal University Hospital

Farnborough Common Orpington United Kingdom BR6 8ND

#### Study participating centre Royal Derby Hospital

Uttoxeter Road Derby United Kingdom DE22 3NE

#### Study participating centre Pinderfields Hospital

Mid Yorkshire Hospitals NHS Trust Wakefield United Kingdom WF1 4DG

# Study participating centre York Teaching Hospital NHS Foundation Trust York United Kingdom YO31 8HE

#### Study participating centre

#### **Princess Royal Hospital**

Orpington United Kingdom BR6 8ND

## Study participating centre James Paget University Hospital

Great Yarmouth United Kingdom NR31 6LA

#### Study participating centre Russells Hall Hospital

The Dudley Group NHS Foundation Trust Dudley United Kingdom DY1 2HQ

#### Study participating centre Peterborough City Hospital

North West Anglia NHS Foundation Trust Peterborough United Kingdom PE3 9GZ

#### Study participating centre Queen's Medical Centre Campus

Nottingham University Hospitals NHS Trust Nottingham United Kingdom NG7 2UH

#### Study participating centre Royal Blackburn Hospital

East Lancashire Hospitals NHS Trust Blackburn United Kingdom BB2 3HH

### Study participating centre University Hospital of North Tees

North Tees and Hartlepool NHS Foundation Trust Stockton-on-Tees United Kingdom TS19 8PE

## Study participating centre The Royal Shrewsbury Hospital

The Shrewsbury and Telford Hospital NHS Trust Shrewsbury United Kingdom SY3 8XQ

#### Study participating centre Doncaster Royal Infirmary

Doncaster and Bassetlaw Teaching Hospitals NHS Foundation Trust Doncaster United Kingdom DN2 5LT

# Study participating centre Royal United Hospital Bath NHS Foundation Trust Bath United Kingdom BA1 3NG

Study participating centre

#### University Hospital of North Durham County Durham and Darlington NHS Foundation Trust Durham United Kingdom DH1 5TW

#### Study participating centre Royal Bolton Hospital Bolton NHS Foundation Tr

Bolton NHS Foundation Trust Bolton United Kingdom BL4 0JR

#### Study participating centre Kingston Hospital

Kingston Hospital NHS Foundation Trust Kingston United Kingdom KT2 7QB

#### Study participating centre Royal Stoke University Hospital

University Hospitals of North Midlands NHS Trust Stoke-on-Trent United Kingdom ST4 6QG

## Study participating centre Countess of Chester Hospital NHS Foundation Trust

Chester United Kingdom CH2 1UL

#### Study participating centre Western Sussex Hospitals NHS Foundation Trust

United Kingdom

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#### Study participating centre Medway Maritime Hospital

Medway NHS Foundation Trust Gillingham United Kingdom ME7 5NY

## Study participating centre Whiston Hospital

St Helens & Knowsley Teaching Hospitals NHS Trust Rainhill United Kingdom L35 5DR

#### Study participating centre Southampton General Hospital

University Hospital Southampton NHS Foundation Trust Southampton United Kingdom SO16 6YD

#### Study participating centre Royal Devon & Exeter Hospital

Royal Devon and Exeter NHS Foundation Trust Wonford United Kingdom EX2 5DW

#### Study participating centre Basildon University Hospital

Basildon and Thurrock University Hospitals NHS Foundation Trust Basildon United Kingdom SS16 5NL

#### Study participating centre Musgrove Park Hospital

Taunton and Somerset NHS Foundation Trust Taunton United Kingdom TA1 5DA

#### Study participating centre Scunthorpe General Hospital

Northern Lincolnshire and Goole NHS Foundation Trust Scunthorpe United Kingdom DN15 7BH

## Study participating centre Royal Lancaster Infirmary

University Hospitals of Morecambe Bay NHS Trust

Lancaster United Kingdom LA1 4RP

**BA21 4AT** 

# Study participating centre Yeovil District Hospital Yeovil District Hospital NHS Foundation Trust Yeovil United Kingdom

# Study participating centre Watford General Hospital West Hertfordshire Hospitals NHS Trust Watford United Kingdom WD18 0HB

# Study participating centre Sandwell General Hospital Sandwell and West Birmingham Hospitals NHS Trust West Bromwich United Kingdom B71 4HJ

#### Sponsor information

#### Organisation

University of East Anglia

#### Sponsor details

University of East Anglia Norwich Research Park Norwich. England United Kingdom NR4 7TJ +44 1603 591460 t.moulton@uea.ac.uk

#### Sponsor type

University/education

#### **ROR**

https://ror.org/026k5mg93

#### Funder(s)

#### Funder type

Charity

#### **Funder Name**

Alzheimer's Research UK

#### Alternative Name(s)

Alzheimer's Research Trust, AlzheimersResearch UK, AlzResearchUK, ARUK

#### **Funding Body Type**

Private sector organisation

#### **Funding Body Subtype**

Trusts, charities, foundations (both public and private)

#### Location

United Kingdom

#### **Results and Publications**

#### Publication and dissemination plan

Planned publication in a high-impact peer reviewed journal.

#### Intention to publish date

30/04/2023

#### Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study are/will be available upon request from Prof Chris Fox, Norwich Medical School, University of East Anglia, Norwich Research Park, Norwich, NR4 7TJ, United Kingdom.

#### IPD sharing plan summary

Available on request

#### Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Protocol article	protocol	07/09/2019	09/12/2020	Yes	No