Development and evaluation of an education programme in rheumatoid arthritis: impact on compliance, function, disease progression and quality of life

Submission date 23/01/2004	Recruitment status No longer recruiting	 Prospectively Protocol
Registration date 23/01/2004	Overall study status Completed	 [] Statistical ana [X] Results
Last Edited 22/02/2008	Condition category Musculoskeletal Diseases	[_] Individual par

y registered

- alysis plan
- rticipant data

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s) Scientific

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Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers 535913

Study information

Scientific Title

Study objectives

Although patient education is assuming an increasingly important role in patient management in many medical disciplines, evaluation of the medium term impact of patient education has not been carried out in rheumatological practice. We wish to know whether a patient education programme will improve compliance with drug, exercise and attendance regimes and the impact of education on disease progression, function and quality of life. We plan a randomised controlled trial of patient education in rheumatoid arthritis based on an ambulant population. We will evaluate the educational benefits of the programme over a four week period and make further outcome assessments for the 12 months following entry into the trial. We hope to show increased compliance with drug regimes and drug monitoring, retention of knowledge and orthodox attitudes to disease management, a reduction in periods of inpatient treatment and unscheduled outpatient attendance. The benefits would accrue primarily to the patients in terms of quality of life and disease progression but would contribute to the cost efficient organisation of the rheumatology department (including number of patient episodes) and in pharmacy costs (currently due, in part, to poor drug compliance and wastage).

Ethics approval required Old ethics approval format

Ethics approval(s) Not provided at time of registration

Study design Randomised controlled trial

Primary study design Interventional

Secondary study design Randomised controlled trial

Study setting(s) Not specified

Study type(s) Not Specified

Participant information sheet

Health condition(s) or problem(s) studied Musculoskeletal diseases: Arthritis (rheumatoid)

Interventions

1. Patient education programme 2. Standard care

Intervention Type

Other

Phase

Not Specified

Primary outcome measure

Compliance with drug, exercise and attendance regimes and disease progression, function and quality of life. The main outcome variables were the modified Larsen radiological score for the hands and the 36-item short form health survey (SF-36) quality of life questionnaire.

Secondary outcome measures

- 1. Health Assessment Questionnaire (HAQ)
- 2. Ritchie Articular Index (RAI)
- 3. Patient Knowledge Questionnaire (PKQ)
- 4. Compliance Questionnaire (CQ)
- 5. Plasma viscosity (PV)
- 6. Pharmaceutical changes and consulting behaviour

Overall study start date 01/01/1992

Completion date 31/12/1994

Eligibility

Key inclusion criteria Patients with rheumatoid arthritis.

Participant type(s) Patient

Age group Adult

Sex Both

Target number of participants 77

Key exclusion criteria

1. Unable to speak or read english

2. Previous education programme

Date of first enrolment 01/01/1992

Date of final enrolment 31/12/1994

Locations

Countries of recruitment England

United Kingdom

Study participating centre University of Leeds Leeds United Kingdom LS2 9NZ

Sponsor information

Organisation NHS R&D Regional Programme Register - Department of Health (UK)

Sponsor details The Department of Health Richmond House 79 Whitehall London United Kingdom SW1A 2NL +44 (0)20 7307 2622 dhmail@doh.gsi.org.uk

Sponsor type

Government

Website http://www.doh.gov.uk

Funder(s)

Funder type

Funder Name

NHS Executive Northern and Yorkshire (UK)

Results and Publications

Publication and dissemination plan Not provided at time of registration

Intention to publish date

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	01/04/1999		Yes	No