

Joint Arrangements for Research

Serious Adverse Event report - NNUH sponsored CTIMP studies

What are you reporting?				
SAE / SAR 🗌			SUSAR* 🗆	
*Note: If you are reporting a SUSAR the randomisation code for that participant will have to be unblinded				
Report Type:		Initial 🔲 Follow-up Report 🗌 Follow-up Report #		
Study information	·			
Study Title: (short)				
Sponsor:		hief Investigator Name: mail Address:		
Eudract Number: (for CTIMPs only)		R&D Reference Number / IRAS Number:		
Protocol title and version number: Version and date of approved Reference Safety Information (RSI) in use at onset of SAE:				
Site Number: (for multi-site studies only)	Site Name:			
Principal Investigator	Name: Email address:			
Date of site becoming aware of the event (dd/mm/yy):				
Participant information				
Participant initials:			Participant Gender:	
Participant Randomisation No:				





Evaluation of Event Please refer to definitions and Appendix 1 &2 in SOP 205 for guide on understanding of					
seriousness, relatedness, expectedness. Event/Reaction: (keywords; e.g. body site, symptoms, severity, treatment)					
	nas, e.g. body sile	e, symptoms, seventy, treatment)			
Date of onset:		Date person completing form became aware of			
(dd/mm/yy)		event: (dd/mm/yy)			
		al abnormality/birth defect			
	□ Resulted	•			
	Life threa				
Criteria for		t hospitalisation/prolongation of hospitalisation			
definition as SAE		· · · · ·			
		Persistent or significant disability			
		vise considered medically significant by the investigator			
Describe event: (A sum		e than one criterion, choose the more/most significant one. I symptoms, diagnosis, treatment of event, concurrent treatment, other relevant			
medical history, including re-ch	allenge details if a	applicable. Please include the point in the study at which the event occurred.)			
		Definitely			
In the investigators o	ninion was				
In the investigators opinion was the event related to a research procedure?					
		□ Not related			
Please specify which	procedure				
if applicable					
		Definitely			
In the investigators o the event related to the					
Investigational Medic	-				
Product?					
		□ Not related			
Action taken with stu	dy				
drug/participant:		Dose temporarily reduced			
		Dose reduced			

R&D SOP Number: SOP 205 Appendix 1 Author/s title: Research Governance Coordinator Date approved: 17/07/2023 Review date: 17/07/2026







				Disc	ontinue	ntinued temporarily			
					ontinue	ntinued			
					articipant withdrawn from trial				
· · · · · · · · · · ·				□ Yes					
If related to IMP was this reaction unexpected (Suspected Unexpected				□ No					
Serious Adverse Reaction – SUSAR)?			□ Not applicable						
				□ Yes					
Did event/reaction abate after			□ No						
stopping drug?				□ Not applicable					
Did event/reaction				□ No					
after reintroduction	on of dru	g?	□ Not applicable						
IMP & concomitant medication information									
Drug details (Daily dose and generic name)	Administration St		Sta	erapy art Date d/mm/yy)	Therapy End Date (dd/mm/	e	Date of last administered Dose prior to SAE onset (dd/mm/yy)	Indications for Use	
Have urgent safe measures been implemented?	ty	If yes, please detail below:							
🗆 No									
□ Not applicable									
Outcome of event									
SAF? resol			Date ev resolve	ed:	Date nationt died				







			NHS FOUND
Recovered with sequalae			
Resulted in death			
Unknown			
	🗆 Coi	oner's inquest	
Cause of death obtained from:	Death certificate		
	🗆 Wo	rking diagnosis	

Contact and signatures
Please supply contact details where further information may be obtained:
Person to contact:
Phone number:
Email address:

Signature (person completing report)	Print name	Date		
PI Signature (if multicentre trial)	Print name	Date		
CI Signature (if not completing report)	Print name	Date		
Please send the completed form to rdsae@nnuh.nhs.uk .				
For F	R&D Office use only			





For SUSAR only:	Date reported to the REC:
	Date reported to MHRA:





For NNUH sponsored trial REVIEW BY NNUH MEDICAL ADVISOR	Medical Advisor comments:
SIGNED:	
Name (print):	
Date:	