# The clinical and cost effectiveness of of a steroid injection versus a night splint for Carpal Tunnel Syndrome



#### **Plain English Summary**

Background and study aims

Carpal Tunnel Syndrome (CTS) is a common condition in which a nerve (known as the median nerve) is squeezed where it passes through the wrist. It can cause pain or aching, tingling or numbness in the affected hand. It may disturb sleep, or affect ability to do day to day things. There have been several studies into the best treatment of patients with severe symptoms of CTS who are referred to a hospital for treatment. However, little is known about the best treatments for patients with mild to moderate symptoms who visit their GP but do not require hospital treatment. This study aims to find out whether a single steroid injection is effective in treating CTS symptoms when compared with a night splint in people suffering with mild to moderate carpal tunnel syndrome.

Who can participate?

Patients aged 18 and over who have been diagnosed with mild to moderate CTS which has been present for at least 6 weeks

What does the study involve?

Each participant is randomly allocated to receive either a single steroid injection or a splint, and is asked to complete up to five questionnaires over 2 years. The steroid is a drug called DepoMedrone and is already widely used to treat CTS. The splint is made of elastic and has an aluminium bar which sits on the palm of the hand. In this study, the splint will be worn at night for 6 weeks. We study the effects of these two treatments over 6 weeks and at 6 months. We also look at whether these 6 weeks of treatment are effective 1 year and 2 years later.

What are the possible benefits and risks of participating? Not provided at time of registration

Where is the study run from?

The study will take place in up to 50 GP practices and hospital clinics across the UK

When is the study starting and how long is it expected to run for? April 2014 to September 2017

Who is funding the study? Arthritis Research UK

Who is the main contact? Jacqueline Gray j.gray@keele.ac.uk

### **Contact information**

**Type(s)** Scientific

**Contact name** Ms Jacqueline Gray

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## Additional identifiers

**EudraCT/CTIS number** 2013-001435-48

**IRAS number** 

ClinicalTrials.gov number NCT02038452

**Protocol/serial number** 16390

## Study information

#### Scientific Title

The clinical and cost effectiveness of of a steroid injection versus a night splint for Carpal Tunnel Syndrome: a pragmatic randomised trial in primary care

#### Acronym

INjection versus SplinTing in Carpal Tunnel Syndrome (INSTinCTS)

#### Study hypothesis

The study aims to find out whether a single steroid injection is effective in treating CTS symptoms when compared with a night splint in people suffering with mild to moderate carpal tunnel syndrome.

### Ethics approval required

Old ethics approval format

Ethics approval(s) 13/NW/0280; First MREC approval date 07/05/2013

**Study design** Randomised; Interventional; Design type: Treatment

**Primary study design** Interventional

**Secondary study design** Randomised controlled trial

**Study setting(s)** GP practice

### Study type(s)

Treatment

#### Participant information sheet

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

#### Condition

Topic: Primary Care, Musculoskeletal disorders; Subtopic: Not Assigned, Musculoskeletal (all Subtopics); Disease: All Diseases, Musculoskeletal Pain Disorders

#### Interventions

Each participant will receive either a single steroid injection or a splint. The steroid is a drug called DepoMedrone 20mg. This drug is already widely used to treat CTS. In this study, one injection will be given. The splint is made of elastic and has an aluminium bar which sits on the palm of the hand. In this study, the splint will be worn at night for 6 weeks. Each participant will be asked to complete up to 5 questionnaires over 2 years. We will study the effects of these 2 treatments over 6 weeks and at 6 months. Subject to further funding, the Study will also look at whether these 6 weeks of treatment are effective 1 year and 2 years later.

Intervention Type

**Phase** Phase IV

Drug/device/biological/vaccine name(s)

#### Depo-medrone

#### Primary outcome measure

Symptom severity and limitations in hand function as assessed by the Boston CTS questionnaire; Timepoint(s): 6 weeks, 6 months, 12 months and 24 months post-randomisation.

#### Secondary outcome measures

Not provided at time of registration

## Overall study start date

18/12/2009

### Overall study end date

31/12/2018

## Eligibility

#### Participant inclusion criteria

1. Male or female aged 18 years or over

2. A clinical diagnosis of unilateral or bilateral CTS as made by a GP or trained clinician according to the diagnostic criteria

- 3. Mild (e.g. intermittent paraesthesia) or moderate (e.g. constant paraesthesia, reversible numbness and / or pain) severity CTS of idiopathic nature
- 4. Symptom duration of episode of at least 6 weeks
- 5. Written informed consent provided by the patient, prior to any trial specific procedures

### Participant type(s)

Patient

#### Age group

Adult

#### Lower age limit

18 Years

#### **Sex** Both

### Target number of participants

Planned Sample Size: 240; UK Sample Size: 240

#### Participant exclusion criteria

1. Steroid injection or night splints for CTS in the affected wrist within preceding 6 months

2. Any previous surgery on the affected wrist

3. Severe CTS exhibiting constant numbness or pain, constant sensory loss, severe thenar muscle atrophy or symptom

severity which requires the patient to be referred for a surgical opinion

- 4. Clinical suspicion of local or systemic sepsis or infection
- 5. Current or previous infection of the affected wrist
- 6. Trauma to the affected hand requiring surgery or immobilisation in the previous 12 months

7. Unable to tolerate the study interventions

8. Unable to understand and complete self report questionnaires written in English 9. Intercurrent illness including, but not limited to: poorly controlled thyroid disease, poorly controlled diabetes mellitus, vibration-induced neuropathy, inflammatory joint disease, suspected complex neurological conditions, any other severe medical illness which in the opinion of the local Principal Investigator (or other authorised clinical delegate) precludes trial participation

10. Pregnant or lactating females

11. Receiving anticoagulants

12. Any history of hypersensitivity to DepoMedrone or any of its excipients (refer to the Summary of Product Characteristics (SPC)

13. Allergy to any of the splint materials (refer to manufacturers specification)

14. Known abuse of drugs or alcohol

15. Involved in ongoing litigation cases for their condition

**Recruitment start date** 17/04/2014

Recruitment end date 01/09/2017

### Locations

**Countries of recruitment** England

United Kingdom

### Study participating centre

**Keele University** Newcastle-Under-Lyme United Kingdom ST5 5BG

### Sponsor information

#### **Organisation** University of Keele (UK)

### Sponsor details

Keele Newcastle England United Kingdom ST5 5BG **Sponsor type** University/education

ROR https://ror.org/00340yn33

### Funder(s)

**Funder type** Charity

**Funder Name** Arthritis Research UK (UK)

Alternative Name(s)

**Funding Body Type** Private sector organisation

**Funding Body Subtype** Other non-profit organizations

**Location** United Kingdom

## **Results and Publications**

**Publication and dissemination plan** Planned publication in a high-impact peer reviewed journal.

Intention to publish date 01/09/2018

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

The datasets generated during and/or analysed during the current study will be available upon request from Dr Linda Chesterton, l.s.chesterton@keele.ac.uk

#### IPD sharing plan summary

Available on request

Study outputs					
Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Protocol article	protocol	06/10/2016		Yes	No
Results article	results	20/10/2018		Yes	No

HRA research summary

28/06/2023 No

No