The use of a portable positive airway pressure device to relieve shortness of breath after exercise

Submission date 22/06/2015	Recruitment status No longer recruiting	 Prospectively registered Protocol
Registration date 06/07/2015	Overall study status Completed	 Statistical analysis plan Results
Last Edited 05/01/2017	Condition category Respiratory	 Individual participant data Record updated in last year

Plain English summary of protocol

Background and study aims Chronic obstructive pulmonary disease (COPD) is the name used to refer to a number of progressive devastating and debilitating lung diseases, which includes chronic bronchitis, emphysema and chronic obstructive airways disease. People that have COPD typically feel breathless after physical activity, have a persistent cough with phlegm and suffer frequently from chest infections. Feeling short of breath can lead to inactivity and muscle deconditioning in patients with COPD. Positive airway pressure (PAP) can relieve shortness of breath and improve exercise tolerance (the amount of exercise that they are able to do). This study examines the effect of PAP delivered by a battery powered handheld device to COPD patients after exercise to relieve shortness of breath.

Who can participate?

Aged at least 18 with diagnosed COPD but able to do mild exercise.

What does the study involve?

Participants are randomly allocated into one of three groups. They are all then asked to do some mild exercise. Those in group 1 are then provided with the portable PAP device to use after the exercise to ease breathlessness. Those in group 2 are provided with what looks like a portable PAP device, but it is nonfunctional (a SHAM device) to use after the exercise. Those in group 3 are asked to perform lip breathing after exercise. The time it takes to return to the shortness of breath experienced prior to exercise for each participant is determined. We anticipate that the time after exercise to return to baseline shortness of breath using the hand held device is shorter than that while using a SHAM device and pursed-lip breathing (PLB) after exercise. As a secondary outcome, we will evaluate the distance walked during exercise.

What are the possible benefits and risks of participating?

Other than possibly gaining a better understanding of their ability to exercise and contributing to general knowledge of strategies to address shortness of breath, participants will not benefit directly from enrolling in the study. Risks of study participation include breathing and general discomfort while exercising. Lung damage from the portable positive airway pressure device is a remote possibility. Participants will be closely monitored during exercise and may request that the exercise activity be terminated. Patients with a history of lung damage from positive pressure devices are not eligible to participate.

Where is the study run from? Pittsburgh Pulmonary Associates Ltd (USA)

When is the study starting and how long is it expected to run for? April 2014 to October 2015

Who is funding the study? Philips Respironics (USA)

Who is the main contact? Mr William Hardy bill.hardy@philips.com

Contact information

Type(s) Public

Contact name Mr William Hardy

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Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers HRC-1510-ENGVB-SS

Study information

Scientific Title

Effect of intermittent positive air pressure in relieve of shortness of breath in COPD patients after exertion.

Study objectives

To determine whether intermittent positive airway pressure given after exercise to patients with COPD can shorten their shortness of breath recovery period

Ethics approval required Old ethics approval format

Ethics approval(s) Allendale Institutional Review Board (AIRB), 26/03/2015

Study design Prospective, interventional trial

Primary study design Interventional

Secondary study design Randomised controlled trial

Study setting(s) Other

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Study type(s)

Treatment

Participant information sheet

Not available in web format, please use contact details to request a participant information sheet

Health condition(s) or problem(s) studied

Chronic obstructive pulmonary disease

Interventions

Participants are asked to undergo exercise session and then use one of the following to alleviate shortness of breath:

- 1. Portable battery powered positive airway pressure (PAP) device
- 2. Portable sham positive airway pressure device

3. Pursed lip breathing

Intervention Type

Device

Primary outcome measure

Time to return to baseline shortness of breath rating after exercise

Secondary outcome measures

Distance walked during exercise

Overall study start date 06/04/2014

Completion date

30/10/2015

Eligibility

Key inclusion criteria

 Age ≥ 18
 Ability to provide consent
 COPD diagnosis with FEV1 < 55 and > 25 of predicted value
 Perceived Shortness of Breath via the Modified Medical Research Counsel Dyspnea questionnaire (rating of 2 or greater)
 Able to follow directions
 Able to tolerate mild physical activity

Participant type(s) Patient

Age group Adult

Lower age limit 18 Years

Sex Both

Target number of participants 30

Key exclusion criteria

 Subjects who are acutely ill, medically complicated or who are medically unstable as determined by the investigator
 Suffering from COPD Exacerbation at time of data collection

Date of first enrolment 06/04/2015

Date of final enrolment 30/10/2015

Locations

Countries of recruitment United States of America

Study participating centre

Sponsor information

Organisation Philips Respironics

Sponsor details 1740 Golden Mile Highway Monroeville United States of America 15146

Sponsor type Industry

Website www.respironics.com

ROR https://ror.org/03kw6wr76

Funder(s)

Funder type Industry

Funder Name Philips Respironics (USA)

Results and Publications

Publication and dissemination plan

Submitted late breaking abstract to the 2015 European Respiratory Society meeting. Upon completing recruitment of approximatlely thirty (30) participants, a peer reviewed publication is anticipated.

Intention to publish date

Individual participant data (IPD) sharing plan

IPD sharing plan summary Not provided at time of registration