

# Is it possible to have a chronic obstructive pulmonary disease (COPD) specific breathlessness service in the Netherlands?

<b>Submission date</b> 01/11/2021	<b>Recruitment status</b> No longer recruiting	<input type="checkbox"/> Prospectively registered <input checked="" type="checkbox"/> Protocol
<b>Registration date</b> 06/11/2021	<b>Overall study status</b> Completed	<input type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
<b>Last Edited</b> 05/11/2021	<b>Condition category</b> Respiratory	<input type="checkbox"/> Individual participant data <input type="checkbox"/> Record updated in last year

## Plain English summary of protocol

### Background and study aims

Chronic obstructive pulmonary disease, or COPD, refers to a group of diseases that cause airflow blockage and breathing-related problems. Refractory (treatment resistant) breathlessness is a devastating symptom in COPD. Breathlessness services (BS), usually collaborations of palliative and respiratory teams, offer a multidisciplinary approach. In the Netherlands, few palliative care teams work in an outpatient setting, and BS do not exist.

The study involves training for people with COPD (chronic obstructive lung disease). The aim is to develop training for patients who are troubled by breathlessness. The training cannot take away the breathlessness but should help patients cope with the symptom.

### Who can participate?

Patients with COPD and refractory breathlessness.

### What does the study involve?

Participants attended at least two sessions with a pulmonologist and a respiratory nurse and one session with a physiotherapist who practiced the breathing techniques with them. The toolkit that was given to patients consisted of a booklet with breathing exercises and a hand-held fan. There was not a set number of sessions.

### What are the possible benefits and risks of participating?

Benefits: patients who participate might improve their coping skills regarding breathlessness. They also help us set up training that is appropriate for patients. There are no risks in participating; however, the intervention takes up some of the participants' time

### Where is the study run from?

Spaarne Gasthuis (Netherlands)

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

April 2019 to November 2020

Who is funding the study?  
Spaarne Gasthuis (Netherlands)

Who is the main contact?  
Kris Mooren, k.mooren@spaarnegasthuis.nl

## Contact information

**Type(s)**  
Scientific

**Contact name**  
Mrs Kris Mooren

**ORCID ID**  
<http://orcid.org/0000-0001-7815-0488>

**Contact details**  
Spaarne Gasthuis  
Boerhaavelaan 22  
Haarlem  
Netherlands  
2035 RC  
+31232245969  
k.mooren@spaarnegasthuis.nl

## Additional identifiers

**EudraCT/CTIS number**  
Nil known

**IRAS number**

**ClinicalTrials.gov number**  
Nil known

**Secondary identifying numbers**  
Nil known

## Study information

**Scientific Title**  
Testing the waters for a COPD-specific breathlessness service in the Netherlands: a feasibility study

**Study objectives**  
Our primary aim was to demonstrate the feasibility of setting up a BS specifically for COPD patients, with the view of undertaking a randomized controlled clinical trial to test the effectiveness of this BS on breathlessness mastery.

## **Ethics approval required**

Old ethics approval format

## **Ethics approval(s)**

Approved 09/08/2019, Medical Ethics Review Committee of the Amsterdam University Medical Centre (Medisch Ethische Toetsingscommissie VUmc, Van der Boechorststraat 7, kamer H-443, Postbus 7057

1007 MB Amsterdam, Netherlands; +31 20 444 5585; no email provided), ref: 2019.199

## **Study design**

Non-randomized single-center feasibility study

## **Primary study design**

Interventional

## **Secondary study design**

Non randomised study

## **Study setting(s)**

Hospital

## **Study type(s)**

Quality of life

## **Participant information sheet**

No participant information sheet available

## **Health condition(s) or problem(s) studied**

Improvement of quality of life in COPD patients with refractory dyspnea

## **Interventions**

For this study, the authors were trained by the Cambridge Breathlessness Intervention Service (CBIS). Their Breathing Thinking Functioning model (BTF) was translated into a Dutch toolkit (Ademen Denken Doen, ADD) with full consent of, and in collaboration with, the Cambridge team. The content of the Dutch intervention is similar to the original. However, the service in Cambridge is delivered by occupational therapists, physiotherapists and a consultant in palliative medicine, whereas the Dutch service is delivered by a pulmonologist, a respiratory nurse and a physiotherapist.

The model helps the patients to understand the vicious circles that influence breathlessness. A well-known example is the dyspnea-anxiety-dyspnea cycle: patients have catastrophic thoughts (I need oxygen, I am going to suffocate) (thinking domain), leading to shallow, exhausting 'dead space' breathing (breathing domain), leading to worsening of the symptom that can spiral into a panic attack. These episodes of breathlessness can lead to avoidance of exercise (functioning domain), in turn causing deconditioning that worsens the symptom.

ADD was set up as a brief intervention: patients had at least two sessions with a pulmonologist and a respiratory nurse, and one session with a physiotherapist who practiced the breathing techniques with them. The toolkit that was given to patients consisted of a booklet with breathing exercises and a hand-held fan. If necessary, extra sessions were scheduled. Since we had no experience with giving the intervention in this setting and specifically for COPD patients,

we decided not to fix the number of sessions. However, the average number of sessions for a future randomized multicenter controlled clinical trial shall be derived from this feasibility study.

**Intervention Type**

Behavioural

**Primary outcome measure**

Feasibility outcome: number of patients completing the intervention measured using patient records at the end of the study

**Secondary outcome measures**

1. Respiratory problems measured using a Dutch translation of the Chronic Respiratory Questionnaire (CRQ), subset mastery domain at baseline and final visit
2. Pulmonary function tests were taken from the patient's medical file at baseline
3. Patient and professional satisfaction measured using a postal survey with multiple choice questions after the final visit

**Overall study start date**

01/04/2019

**Completion date**

01/11/2020

**Eligibility****Key inclusion criteria**

Patients with COPD (diagnosed by a pulmonologist, post-bronchodilator FEV1/FVC below the lower limit of normal) were eligible for the study if they experienced refractory dyspnea (troubled by breathlessness despite optimization of COPD treatment) and were able to visit the outpatient clinic.

**Participant type(s)**

Patient

**Age group**

Adult

**Sex**

Both

**Target number of participants**

20

**Total final enrolment**

19

**Key exclusion criteria**

1. Terminal phase
2. Cognitive impairment
3. Unable to speak Dutch

**Date of first enrolment**

01/10/2019

**Date of final enrolment**

01/11/2020

## **Locations**

**Countries of recruitment**

Netherlands

**Study participating centre**

**Spaarne Gasthuis**

Boerhaavelaan 22

Haarlem

Netherlands

2035RC

## **Sponsor information**

**Organisation**

Spaarne Gasthuis

**Sponsor details**

Boerhaavelaan 22

Haarlem

Netherlands

2035 RC

+31 232240000

wetenschapsbureau@spaarnegasthuis.nl

**Sponsor type**

Hospital/treatment centre

**Website**

<https://spaarnegasthuis.nl>

## **Funder(s)**

**Funder type**

Hospital/treatment centre

**Funder Name**  
Spaarne Gasthuis

## Results and Publications

### Publication and dissemination plan

Planned publication in a peer-reviewed COPD journal.

### Intention to publish date

01/12/2021

### Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request (k.mooren@spaarnegasthuis.nl)

### IPD sharing plan summary

Available on request

### Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<a href="#">Protocol file</a>	version 4	01/08/2019	05/11/2021	No	No