

Participant Identification Number:

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Title of Trial: Vital BE Trial; A Study of the Safety, Tolerability and Efficacy of 2 doses of Cayston (Aztreonam Lysine) compared to placebo in patients with bronchiectasis

IRAS ref number: 252929

Name of the Researcher: Dr Tom Bewick

Sponsors: University of Dundee and NHS Tayside

Please initial box

1. I confirm that I have read the information sheet/leaflet dated.....
(version.....) for the above trial. I have had the opportunity to consider
the information, ask questions and have had these answered
satisfactorily.

2. I understand that my participation is voluntary and that I am free to
withdraw at any time without giving any reason, without my medical care
or legal rights being affected.

3. I understand that personal data about me and research data collected
during the trial will be stored by the University of Dundee.

4. I understand that relevant sections of my medical records and data
collected during the trial, may be looked at by the Researcher and/or
research team, the Sponsors or regulatory authorities where it is relevant
to my taking part in this research. I give permission for the Researcher
and/or research team, the Sponsors and regulators to have access to my
records and data.

5. I understand that my research data collected by the Researcher and/or
research team in this trial may be used to support other research in the
future, and may be shared anonymously with other researchers or
collaborators, including commercial organisations.

INFORMED CONSENT FORM

6. I understand that my tissue (blood, nasal swab and sputum sample) collected by the Researcher and/or research team in this trial will be stored and that any excess may be shared anonymously with other researchers collaborating with the Sponsors to support future research - including commercial organisations. I hereby gift such tissue to the Sponsors.
7. I agree to my General Practitioner being informed of my participation in the trial and of any clinical findings made during the trial.
8. I agree to take part in the above trial.
9. *OPTIONAL QUESTION:* I agree to be contacted by the Researcher and/or research team in the event that I may be suitable for further research projects in the future. YES NO

Name of Participant
(CAPITAL LETTERS)

Date

Signature

Name of Person
taking consent (CAPITAL LETTERS)

Date

Signature

1 for participant, 1 to be kept in medical records and 1 for trial file.