

Effects of a pain consult and patient education and monitoring: a prospective study in oncology patients

Submission date 28/04/2006	Recruitment status No longer recruiting	☐ Prospectively registered ☐ Protocol
Registration date 28/04/2006	Overall study status Completed	☐ Statistical analysis plan ☐ Results
Last Edited 10/08/2009	Condition category Signs and Symptoms	☐ Individual participant data ☐ Record updated in last year

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
NTR613; EMC 2005 - 257

Study information

Scientific Title

Study objectives

It is hypothesised that a pain consult at the specialized pain clinic in combination with Patient Education Program is more effective in reducing average pain intensity compared to a pain consult alone. A pain consult at the specialized pain clinic is more effective in reducing average pain intensity compared to standard care.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Ethics approval received from the local medical ethics committee

Primary study design

Interventional

Study design

Randomised controlled trial

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Cancer pain

Interventions

1. Standard care
2. Second opinion pain consult at the specialist pain clinic
3. Second opinion pain consult combined with Pain Education Program and monitoring by nurse specialists

Intervention Type

Other

Phase

Not Specified

Primary outcome(s)

Average pain reduction measured by numeric rating scale during the study period.

Key secondary outcome(s)

Effect of the interventions after 2, 4 and 8 weeks on:

1. Adherence to ATC analgesics
2. Worst pain reduction
3. Average pain reduction
4. Present pain reduction
5. Proportion of patients with clinically relevant pain reduction
6. Pain interference
7. Quality of life
8. Reduction of side effects

- 9. Adequacy of pain treatment
- 10. Pain knowledge

Completion date

31/12/2007

Eligibility

Key inclusion criteria

- 1. Cancer-related pain or cancer treatment related pain for at least two weeks
- 2. Nociceptive pain
- 3. Average pain intensity score of 4 or more
- 4. Accessibility by telephone
- 5. A life expectancy of at least three months
- 6. Informed consent

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Sex

All

Key exclusion criteria

- 1. Neuropathic pain
- 2. Residing in nursing home or retirement home
- 3. Pain not treated with oral medication
- 4. Radiotherapy in the past two weeks

Date of first enrolment

01/02/2006

Date of final enrolment

31/12/2007

Locations

Countries of recruitment

Netherlands

Study participating centre

Erasmus Medical Center

Rotterdam

Netherlands
3008 AE

Sponsor information

Organisation

Erasmus Medical Center (Netherlands)

ROR

<https://ror.org/018906e22>

Funder(s)

Funder type

University/education

Funder Name

Erasmus Medical Center (Netherlands)

Alternative Name(s)

Erasmus Medical Center, Erasmus MC, Erasmus Universitair Medisch Centrum, Erasmus University Medical Center, Universitair Medisch Centrum Rotterdam, Erasmus Universitair Medisch Centrum Rotterdam, EMC

Funding Body Type

Government organisation

Funding Body Subtype

Universities (academic only)

Location

Netherlands

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Not provided at time of registration