# Multiple combination bactericidal antibiotics testing for acute exacerbations of cystic fibrosis associated with multi-resistant Burkholderia cepacia and Pseudomonas aeruginosa infection

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# Plain English summary of protocol

Not provided at time of registration

# **Contact information**

# Type(s)

Scientific

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# Additional identifiers

Protocol serial number MCT-44147

# Study information

#### Scientific Title

Multiple combination bactericidal antibiotics testing for acute exacerbations of cystic fibrosis associated with multi-resistant Burkholderia cepacia and Pseudomonas aeruginosa infection: a randomised controlled trial

#### **Study objectives**

The objective of this clinical trial is to prospectively assess whether the use of combination antibiotic therapy, directed by results from multiple combination, bactericidal antibiotic testing (MCBT), improves bacteriologic and clinical outcomes in patients with acute pulmonary exacerbations of cystic fibrosis who are infected with multiple resistant bacteria.

#### Ethics approval required

Old ethics approval format

#### Ethics approval(s)

Approval gained from the Ottawa Hospital Research Ethics Board in Spring 2000

### Study design

Randomised controlled trial

#### Primary study design

Interventional

#### Study type(s)

Treatment

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

Pulmonary exacerbation in adult patients with cystic fibrosis (CF)

#### **Interventions**

Patients randomised to the control group will receive a 14 days course of any two intravenous antibiotics ± one inhaled antibiotic (tobramycin/TOBI) chosen by their physicians based on usual culture and sensitivity testing.

Patients randomised to MCBT-directed therapy group will receive a 14 days course of any two intravenous antibiotics ± one inhaled antibiotic (tobramycin/TOBI) chosen based on MCBT culture and sensitivity testing.

### Intervention Type

Drug

#### Phase

Not Applicable

# Drug/device/biological/vaccine name(s)

**Antibiotics** 

#### Primary outcome(s)

The time (days) from randomisation until the patient's next pulmonary exacerbation.

#### Key secondary outcome(s))

- 1. Mean changes in sputum bacterial densities for day zero to day 14
- 2. Changes in pulmonary function, pre-bronchodilator forced expiratory volume in one second (FEV1), and forced vital capacity (FVC)
- 3. Changes in oxygenation from day zero to day 14
- 4. The proportion of antibiotic treatment failures in both treatment groups within 14 days
- 5. Changes in subjective dyspnoea score from day zero to day 14
- 6. Length of hospital stay, for patients admitted to hospital
- 7. Adverse effects related to antibiotic therapy

#### Completion date

15/02/2005

# **Eligibility**

#### Key inclusion criteria

- 1. Age greater than or equal to 12, either sex
- 2. A confirmed diagnosis of cystic fibrosis (a sweat chloride value higher than 60 mmol/litre or two disease-causing mutations)
- 3. Chronically colonised with multi-resistant P. aeruginosa, or S. maltophilia, or A. xylosidans (at least two sputum cultures within the last 12 months which have grown these multi-resistant bacteria, one of which must have been obtained within six months of randomisation)
- 4. Patients must be known to be chronically colonised with Burkholderia cepacia bacteria (at least two sputum cultures within the last 12 months which have grown Burkholderia cepacia, one of which must have been obtained within six months of randomisation)
- 5. Patients must be able to spontaneously produce sputum for culturing

#### Participant type(s)

Patient

## Healthy volunteers allowed

No

## Age group

Adult

#### Sex

All

#### Key exclusion criteria

- 1. Unable to give informed consent
- 2. Previous lung transplant recipients
- 3. Patients with severe pulmonary exacerbations who require admission to an Intensive Care Unit (ICU) and/or mechanical ventilatory support
- 4. Patients who are already receiving continuous home intravenous antibiotic therapy
- 5. Pregnant patients

#### Date of first enrolment

03/08/2000

### Date of final enrolment

15/02/2005

# Locations

#### Countries of recruitment

Canada

Study participating centre The Ottawa Hospital Ottawa, Ontario Canada K1H 8L6

# Sponsor information

### Organisation

Ottawa Hospital Research Institute (Canada)

#### **ROR**

https://ror.org/03c62dg59

# Funder(s)

## Funder type

Research organisation

#### **Funder Name**

Canadian Institutes of Health Research (CIHR) (Canada) - http://www.cihr-irsc.gc.ca (ref: MCT-44147)

#### Funder Name

Other funders:

#### **Funder Name**

1. Canadian Cystic Fibrosis Foundation (Canada)

#### Funder Name

2. Astra Zeneca Canada Inc. (Canada)

# **Results and Publications**

Individual participant data (IPD) sharing plan

# IPD sharing plan summary

Not provided at time of registration

## **Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	results	01/08/2005		Yes	No