

Clinical study of the volar locking plate for distal radial fractures

Submission date 17/05/2009	Recruitment status No longer recruiting	☐ Prospectively registered
		☐ Protocol
Registration date 08/09/2009	Overall study status Completed	☐ Statistical analysis plan
		☒ Results
Last Edited 07/10/2013	Condition category Musculoskeletal Diseases	☐ Individual participant data

Plain English summary of protocol
Not provided at time of registration

Contact information

Type(s)
Scientific

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Additional identifiers

Protocol serial number
07OR003

Study information

Scientific Title
Unstable fractures of the distal radius: a randomised prospective clinical study comparing their treatment with volar locking plate and conventional method

Study objectives

This study will compare the outcome of displaced distal radial fractures when treated with a volar locking plate (the Distal Volar Radius [DVR®]) or the conventional method which involves percutaneous wires +/- an external fixator.

Our primary research objective is to determine whether the use of volar locking plates improves functional outcome and allows for an earlier return to normal activities and work.

As a secondary objective, we aim to determine through economic evaluation, whether the use of volar locking plates for distal radial fractures is of financial benefit to the health service and society in general.

Ethics approval required

Old ethics approval format

Ethics approval(s)

North Nottinghamshire Research Ethics Committee approved on the 6th September 2007 (ref: 07/H0407/39)

Primary study design

Interventional

Study design

Pragmatic randomised single centre controlled parallel group surgical trial

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Distal radius fracture

Interventions

The intervention consists of surgical fixation of the distal radius fracture in a patient who fulfils the trial inclusion and exclusion criteria with a radial volar locking plate or the established conventional method involving percutaneous wires +/- an external fixator.

The health technology under assessment is the radial volar locking plate, a type of orthopaedic implant with locking screws which aid the reduction and stabilisation of distal radius fractures. We compare this to the established conventional methods (percutaneous wires and/or an external fixator device).

The plate chosen for this trial is the Distal Volar Radius or DVR® plate, which is in common use across the United Kingdom. Instrumentation also includes smooth 1.6 mm Kirschner wires and the standard AO external fixator, as appropriate. All are currently in use within the NHS, CME licensed and will not require MHRA authorisation.

The total duration of follow-up for each participant is one year post-surgery.

Intervention Type

Other

Phase

Not Applicable

Primary outcome(s)

Function of the limb following a surgically treated distal radius fracture. As a measure of function we selected the Hand Health Profile forming part of the Patient Evaluation Measure (PEM). Expected duration of patient participation will be one year. Follow up and data collection will be performed at 6 weeks, 12 weeks and 1 year.

Key secondary outcome(s)

1. Clinical measurements: grip strength and range of motion
2. Radiographic parameters: radial length, palmar tilt, radial inclination and articular gaps/steps less or equal to 2 mm
3. Quality of life: as assessed via the EUROQUOL EQ-5D and 12-item short form health survey (SF-12) scores

Expected duration of patient participation will be one year. Follow up and data collection will be performed at 6 weeks, 12 weeks and 1 year.

Completion date

30/08/2010

Eligibility**Key inclusion criteria**

1. Fractures which the referring physician considers require operative intervention
2. Configuration is such that the fracture would be amenable to stabilisation via volar locking plate (not massively comminuted)
3. Adults (skeletally mature) with high demand requirements of their wrist in whom the radiological appearance of the bone suggests that it is robust enough to tolerate internal fixation; and in whom the fracture pattern at presentation fulfils the criteria as described below
4. Fractures of the distal radius which are:
 - 4.1. Dorsally displaced extra-articular fractures (with or without an undisplaced intra-articular component) with dorsal cortical comminution as seen on the lateral radiograph
 - 4.2. Displaced intra-articular fractures with an articular step or gap in the radio-carpal joint surface
5. Skeletally mature adults, minimum age 16 years and above, either sex

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Sex

All

Key exclusion criteria

1. Patients with concomitant systemic diseases (diabetes with vascular or neurological complications, advanced cardiac, pulmonary or neurological disease)
2. Proximal metaphyseal fractures (more than one inch or 2.5 centimetres from the articular surface)
3. Open fractures
4. Smith's and volar Barton's configuration
5. Previous fractures of the distal radius of the same or contra-lateral limb less than six months old
6. Significant pre-existing radiological abnormality
7. Multiply injured
8. Bilateral injuries
9. Patients who are unable to consent for themselves to treatment
10. Patients who may have difficulties in adequate understanding of English

Date of first enrolment

11/02/2008

Date of final enrolment

30/08/2010

Locations

Countries of recruitment

United Kingdom

England

Study participating centre

Academic Department of Orthopaedic & Accident Surgery

Nottingham

United Kingdom

NG7 2UH

Sponsor information

Organisation

Nottingham University Hospitals NHS Trust (UK)

ROR

<https://ror.org/05y3qh794>

Funder(s)

Funder type

Government

Funder Name

Nottingham University Hospitals NHS Trust (UK)

Funder Name

Nottingham University Hospitals Charity via the Hand Research Fund (UK)

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	results	02/10/2013		Yes	No