EUROPEAN UNION

	I.1. Consignor						I.2. IMSOC Reference				
	Name						I.2.a. Local Reference				
							1.2.a. Local Reference				
	Address										
	Country ISO Code										
	I.5. Consignee						I.3. Central co	mpetent a	uthority		
						I.4. Local com					
H	Address										
Ĕ	Country ISO Code										
E.											
ısi	I.7. Country of origin ISO Code					I.9. Country of	f destinatio	on		ISO Code	
<u>o</u>											
Part I : Details of consignment	I.8. Region of origi	n				Code	I.10. Region o	fdestinatio	n		
s o	I.11. Place of Dispa						I.12. Place of o				
ail	-					acountation.	L				
et:	Name					Name					
Ω	Address						Address	Approval Number			
Ξ	Approval Number	ſ		100	2.1.		Country ISO Code				
art	Country			ISO (lode		Country			ISO Code	
Ę,	I.13. Place of Load	ing					I.14. Date and time of departure				
	Name	0							1		
	Address										
		-									
	Approval Number	Ľ		100	2						
	Country			ISO (lode						
	I.15. Means of Trai	nsport					I.16 Entry Point				
	Mode	-	nal	Identificatio	m						
	transport										
		document									
							-				
							-				
							-				
								. ,			
	I.18. Transport conditions					I.17. Accompa	inying doci	uments			
	Ambient Controlled Frozen Chilled Chilled					Commercial document Date of issue reference					
							Country	Country Place of issue			
							Issue				
	.19. Container No / Seal No										
	I.20. Certified as										
	Artificial reproduction Breeding										
	I.21. For transit through a third country					I.22. For transit through Member State(s)					
	Country	-		ISO Code							
	EU Exit Authority BCP code					Country ISO Code					
	EU Entry Authority			BCP code							
	I.24. Total quantity					I.25. Total gross weight					
	1 2						, s	0			
	I.28. Description of consignment										
	I. 05 PRODUCTS OF ANIMAL ORIGIN, NOT ELSEWHERE SPECIFIED OR INCLUDED										
	0511 Animal products not elsewhere specified or included; dead animal 051199 Other					ls of Chapter 1	or 3, unfi	t for human consun	nption		
	05119985 Other										
			Specie	20		Identification	number	Identifica	ation mark	Nature of com	modity
	Commodity		opeere	.5		lucitimention	number	Identified			intourty
	Commodity										
					Quantity Date of collection/produ				Anticon Manufacturing plant		
	Commodity Quantity				Date of c	collection/produ			Manufacturing pla	nt	
					Date of c	ollection/produ			Manufacturing pla		
					Date of c	collection/produ			Manufacturing pla	nt 	
					Date of c	collection/produ			Manufacturing pla	<u>nt</u>	
					Date of c	collection/produ			Manufacturing pla	ML	
					Date of c	collection/produ			Manufacturing pla	<u>nu</u>	
					Date of c	ollection/produ			Manufacturing pla	<u></u>	
					Date of c	ollection/produ			Manufacturing pla	<u></u>	
					Date of c	ollection/produ			Manufacturing pla	<u></u>	

(GB) In vitro bovine embryos (approved semen centres) from EU countries GBHC008E (v3.1)

JROPEAN	UNION	< <i>/</i>		countries GBHC008E (v3.1)			
II. Health inf	formation						
I, the und	ersigned, off	icial veterinarian of	(exporting country) (2) ce	ertify that:			
II.1.	-	os to be exported		,			
	II.1.1.	-	orting country, which according	g to official findings:			
	II.1.1.1	was free from rinderpest d	luring the 12 months immediat	ely prior to their production;			
(1)either	○ [II.1.1.2.	was free from foot-and-mouth disease and lumpy skin disease during the 12 months immediately prior to their production and did not carry out vaccination against foot-and-mouth or lumpy skin disease during that period.]					
(1)either (1)or	○ [II.1.1.2.	was not free from foot-and-mouth disease or lumpy skin disease during the 12 months immediately prior to their production or carried out vaccination against foot-and-mouth disease or lumpy skin disease during that period, and					
	-	the embryos were produce	ed without penetration of the Z	ona pellucida			
	-	the embryos were stored u their production	under approved conditions for	at least 30 days immediately after			
	-	and-mouth disease or lump animal of a susceptible spe	py skin disease during the 30 da	ot-and-mouth disease or lumpy			
II.1.2	were prod	uced by the embryo product	tion team (3) which:				
	-	has been approved in acco	ordance with Chapter I of Anne	x A to Directive 89/556/EEC;			
	-	carried out the production with Chapter II of Annex A		port of the embryos in accordance			
	-	is subject to inspection by	an official veterinarian at least	t twice a year			
II.2.	an area of occurrence fever, cont their collec	s used in the production of the embryos to be exported were collected on premises situated in at least 10km radius centred on them, which according to official findings there was no of foot-and-mouth disease, epizootic haemorrhagic disease, vesicular stomatitis, Rift Valley agious bovine pleuropneumonia or lumpy skin disease in the 30 days immediately prior to tion and until their dispatch to Great Britain, in case of fresh embryos, or during the 30 days tion, in case of embryos subject to a mandatory storage for at least 30 days in accordance II.2.2.					
II.3.	the day of 10km radi and-mouth	e time of collection of the oocytes until 30 days thereafter or, in the case of fresh embryos, until of dispatch, the embryos to be exported were stored on premises situated in an area of at least dius centred on them, on which according to official findings there was no occurrence of foot- uth disease, vesicular stomatitis, Rift Valley Fever, contagious bovine pleuropneumonia or kin disease.					
II.4.	the donors	of oocytes used in the prod	uction of the embryos to be exp	ported:			
	II.4.1.	within a 10-km radius of w foot-and-mouth disease, bl	which according to official finding	lection of the oocytes, on premises ngs, there was no occurrence of agic disease, vesicular stomatitis, lumpy skin disease;			
	II.4.2.	showed no clinical signs of	f disease on the day of collection	n;			
	II.4.3.	spent the six months imme country in no more than tw		in the territory of the exporting			
	-	which, according to officia	al findings, were free from tube	erculosis during that time,			
	-	which, according to officia	al findings, were free from bruc	cellosis during that time,			
	-		ootic bovine leukosis or in whic s during the previous three year	ch no animal showed clinical signs rs,			
	-		ıl showed clinical signs of infect pustular vulvo-vaginitis during				
(1)either	○ [II.4.4.	were kept in a bluetongue during, collection of the oo	virus-free country or zone for a	at least 60 days prior to, and			

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БU	OROPEAN U	JINION							
	II. Health info	rmation							
Part II: Certification	(1)or • [II.4.4. were kept during a seasonally free period or protected from the vector for at least 60 days prior to, and during, the collection of the oocytes, and the embryos were produced without penetration of the zona pellucida, except if the donors underwent a serological test to detec antibodies to the bluetongue virus group, carried out in accordance with the WOAH (formerly OIE) Manual of Diagnostic Test and Vaccines for Terrestrial Animals between 21 and 60 days after collection and giving negative results and the embryos were stored for at least 30 days.]								
	(1)or	○ [II.4.4.	underwent a serological test to detect antibodies to the bluetongue virus group carried out in accordance with the WOAH (formerly OIE) Manual of Diagnostic Tests and Vaccines for Terrestrial Animals between 21 and 60 days after collection and giving negative results, and the embryos were stored for at least 30 days.]						
	(1)or	○ [II.4.4.	OIE) Manual of Diagnostic Tests an taken on the day of collection or the	identification test, carried out in accordance with the WOAH (formerly nostic Tests and Vaccines for Terrestrial Animals on a blood sample collection or the day of slaughtering and giving negative results – the n produced, in the latter case, without penetration of the zona					
	II.5.	The embryos to be exported were conceived by in vitro fertilisation using semen coming from semen collection or storage centres approved for the collection, processing and/or storage of semen by the competent authority of a third country or a part thereof listed in a document relating to 'bovine semen' published on gov.uk, in accordance with Decision 2011/630/EU (4) or by the competent authority of Great Britain.							
	Notes								
References to European Union legislation within this certificate are references to direct EU legislation w been retained in Great Britain (retained EU law as defined in the European Union (Withdrawal) Act 201 be viewed on the UK legislation website (legislation.gov.uk).									
	References to Great Britain in this certificate include Channel Islands and Isle of Man.								
	In accordance with Article 3(a) of Directive 89/556/EEC, the in vitro produced bovine embryos using semen from semen centres approved by the exporting country, imported under the conditions laid down in this certificate are excluded from further export to an EU member State; Liechtenstein; Norway and Switzerland Part I:								
	Box	Person responsible for the load in Great Britain: this box is to be filled in only if it is a certificate for transit commodity.							
	Box reference I.11:	Place of origin shall correspond to the embryo production teams from which the embryos are dispatch to Great Britain and listed in accordance with Article 8(2) of Directive 89/556/EEC.							
	Box reference I.22:	Number of packages shall correspond to the number of containers. ce							
	Box reference I.23:	Identification of container and seal number shall be indicated. e							
	Box reference I.26:	Fill in according to whether it is a transit or an import certificate.							
	Box reference I.27:	Fill in acco	rding to whether it is a transit or an	import certificate.					
	Box reference I.28:	Species: select amongst 'Bos taurus', 'Bison bison' or 'Bubalus bubalis' as appropriate. e							
		• • • •							

Category: select 'in vitro produced embryos'.

Dam identity shall correspond to the official identification of the animal.

EUROPEAN UNION

(GB) In vitro bovine embryos (approved semen centres) from EU

EUROPEAN UNION Countries GBHC008E (v3									
	II. Health info	rmation							
Part II: Certification	Date of free Approval r	 Delete as appropriate. Only third countries listed in a document relating to 'bovine embryos' published on gov.uk, in accordance with Decision 2006/168/EC.(5) 							
art II:	(4)	Only third countries listed in a document relating to 'bovine semen' published on gov.uk, in accordance with Implementing Decision 2011/630/EU. (5)							
	(5)	Documents relating to 'bovine semen' and 'bovine embryos' published by the Secretary of State, with the consent of the Scottish and Welsh Ministers, may be found here:							
	EU and EFTA states approved to export animals and animal products to Great Britain - data.gov.uk								
	Non-EU countries approved to export animals and animal products to Great Britain - data.gov.uk								
	The signature and the stamp must be in a different colour to that of the printing.								
	Certifying Officer								
	Name (in cap Date of signa		Qualification and title Signature						
	Stamp		0						