Secondary prevention of problems in health status in patients with COPD by early detection, motivational intervention to engage in treatment by the patient and by individualized treatment

Gepubliceerd: 08-06-2009 Laatst bijgewerkt: 18-08-2022

In a sample of COPD patients with clinically relevant problems in four main domains of health status (physiological functioning, symptoms, functional impairment and quality of life) a motivational intervention conducted by a pulmonary nurse will...

Ethische beoordeling Positief advies **Status** Werving gestopt

Type aandoening -

Onderzoekstype Interventie onderzoek

Samenvatting

ID

NL-OMON24431

Bron

Nationaal Trial Register

Aandoening

Chronic Obstructive Pulmonary Disease (COPD), Health Status, Quality of Life, Intervention, Patient-tailored treatment, gezondheidstoestand, kwaliteit van leven, interventie, zorg op maat

Ondersteuning

Primaire sponsor: PICASSO-Partners in Care Solutions

Radboud University Nijmegen Medical Centre, department of Medical Psychology and department of Pulmonary Diseases

Sponsor / Initiator: Radboud University Nijmegen Medical Centre, Department of Medical

Psychology & Department of Pulmonary Diseases, Postbus 66, Nijmeegsebaan 31, 6560 AB Groesbeek

Overige ondersteuning: PICASSO-Partners in Care Solutions

Onderzoeksproduct en/of interventie

Uitkomstmaten

Primaire uitkomstmaten

- 1. Physiological Functioning: TLC%p predicted, RV% predicted, FEV1% predicted, BMI; < br>
- 2. Symptoms:

- B. Global Dyspnoea Activity and Global Dyspnoea Burden; <br
- C. Dyspnoea Emotions Questionnaire;

- D. Frustration and Anxiety;

- E. Checklist Individual Strength; <br
- F. Fatigue. < br>
- 3. Quality of Life:

- A. Beck Depression Inventory Primary Care; <br
- B. Satisfaction with Life Scale;

- 4. Functional Impairment:

- A. Quality of Life for Respiratory Illness Questionnaires;

- B. General Activities;

- C. Sickness Impact Profile;

- D. Home Management and Ambulation. <br
- 5. The number of additional treatments in both groups.

Toelichting onderzoek

Achtergrond van het onderzoek

Chronic Obstructive Pulmonary Disease (COPD) is a disease state characterized by airflow that is not fully reversible. Besides problems in phsyiological functioning, the patient also can experience symptoms, functional impairment and a diminished quality of life. Problems in the three latter domains of health status are hardly recognized in usual care, and remain untreated until escalated. This is mainly caused by two phenomena: doctor delay and patient delay. Doctor delay: the physician does not directly identify symptoms, functional impairment, and problems in quality of life. Patient delay: the patient does not report problems in these health status domains. What is necessary is a screening instrument that can be used in routine care and identifies patients with problems in the four domains of

health status. If clinically relevant problems exists, and additional treatment is recommended, an intervention by the pulmonary nurse is indicated. This intervention is directed at increasing awareness of existing problems and motivating the patient for additional treatment. By means of the screening and intervention, problems in health status are detected and treated early, before escalation. Treatment is patient-tailored, based on the existing problems in the four domains of health status, eventually leading to an improved health status. A randomized controlled trial is conducted to test this hypothesis, comparing (1) patients with no clinically relevant problems in health status (usual care as delivered by the outpatient clinic), (2) patients with clinically relevant problems in health status (usual care as delivered by the outpatient clinic), (3) patients with clinically relevant problems in health status receiving an intervention by a pulmonary nurse.

Doel van het onderzoek

In a sample of COPD patients with clinically relevant problems in four main domains of health status (physiological functioning, symptoms, functional impairment and quality of life) a motivational intervention conducted by a pulmonary nurse will lead to patient-tailored treatment and an improved health status.

Onderzoeksopzet

T1: group II/III;

T2 group I/II/III.

1. T0 (baseline);
2. T1 (after 6 months);
3. T2 (after 12 months).
Each timepoint measurement of primary and secondary outcomes.
T0: group I/II/III;

Onderzoeksproduct en/of interventie

Based on the indepent clinical interpretation of the PatientProfileChart by three professionals, patients are assigned to one of the following groups:

- 1. Patients with no clinically relevant problems in health status (group I);
- 2. Patients with clinically relevant problems in health status (group II/III).

Patients with clinically relevant problems in health status are randomized to a control group (group II: usual care as delivered by the outpatient clinic) and the experimental group (group III: intervention conducted by a pulmonary nurse, directed at increasing awareness of problems in health status, increasing motivation to engage in additional treatment, and improving health status).

Contactpersonen

Publiek

Postbus 66 Leonie Daudey Radboud University Nijmegen Medical Centre, Department of Medical Psychology and Department of Pulmonary Diseases

Groesbeek 6560 AB The Netherlands 0031-24-6859558

Wetenschappelijk

Postbus 66 Leonie Daudey Radboud University Nijmegen Medical Centre, Department of Medical Psychology and Department of Pulmonary Diseases

Groesbeek 6560 AB The Netherlands 0031-24-6859558

Deelname eisen

Belangrijkste voorwaarden om deel te mogen nemen (Inclusiecriteria)

Diagnosis COPD according to GOLD-criteria.

Belangrijkste redenen om niet deel te kunnen nemen

(Exclusiecriteria)

- 1. Not competent enough in understanding Dutch language;
- 2. Participation in pulmonary rehabilitation program within the previous six months.

Onderzoeksopzet

Opzet

Type: Interventie onderzoek

Onderzoeksmodel: Parallel

Toewijzing: Gerandomiseerd

Blindering: Open / niet geblindeerd

Controle: Actieve controle groep

Deelname

Nederland

Status: Werving gestopt

(Verwachte) startdatum: 01-09-2007

Aantal proefpersonen: 120

Type: Werkelijke startdatum

Ethische beoordeling

Positief advies

Datum: 08-06-2009

Soort: Eerste indiening

Registraties

Opgevolgd door onderstaande (mogelijk meer actuele) registratie

Geen registraties gevonden.

Andere (mogelijk minder actuele) registraties in dit register

Geen registraties gevonden.

In overige registers

Register ID

NTR-new NL1733 NTR-old NTR1844

CCMO NL15356.091.06

ISRCTN wordt niet meer aangevraagd.

Resultaten

Samenvatting resultaten

N/A