Loughborough University The Centre for Biological Engineering

Acquisition and Receipt of Biological Materials

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: HTA-PR-FORM/007

Version No:

1.0

Issue Date:

ARF No: CBE/ARF/000 66

PART A: To be completed by the	Receiver (a separate	e form must be o	completed for	each sample	type)			
A1. Details of Sample/Specimen								
Type/ID: (eg primary cell, cell line, tissu fluid, excreta, biological agent)	e, body Human Coro	Human Cord Blood (Ficolled) CD34+ isolation				⊠Human □Animal		
Format / Quantity: (eg vials, slides, e	tc) 2x50mL tub	2x50mL tube containing ~21mL ficol unit. 2 donors total						
Tissue site/Organ source:	Cord Blood	Cord Blood						
Batch N°:								
Is the sample/specimen considered to be Relevant Material under the Human Tissue Act (HTA)? If No, go to section A2.						⊠Yes □No		
Is the material obtained from an HTA licenced Tissue Bank with REC approval for generic research use?								
Is the material obtained for storage and use under a project specific NHS REC approval?								
Is the material licensable under the HTA? Indicate source below:								
☑HTA licensed organisation ☑C	ommercial Supplier	☐ Imported (fro	om outside Eng	gland, Wales c	or N.Ireland)			
If Yes, list lot numbers (or other identifier) & the corresponding assigned unique sample ID	Lot N°: G2212222	04742, G2212	22204720	Assigned Unique ID (Procuro): S00271852				
If Yes, provide Project Ref N°:	BRA060 & B	BRA060 & BRA010						
If Yes, provide the name of the PI	mas				,			
A2. Details of Receipt			6					
Date/Time of receipt	Date	05/10/2022	05/10/2022		16:20			
ID of Receiver	Name:	Jon Harriman		Dept:	СВЕ			
ID of Supplier/Provider	Name:	Anthony Nola	Anthony Nolan		UK	80		
PART B: To be completed by the	Receiver							
B1. Inspection and Quarantine								
Has a biological risk assessment for the use of this material been approved?		⊠Yes □No	Ref Number: BRA060 & BRA010					
Physical integrity of the material(s) acceptable?		⊠Yes □No	If No, describe action taken					
Quantity received correct?		.⊠Yes □No	If No, describe action taken					
Labelling correct and legible?	⊠Yes □No	If No, describe action taken						
Is the relevant documentation attached to this form? If No, add reference or details to ensure traceability	C of A or equivalen	t evidence of	⊠Yes □No □N/A					
		eements to enable transfer of		⊠Yes □No □N/A				
		Details/evidence/assurance of		⊠Yes □No □N/A				

	Other (describe) □Yes □No □N/A						
Temporary storage - Quarantine location (as applicable)	Building/Room	JH) into 1x 1mL vials @3.8		Processed immediately (H21, E6. Frozen via passive cooling ntil 06/10/22 and transferred				
	Storage Unit ID	Management of the Control of the Con						
	Within storage	thin storage unit location ID Bank 7 Rack 5 Box A 1						
	Date/Time of qu	uarantine	06/10/2022 12:00					
Submitted by:	Signature:	1	Date:					
PART C: To be completed by the departmental Quality Manager								
C1. Quality Assurance Checks				Ves □No □N/A				
Has the sample/specimen been								
Has the donor been screened for	☑Yes □No □N/A							
Has the sample/specimen been	□Yes □No □N/A							
Is there evidence that the suppl or other recognised certification	□Yes □No □N/A							
For HTA licensable material, is t and use of the material under the	√Yes □No □N/A							
Is there sufficient evidence to s	□Yes ☑No □N/A							
C2. Approval for release from quarantine								
Can the material be released from released for processing?	⊡Yes □No							
If No, provide recommendations for deposition of the material and the results of any action relating to non-conforming material.		☐ Accept as is, but with extra controls						
		☐ Rework or rep specified require	rocess to meet the ments					
		☐Test to meet s	pecified requirements	. ~.				
		☐Return to supp	olier/provider					
		□Disposal						
If Yes, provide details of storage location (as applicable)		Building/Room		-				
		Storage Unit ID		~				
		Within storage u	nit location ID	As above.				
		Database Refere	nce					
		Date/Time of transfer						
Approved by: Signature:			C 1/	Date:				
		*	C. Kavargh	10(10/22				