

Clinical study of a novel anatomic cuff for forearm crutches

Submission date 07/02/2017	Recruitment status No longer recruiting	☐ Prospectively registered ☐ Protocol
Registration date 14/02/2017	Overall study status Completed	☐ Statistical analysis plan ☒ Results
Last Edited 16/03/2017	Condition category Musculoskeletal Diseases	☐ Individual participant data

Plain English summary of protocol

Background and study aims

Osteoarthritis (OA) is the most common type of arthritis and affects millions of people worldwide. It occurs when the protective cartilage on the end of bones wears away. The bones then rub against one another, causing stiffness, pain and a reduction in the range of movement. In some, movement can be limited so severely that they need to use walking aids such as crutches to get around. Using forearm walking aids (crutches that attach at the forearm) often leads to pain in the forearms. Special handles called anatomic handles have been developed to increase the size of the contact area and reduce the pressure, but for many this is ineffective. One third of the load is transferred via the forearm. In contrast to the hand, the ulnar (outer, lower arm bone) is not well protected and this can often lead to compression of the ulna nerve, leading to pain. This study aims to look at the effectiveness of a newly developed anatomic cuff for forearm crutches which aims to protect the ulnar bone and distributing the load towards surrounding muscles and soft tissue.

Who can participate?

Patients with OA who have used forearm crutches on both sides for at least eight weeks.

What does the study involve?

All participants are given the special crutches with the anatomic cuff to use for four weeks. The length of the crutches is adjusted depending on participant height. At the start of the study and then after four weeks, participants complete questionnaires to assess their forearm pain and general health.

What are the possible benefits and risks of participating?

Participants may benefit from lower pain levels in their forearms than if they used conventional crutches. There are no notable risks involved with participating.

Where is the study run from?

University Hospital Basel (Switzerland)

When is the study starting and how long is it expected to run for?

January 2014 to December 2016

Who is funding the study?
University Hospital Basel (Switzerland)

Who is the main contact?
Professor Thomas Hügle
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Contact information

Type(s)

Public

Contact name

Prof Thomas Hügle

Contact details

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Additional identifiers

Protocol serial number

2014/1

Study information

Scientific Title

Prospective clinical evaluation of a novel anatomic cuff for forearm crutches in patients with osteoarthritis

Study objectives

Anatomic cuffs with ulnar protection are less associated with pain and discomfort in long-term users of forearm crutches.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Ethikkommission beider Basel (EKBB) Switzerland, 03/06/2014, ref: E350/12

Study design

Prospective non-randomised study

Primary study design

Interventional

Study type(s)

Quality of life

Health condition(s) or problem(s) studied

Osteoarthritis

Interventions

At baseline, all participants receive forearm crutches with an anatomic cuff (Ulnar Pro®, Rebotec, Quakenbrück, Germany) with anatomic hand grip (model soft, Rebotec, Quakenbrück, Germany) to use for four weeks. The length of the crutches are adjusted by the study team to the patients' hand height during stance with the arms positioned at 20 to 30° elbow flexion. Clinical data are collected at baseline and at 4-weeks follow-up using questionnaires.

Intervention Type

Device

Primary outcome(s)

Pain along the forearm is measured using a questionnaire based on a unipolar 9-point Likert-scale designed for the purpose of this study at baseline and 4 weeks

Key secondary outcome(s)

General health is assessed using 36-Item Short Form Survey (SF-36) at baseline and 4 weeks

Completion date

31/12/2016

Eligibility**Key inclusion criteria**

1. Suffering from osteoarthritis
2. Have used bilateral forearm crutches for at least 8 weeks prior to the study

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Mixed

Sex

All

Key exclusion criteria

1. Infections of hands or forearms
2. Amputation
3. Neuropathy (e.g. due to diabetes or syringomyelia)
4. Active rheumatic diseases
5. Upper limb injuries

Date of first enrolment

01/03/2014

Date of final enrolment

01/10/2014

Locations

Countries of recruitment

Switzerland

Study participating centre

University Hospital Basel

Petersgraben 4

Basel

Switzerland

4045

Sponsor information

Organisation

University Hospital Basel

ROR

<https://ror.org/04k51q396>

Funder(s)

Funder type

Hospital/treatment centre

Funder Name

University Hospital Basel

Results and Publications

Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study are/will be available upon request from Thomas.huegle@unibas.ch

IPD sharing plan summary

Available on request

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	results	14/03/2017		Yes	No