

A 2x2 factorial randomised open label trial to determine the CLinical and cost-Effectiveness of hypertonic saline (HTS 6%) and carbocisteine for Airway cleaRance versus usual care over 52 weeks in bronchiectasis

Submission date 25/06/2018	Recruitment status No longer recruiting	☐ Prospectively registered ☒ Protocol
Registration date 06/07/2018	Overall study status Completed	☐ Statistical analysis plan ☒ Results
Last Edited 30/09/2025	Condition category Respiratory	☐ Individual participant data

Plain English summary of protocol

Background and study aims

Bronchiectasis is a condition where the lungs' airways become widened, leading to a build-up of mucus that can make them more vulnerable to infection. Patients with bronchiectasis suffer from a persistent cough, daily sputum (saliva and mucus) production and recurrent exacerbations, affecting quality of life. This study is concerned with sputum production and exploring which combinations of treatments (hypertonic saline (HTS) and carbocisteine) might help patients bring up their sputum more easily. HTS is salty water that is delivered into the airways via a nebuliser, which converts it into a mist. This treatment takes ten minutes. Carbocisteine is taken as a capsule. Research has shown that these treatments can make it easier for patients to cough up sputum, resulting in potentially fewer exacerbations and hospital admissions, which could improve quality of life for people with bronchiectasis. These treatments are currently used in clinical care but there is not enough evidence to recommend them as part of standard care. The aim of this study is to find out whether HTS and/or carbocisteine result in better outcomes than usual care in patients with bronchiectasis.

Who can participate?

Patients with bronchiectasis

What does the study involve?

Participants are randomly allocated to one of four combinations of treatments: HTS and standard care, carbocisteine and standard care, a combination of HTS and carbocisteine with standard care, and standard care alone. Participants have to attend six additional appointments as part of the study over a 2-year period with five in year one. At the start of the study, information is collected about participants' smoking status, bronchiectasis characteristics, medications, medical history, exacerbation history, antibiotic use, and any airway clearance techniques used. A physical exam, vital signs and urine pregnancy test are completed. A

questionnaire is completed with patients to explore any respiratory symptoms they may be experiencing since their last clinic visit. A lung function test is also performed and a range of health-related quality of life questionnaires are completed. At subsequent visits, lung function tests are repeated, exacerbation history is explored, vital signs are obtained and questionnaires are completed again. The nebulizer that patients use to deliver the hypertonic saline can record data about how the patient uses the nebuliser. This usage data is not reviewed with the patient until after they have completed the first year of the study. All patients are also be given a spirometer to perform lung function tests at home on a weekly basis. In addition, patients are asked to use their spirometers at the start of a suspected exacerbation. Patients are asked to bring their spirometer to each study visit. At the 52-week visit the patient is also asked to complete a questionnaire about using the spirometer and nebuliser. Follow-up at 104 weeks collects information about the patient's quality of life, any exacerbations they have had or any antibiotics they have taken for exacerbations. If the patient is already taking part in studies that are currently collecting information on bronchiectasis patients are asked for permission to use this information. Otherwise they are invited to the study site for an additional visit.

What are the potential benefits and risks of participating?

Carbocisteine may help reduce exacerbations and hospitalisations for people with bronchiectasis, improving their overall quality of life. All participants in the study receive a device to test their lung function at home and are also monitored regularly throughout the study. This may improve the overall care they receive through earlier recognition of exacerbations. Ultimately it is hoped that this research will help guide the treatment of bronchiectasis for people in the future and this may benefit patients in the long term outside of the study. Depending on the group allocation, a participant's treatment may be altered. A small number of people experience minor side effects from nebulising HTS such as wheezing, but the likelihood of this will be minimised by patients completing a drug response assessment on entry into the study. A small number of patients experience some minimal side effects (stomach discomfort, vomiting, skin rashes and allergies) associated with carbocisteine. Lung function procedures may cause the participant to cough, experience shortness of breath, or feel lightheaded, but there is no pain expected with these tests. Treatment will be provided if this occurs. This study will take place outside of the normal clinical management of participants. Participants will be required to attend up to six additional visits lasting about 2 hours. Appointments will be scheduled at a time convenient to participants, where possible in an attempt to minimise this burden.

