

Can a new breathing device help people in hospital with severe flare ups of chronic lung disease clear mucus and breathe more easily?

Submission date 26/04/2026	Recruitment status Recruiting	<input type="checkbox"/> Prospectively registered <input checked="" type="checkbox"/> Protocol
Registration date 05/05/2026	Overall study status Ongoing	<input type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
Last Edited 30/04/2026	Condition category Respiratory	<input type="checkbox"/> Individual participant data <input checked="" type="checkbox"/> Record updated in last year

Plain English summary of protocol
Not provided at time of registration

Contact information

Type(s)

Principal investigator, Public, Scientific

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Additional identifiers

Study information

Scientific Title

Feasibility and benefits of an oscillatory negative-pressure device for secretion mobilization in patients hospitalized with acute COPD exacerbation - a randomized controlled trial

Study objectives

The aim of this study was to assess the feasibility and clinical effects of adding an oscillatory negative-pressure device to conventional respiratory physiotherapy in patients hospitalized for an acute exacerbation of COPD with secretion retention, compared with respiratory physiotherapy combined with a sham procedure.

Ethics approval required

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Ethics approval(s)

approved 14/05/2024, Comissão de ética da ULSAAVE (Hospital da Senhora da Oliveira, Guimarães. Comissão de Ética Rua dos Cutileiros, Creixomil, Guimarães, 4835-044, Portugal; +351 253540330; comissaoetica@ulsaave.min-saude.pt), ref: 45/2024

Primary study design

Interventional

Allocation

Randomized controlled trial

Masking

Blinded (masking used)

Control

Placebo

Assignment

Parallel

Purpose

Device feasibility

Study type(s)

Health condition(s) or problem(s) studied

Patients hospitalized with acute exacerbation of COPD

Interventions

Participants will be recruited by the attending physician in collaboration with a physiatrist, who confirmed the indication for respiratory rehabilitation. An initial interview will be conducted by one of the investigators to verify eligibility criteria and obtain written informed consent.

After consent is obtained, the baseline assessment (Assessment 0) will be performed and recorded in the Case Report Form. In this first assessment, demographic data, clinical history and clinical evaluation (Vital signs, respiratory symptoms, CAT, Leicester acute) will be registered.

Participants will then be randomized to either the intervention group or the control group using appropriate randomization software (Randomizer®) making a computer-generated allocation sequence.

The control group will receive conventional respiratory physiotherapy (breathing retraining, controlled inspiratory slow breathing, Slow open glottis expirations, huffing and cough) combined with a sham procedure using the Semiox® device, while the intervention group will receive similar respiratory physiotherapy combined with the Semiox® oscillatory negative-pressure device.

Interventions will be carried out once a day by the principal investigator, an experienced physiotherapist, with the agreement of the attending physician.

Intervention Type

Device

Phase

Phase II/III

Drug/device/biological/vaccine name(s)

Semiox® oscillatory negative-pressure device.

Primary outcome(s)

1. Sputum Weight (g) measured using a precision balance at 1 hour after treatment

Key secondary outcome(s)

1. COPD Impact (CAT) measured using CAT score at Every 3 days until study discharge

2. Cough-Related Quality of Life (LCQ-acute) measured using Leicester cough questionnaire acute score at Every 3 days until study discharge

3. Length of hospital stay (days) measured using Patient records at End of hospitalization

Completion date

31/12/2026

Eligibility

Key inclusion criteria

1. Patient with confirmed diagnosis of COPD
2. Hospitalized with respiratory onset
3. With an increase of his usual secretion production

Healthy volunteers allowed

No

Age group

Mixed

Lower age limit

18 years

Upper age limit

100 years

Sex

All

Total final enrolment

0

Key exclusion criteria

1. Hemodynamic instability requiring vasopressor support
2. Severe respiratory failure defined as $\text{PaO}_2/\text{FiO}_2 < 100$
3. Respiratory failure requiring a fraction of inspired oxygen (FiO_2) $> 60\%$
4. Altered level of consciousness or inability to cooperate with the intervention
5. Untreated or undrained pneumothorax
6. Massive hemoptysis
7. Unstable cardiovascular disease (e.g., acute coronary syndrome, uncontrolled arrhythmias)
8. Any clinical condition requiring strict bed rest or contraindicating participation in respiratory physiotherapy

Date of first enrolment

01/11/2024

Date of final enrolment

20/12/2026

Locations

Countries of recruitment

Portugal

Sponsor information

Organisation

Unidade Local de Saude do Alto Ave

ROR

<https://ror.org/049cs4442>

Funder(s)

Funder type

Funder Name

Investigator initiated and funded

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Not expected to be made available

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Protocol file			30/04/2026	No	No