

Participant name:	
Hospital ID/CHI:	
Date of Birth:	

aztreonam lysine ir		Date of Birth:				
Value of inhaled Sponsor Chief Investigate	d treat	ment with Aztreonam lysine in bro University of Dundee-I Professor James Chal	NHS Tays		ΓAL- BE	.
IRAS number		252929				
Principal Investion Contact number Contact email	gator					
Visit number Date of visit:	Visit	5				
The above partici phone call.	pant h	as agreed to take part in the VitalBE	clinical tr	ial and h	as atte	nded for follow-up
Confirm Participa Visit has	ed part ant has been	the following has been completed: ticipant's identity s verbally given their consent to conti carried out as per protocol pant bring a previously completed Qo			e?	
Vital signs Please enter the Blood pressure Oxygen saturation		s of the following assessments: mmHg m air) %	Pulse Tympani	c tempe	rature	bpm C
If female but not Post-me permane If female and of Has the form of a	of chil nopau ent ste childbe partici a medi	rilisation	n confirm] se a	yes no
nebulise inhaled :	d salb salbuta		mber of բ	ouffs	\neg	
Research blood	sampl	e obtained?	Yes	No		
Chantanas::-		nample obtained?	Vac	Na		
		sample obtained? Iture & Sensitivity?	Yes Yes	No No	-	
		tory Questionnaire being	Yes	No	1	
completed?			1			
Is Quality of Life completed?	Bronc	hiectasis Questionnaire being	Yes	No		
- John Protog :				<u> </u>	ĺ	



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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

Trial	med	licat	tion
HILL	11164	IICU	LIVII

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	t 🔿	confirm	١.

Signature: Name: Job title: Date:

Participant has been given trial medication, nebuliser and handset	
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If the participant was withdrawn from the trial at this visit state	
reason:	