Where is the study run from?

Belfast City Hospital (UK) (lead centre)

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

July 2017 to September 2024

Who is funding the study?

National Institute for Health and Care Research (NIHR) (UK)

Who is the main contact?

Prof. Judy Bradley

judy.bradley@qub.ac.uk

Contact information

Type(s)

Scientific

Contact name

Prof Judy Bradley

ORCID ID

<https://orcid.org/0000-0002-7423-135X>

Contact details

School of Medicine, Dentistry and Biomedical Sciences

NI Clinical Research Facility

U Floor

Belfast City Hospital

Lisburn Road

Belfast

United Kingdom

BT9 7AB

+44 (0)28 9504 0342

judy.bradley@qub.ac.uk

Type(s)

Public

Contact name

Mr Andrew Jackson

Contact details

Northern Ireland Clinical Trials Unit

1st Floor Elliot Dynes

The Royal Group of Hospitals

Grosvenor Road

Belfast

United Kingdom

BT12 6BA

-

CLEAR@nictu.hscni.net

Type(s)

Public

Contact name

Mr Andrew Jackson

Contact details

NICTU

1st Floor Elliot Dynes Building

Royal Hospitals

Grosvenor Road

Belfast

United Kingdom

BT12 6BA

+44 (0) 28961 51447
AndrewX.Jackson@nctu.hscni.net

Additional identifiers

Clinical Trials Information System (CTIS)
2017-000664-14

Integrated Research Application System (IRAS)
214254

ClinicalTrials.gov (NCT)
NCT04140214

Protocol serial number
CPMS 37574; 16178SE-AS

Study information

Scientific Title

A 2x2 factorial randomised open label trial to determine the CLinical and cost-Effectiveness of hypertonic saline (HTS 6%) and carbocisteine for Airway cleaRance versus usual care over 52 weeks in bronchiectasis

Acronym
CLEAR

Study objectives

HTS (6%) and/or the oral mucolytic carbocisteine will result in better outcomes than usual care over 52 weeks in patients with bronchiectasis.

Ethics approval required
Old ethics approval format

Ethics approval(s)
North East – Tyne & Wear South Research Ethics Committee, 20/12/2017, ref: 17/NE/0339

Study design
Randomized; Interventional; Design type: Treatment, Drug

Primary study design
Interventional

Study type(s)
Treatment

Health condition(s) or problem(s) studied
Bronchiectasis

Interventions

Current interventions as of 06/08/2025:

Treatment allocation will be assigned using an automated randomisation process. Eligible participants will be allocated to one of the four treatment groups (three intervention groups or one standard care group) in a 1:1:1:1 ratio using a central randomisation system. Randomisation will be stratified by site, to minimise baseline imbalances in antibiotic use due to exacerbations in the last year (2-3 times, >3 times) and based on current use of macrolides (yes, no).

Intervention 1: Standard care and twice daily nebulised HTS (6%) over 52 weeks

Intervention 2: Standard care and carbocisteine (750 mg three times per day until visit 3 reducing to 750 mg twice per day) over 52 weeks

Intervention 3: Standard care and a combination of twice daily nebulised HTS (6%) and 750 mg of carbocisteine three times per day until visit 3* reducing to 750 mg twice per day) over 52 weeks

Intervention 4: Control: standard care over 52 weeks

The study treatment period is 52 weeks.

Previous interventions:

Treatment allocation will be assigned using an automated randomisation process. Eligible participants will be allocated to one of the four treatment groups (three intervention groups or one standard care group) in a 1:1:1:1 ratio using a central randomisation system. Randomisation will be stratified by site, to minimise baseline imbalances in antibiotic use due to exacerbations in the last year (2-3 times, >3 times) and based on current use of macrolides (yes, no).

Intervention 1: Standard care and twice daily nebulised HTS (6%) over 52 weeks

Intervention 2: Standard care and carbocisteine (750 mg three times per day until visit 3 reducing to 750 mg twice per day) over 52 weeks

Intervention 3: Standard care and a combination of twice daily nebulised HTS (6%) and 750 mg of carbocisteine three times per day until visit 3* reducing to 750 mg twice per day) over 52 weeks

Intervention 4: Control: standard care over 52 weeks

The study treatment period is 52 weeks, after which follow-up will take place at week 104.

Intervention Type

Drug

Phase

Not Applicable

Drug/device/biological/vaccine name(s)

Hypertonic saline, carbocisteine

Primary outcome(s)

Mean number of exacerbations over 52 weeks; exacerbations diagnosed as they occur via interview with patients and completion of the Respiratory and Systemic Symptoms questionnaire (RSSQ)

Key secondary outcome(s)

1. Disease-specific health-related quality of life (HRQoL) at 52 weeks measured using the respiratory symptoms domain of quality of life with bronchiectasis (QoL B) questionnaire completed at baseline, 2 weeks, 8 weeks, 26 weeks and 52 weeks

2. Time to next exacerbation post-randomisation measured in days, with exacerbations

diagnosed as they occur via interview with patients and completion of the RSSQ

3. Number of days of antibiotics related to exacerbations over 52 weeks assessed via interview with patients at 2, 8, 26 and 52 weeks

4. Generic health-related quality of life (HRQoL) measured using the EQ-5D-5L questionnaire at baseline, 2 weeks, 8 weeks, 26 weeks and 52 weeks

5. Health Service use over 52 weeks assessed via completion of a questionnaire completed at baseline, 2 weeks, 8 weeks, 26 weeks and 52 weeks

6. Quality Adjusted Life Years (QALYs) over 52 weeks calculated by assessment of generic HRQoL measured using the EQ-5D-5L questionnaire completed at baseline, 2 weeks, 8 weeks, 26 weeks and 52 weeks

7. Health impairment measured using the St George's Respiratory Questionnaire (SGRQ) at baseline, 2 weeks, 8 weeks, 26 weeks and 52 weeks

8. Patient preferences for treatment measured using a Treatment Satisfaction Questionnaire at 2, 8, 26, and 52 weeks

9. Adverse events over 52 weeks recorded as they occur by the PI or designee via interview with patients

10. Lung function over 52 weeks measured by spirometry testing at baseline, 2 weeks, 8 weeks, 26 weeks and 52 weeks

11. Adherence to HTS and carbocysteine over 52 weeks assessed via interview with patients at 2, 8, 26, and 52 weeks

Completion date

30/09/2024

Eligibility

Key inclusion criteria

Current inclusion criteria as of 06/10/2022:

1. Diagnosis of BE on high resolution computed tomography (HRCT)/computed tomography (CT) scans

2. BE must be the primary respiratory diagnosis

3. One or more pulmonary exacerbations in the last year requiring antibiotics*

4. Production of daily sputum

5. Stable for 14 or more days before the first study visit with no changes to treatment

6. Willing to continue any other existing chronic medication throughout the study

7. Female subjects must be either surgically sterile, postmenopausal or agree to use effective contraception during the treatment period of the trial

*This can include patient-reported exacerbations

Previous inclusion criteria:

1. Diagnosis of BE on high resolution computed tomography (HRCT)/computed tomography (CT) scans

2. BE must be the primary respiratory diagnosis

3. Two or more pulmonary exacerbations in the last year requiring antibiotics*

4. Production of daily sputum

5. Stable for 14 or more days before the first study visit with no changes to treatment

6. Willing to continue any other existing chronic medication throughout the study

7. Female subjects must be either surgically sterile, postmenopausal or agree to use effective contraception during the treatment period of the trial

*This can include patient-reported exacerbations

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

18 years

Sex

All

Total final enrolment

288

Key exclusion criteria

1. Age <18 years old
2. Patients with CF
3. Patients with COPD as a primary respiratory diagnosis
4. Current smokers, female ex-smokers with greater than 20 pack years and male ex-smokers with greater than 25 pack years
5. FEV1<30%
6. If being treated with long term macrolides, on treatment for less than 1 month before joining study
7. Patients on regular isotonic saline
8. Treatment with HTS, carbocysteine or any mucolytics within the past 30 days
9. Known intolerance or contraindication to HTS or carbocysteine.
10. Hypersensitivity to any of the active ingredients or the excipients of carbocysteine
11. Active peptic ulceration
12. Any heredity galactose intolerance, the Lapp-Lactase deficiency or glucose-galactose malabsorption.
13. Patients unable to swallow oral capsules.
14. Women who are pregnant or lactating
15. Participation in another Clinical Trial of an Investigational Product within 30 days

