

Participant name:

| | Hospital ID/CHI: | |
|---|--|---|
| Value of inhaled treatment with aztreonam lysine in bronchiectasis | Date of Birth: | |
| Value of inhaled treatment wit Sponsor Chief Investigator IRAS number | h Aztreonam lysine in bronchi University of Dundee-NHS ⁻ Professor James Chalmers 252929 | |
| Principal Investigator Contact number Contact email | | |
| Visit number Date of visit: | | |
| The above participant has agreed phone call. | l to take part in the VitalBE clinic | al trial and has attended for follow-up |
| | | |

Please tick to indicate the following has been completed:

Confirmed participant's identity Participant has verbally given their consent to continue in the trial Visit has been carried out as per protocol

Vital signs

| Please enter the results of the following assessments: | |
|--|------------------------|
| | Pulse bpm |
| Oxygen saturation (room air) % | Tympanic temperature C |
| Pregnancy test Is the participant a woman of child bearing potential? yes If female but not of child bearing potential how has this beer Post-menopausal Date of last period permanent sterilisation Date of last period If female and of childbearing potential: Has the participant agreed to either abstain from se form of a medically approved birth control method? result of pregnancy test: negative positive Spirometry What method of bronchodilation was used? nebulised salbutamol Dose mg | n confirmed? |
| Research blood sample obtained? | Yes No |
| Spontaneous sputum sample obtained? | Yes No |
| Viral Nasal Swab obtained? | Yes No |
| Sputum sample for Culture & Sensitivity? | Yes No |
| Is St George's Respiratory Questionnaire being | Yes No |
| completed? | |
| Is Quality of Life Bronchiectasis Questionnaire being completed? | Yes No |
| Is Bronchiectasis Health Questionnaire being completed? | Yes No |



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| name: | |
|-------|--|
| CHI: | |
| h: | |
| | |

The following must have source data documented in the medical notes. If not documented elsewhere these should be written in the notes.

Changes to concomitant medications/respiratory medications since last visit

Any adverse events since last visit

Any pulmonary exacerbations since last visit

Any other notable findings and actions taken

If the participant was withdrawn from the trial at this visit state reason: