For office use

QC: No

v1.6 06 December 2021

Short project title*:	RECAST-3									
IRAS project ID* (or REC reference if no IRAS project ID is available):	282606									
Sponsor amendment reference number*:	MA_05_24									
Sponsor amendment date* (enter as DD/MM/YY):	09 July 2024									
Briefly summarise in lay language the main changes proposed in this amendment. Explain the purpose of the changes and their significance for the study. If the amendment significantly alters the research design or methodology, or could otherwise affect the scientific value of the study, supporting scientific information should be given (or enclosed separately). Indicate whether or not additional scientific critique has been obtained (note: this field will adapt to the amount of text entered)*:	Minor change to stu Information sheets: intervention doses (o space for local PALS	wording amended to correction to align w	ith existing approve	ed protocol), and						
				Specific stu	ıdy					
Project type (select):		Research tissue ban								
			Research database							
Has the study been reviewed by a UKECA-recognised Res Committee (REC) prior to this amendment?:	search Ethics	Y	es		No					
What type of UKECA-recognised Research Ethics Commit	ttoo (PEC) raviow	NHS/HSC REC								
is applicable? (select):	itee (NEC) leview			Ministry of D	efence (MoDRE)					
Is all or part of this amendment being resubmitted to the Ro Committee (REC) as a <b>modified amendment</b> (i.e. a subst amendment previously given an unfavourable opinion)?		Y	es		No					
Where is the NHS/HSC Research Ethics Committee (REC	that reviewed	England	Wales	Scotland	Northern Irelar					
the study based?:	, macronous	Yes	No	No	No					
Was the study a clinical trial of an investigational medicinal OR does the amendment make it one?:	product (CTIMP)	Y	es	No						
Was the study a clinical investigation or other study of a medoes the amendment make it one?:	edical device OR	Y	es	No						
Does this clinical investigation or other study of a med require a Notice of No Objection from MHRA Devices			Yes	No No						
Did the study involve the administration of radioactive subsrequiring ARSAC review, OR does the amendment introdu		Y	es							
Did the study involve the use of research exposures to ioni (not involving the administration of radioactive substances) amendment introduce this?:		Y	es							
Did the study have Radiation Assurance OR is Radiation being sought for the first time because of this amendme			Yes		No					
Did the study involve adults lacking capacity OR does the a introduce this?:	amendment	Y	es		No					
Did the study involve access to confidential patient informa direct care team without consent OR does the amendment		Y	es	No						
Did the study involve prisoners or young offenders who are supervised by the probation service OR does the amendmenthis?:		Y	es	No						
Did the study involve children OR does the amendment int	roduce this?:	Y	es		No					
Did the study involve NHS/HSC organisations prior to this a		Y	es		No					
Did the study involve non-NHS/HSC organisations OR doe amendment introduce them?:	es the	Y	es	No						
		England	Wales	Scotland	Northern Irelar					
Lead nation for the study:		Yes	No	No	No					
Which nations had participating NHS/HSC organisations pramendment?		Yes	Yes	Yes	Yes					
Which nations will have participating NHS/HSC organisation	ns after this		Yes	Yes	Yes					

Section 2: Summary of change(s)

Please note: Each change being made as part of the amendment must be entered separately. For example, if an amendment to a clinical trial of an investigational medicinal product (CTIMP) involves an update to the Investigator's Brochure (IB), affecting the Reference Safety Information (RSI) and so the information documents to be given to participants, these should be entered into the Amendment Tool as three separate changes. A list of all possible changes is available on the "Glossary of Amendment Options" tab. To add another change, click the "Add another change" box.

Change 1									
Area of change (select)*:	Study Documents								
Specific change (select - only available when area of change is selected first)*:	questionnaires, letters	to study documents (e.g. information sheets, consent forms, rs) that can be implemented within existing resource in place at ations - Please specify in the free text below							
Further information In particular, please describe why this change can be implemented within the existing resource in place at the participating organisations (free text - note that this field will adapt to the amount of text entered)*	Information sheets: wording amended to indicate that there will be a gap of >4 hours between intervention doses (correction to align with existing approved protocol), and updated to provid space for local PALS and research team contact details to be documented.								
Applicability:	England	Wales	Scotland	Northern Ireland					
Where are the participating NHS/HSC organisations locate by this change?*:	No	No	Yes	No					
Will all participating NHS/HSC organisations be affected by some? (please note that this answer may affect the categoriange):	A	di .	Some						
				Add anot	her change				

### Section 3: Declaration(s) and lock for submission

### Declaration by the Sponsor or authorised delegate

- I confirm that the Sponsor takes responsibility for the completed amendment tool
- · I confirm that I have been formally authorised by the Sponsor to complete the amendment tool on their behalf

Name [first name and surname]*:	Alison Thorpe
Email address*:	sponsor@nottingham.ac.uk

#### Lock for submission

Please note: This button will only become available when all mandatory (\*) fields have been completed. When the button is available, clicking it will generate a locked PDF copy of the completed amendment tool which must be included in the amendment submission. Please ensure that the amendment tool is completed correctly before locking it for submission.

## Lock for submission

After locking the tool, proceed to submit the amendment online. The "Submission Guidance" tab provides further information about the next steps for the amendment.

# Section 4: Review bodies for the amendment

Please note: This section is for information only. Details in this section will complete automatically based on the options selected in Sections 1 and 2.

		Review bodies																	
	UK wide:				Eng	England and Wales:			Scotland:				Northern Ireland:						
Change 1:	REC	Competent Authority MHRA - Medicines	Competent Authority MHRA - Devices	ARSAC	Radiation Assurance	(3) UKSW Governance	REC (MCA)	CAG	HMPPS	HRA and HCRW Approval	REC (AWIA)	PBPP	SPS (RAEC)	(3) National coordinating function	HSC REC	HSC Data Guardians	Prisons	National coordinating function	Category:
						(1)								(1)					
Overall reviews for the amendme	nt:																		
Full review:						Ν								N					
Notification only:						Υ								Υ					
Overall amendment type:	Non-substantial, no study-wide review required																		
Overall Category:	С																		