Date of first enrolment

27/06/2018

Date of final enrolment

30/09/2023

Locations**Countries of recruitment**

United Kingdom

England

Northern Ireland

Scotland

Wales

Study participating centre

Altnagelvin Area Hospital

Glenshane Road

Londonderry

United Kingdom

BT47 6SB

Study participating centre

Royal Free Hospital

Pond St

Hampstead

London

United Kingdom

NW3 2QG

Study participating centre

Royal Infirmary of Edinburgh

51 Little France Crescent

Edinburgh

United Kingdom

EH16 4SA

Study participating centre

Freeman Hospital

Freeman Road

High Heaton

Newcastle upon Tyne

United Kingdom

NE7 7DN

Study participating centre

Royal Brompton Hospital

Sydney St

Chelsea

London

United Kingdom
SW3 6NP

Study participating centre

Belfast City Hospital

Lisburn Road
Belfast
United Kingdom
BT9 7AB

Study participating centre

Ninewells Hospital and Medical School

Ninewells Avenue
Dundee
United Kingdom
DD1 9SY

Study participating centre

Princess Alexandra Hospital

Hamstel Road
Harlow
United Kingdom
CM20 1QX

Study participating centre

The Ulster Hospital

Upper Newtownards Road
Dundonald
United Kingdom
BT16 1RH

Study participating centre

Southampton General Hospital

Tremona Road
Southampton
United Kingdom
SO16 6YD

Study participating centre

Craigavon Area Hospital

Lurgan Rd
Craigavon
United Kingdom
BT63 5QQ

Study participating centre**University Hospitals of Morecambe Bay NHS Foundation Trust**

Westmorland General Hospital
Burton Road
Kendal
United Kingdom
LA9 7RG

Study participating centre**University Hospital Birmingham**

Queen Elizabeth Hospital
Edgbaston
Birmingham
United Kingdom
B15 2TH

Study participating centre**Churchill Hospital**

Churchill Hospital
Old Road
Headington
Oxford
United Kingdom
OX3 7LE

Study participating centre**Royal Gwent Hospital**

Cardiff Road
Newport
United Kingdom
NP20 2UB

Study participating centre**Cardiff & Vale**

Heath Park Way

Cardiff
United Kingdom
CF14 4XW

Study participating centre
Stoke Mandeville Hospital
Mandeville Road
Aylesbury
United Kingdom
HP21 8AL

Study participating centre
Blackpool Victoria Hospital
Whinney Heys Road
Blackpool
United Kingdom
FY3 8NR

Study participating centre
North Tyneside General Hospital
Rake Lane
North Shields
United Kingdom
NE29 8NH

Study participating centre
Bradford Teaching Hospitals NHS Foundation Trust
Bradford Royal Infirmary
Duckworth Lane
Bradford
United Kingdom
BD9 6RJ

Study participating centre
Milton Keynes University Hospital
Milton Keynes Hospital
Standing Way
Eaglestone
Milton Keynes
United Kingdom
MK6 5LD

Study participating centre
Sandwell General Hospital
Lyndon
West Bromwich
United Kingdom
B71 4HJ

Study participating centre
South West Acute Hospital
124 Irvinestown Rd
Enniskillen
United Kingdom
BT74 6DN

Sponsor information

Organisation
Belfast Health & Social Care Trust

ROR
<https://ror.org/02tdmfk69>

Funder(s)

Funder type
Government

Funder Name
NIHR Evaluation, Trials and Studies Co-ordinating Centre (NETSCC); Grant Codes: 15/100/01

Results and Publications

Individual participant data (IPD) sharing plan
The datasets generated and/or analysed during the current study during this study will be included in the subsequent results publication.

IPD sharing plan summary

Other

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	protocol	28/09/2025	30/09/2025	Yes	No
Protocol article		19/12/2019	23/12/2019	Yes	No
HRA research summary	Participant information sheet		28/06/2023	No	No
Participant information sheet		11/11/2025	11/11/2025	No	Yes
Study website	Study website	11/11/2025	11/11/2025	No	Yes