	UNITED KINGDOM				lth certificate to the EU
	I.1 Consignor/Exporter		I.2 Certificate ref	erence	I.2a
	Name				
	Address		I.3 Central Comp	etent Authority	
			DEPARTMENT FOOD & RURAL	OR ENVIRONMENT, AFFAIRS	
			I.4 Local Compet	ent Authority	
	Country	ISO country code	ANIMAL AND P	LANT HEALTH AGENCY	
ŀ	I.5 Consignee/Importer		I.6 Operator resp	onsible for the consignment	V
	Name		Name	-	
Ħ	Address		Address		
Part I: Description of consignment			. 1.0.0		
n 0	Country	ISO country code	Country	IS	O country code
tio	I.7 Country of origin	ISO country code	I.9 Country of de	stination	ISO country code
rip		\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \			
esc	I.8 Region of origin	Code	I.10 Region of des	stination	Code
D			O		
τI	I.11 Place of dispatch	Registration/Approval No	I.12 Place of desti	ination F	Registration/Approval No
Paı					
	Name		Name		
	Address		Address		
			*		
	Country	ISO country code	Country	IS	O country code
	I.13 Place of loading		I.14 Date and tim	e of departure	
		· /			
	I.15 Means of transport		I.16 Entry Borde	r Control Post	
	☐ Aircraft ☐	Vessel	I.17		
			1.17		
	□ Railway □	Road vehicle			
	Identification				
ŀ	I.18 Transport conditions	□ Ambient	☐ Chilled	□ Frozei	n
ŀ	I.19 Container number/Seal number				7
	Container No		Seal No		
ŀ	I.20 Certified as or for			*	
	☐ Germinal products				
	I.21		I.22	or internal market	
			1.22		
ļ	Third country	ISO country code	I.23	100	
	I.24 Total number of packages	I.25 Total quantity	•	I.26	

UNIT	TED KINGDOM	II.a	Certificate reference
I.27	Description of consignment		
1			

I.27 1	Description of consi	gnment		<u> </u>	
1	CN code	Species	Subspecies/Category	Identification number	Quantity
	Туре	Approval or registration nu	mber of plant/establishment/centre	Identification mark	Date of collection/production
2	CN code	Species	Subspecies/Category	Identification number	Quantity
	Туре	Approval or registration nu	mber of plant/establishment/centre	Identification mark	Date of collection/production
3	CN code	Species	Subspecies/Category	Identification number	Quantity
	Туре	Approval or registration nu	mber of plant/establishment/centre	Identification mark	Date of collection/production
4	CN code	Species	Subspecies/Category	Identification number	Quantity
	Туре	Approval or registration nu	mber of plant/establishment/centre	Identification mark	Date of collection/production
5	CN code	Species	Subspecies/Category	Identification number	Quantity
	Туре	Approval or registration nu	mber of plant/establishment/centre	Identification mark	Date of collection/production

			Cui		model bo v-m-vitto-Evib-b-Eivi Ki			
UNITI	UNITED KINGDOM			II.a	Certificate reference			
	II. Health	information		-				
		I, the unders	igned, official veterinarian of certify that:					
		$(exporting\ country)^{(1)}$						
	П.1.	The embryos	he embryos to be exported					
		II.1.1.	were produced in the exporting country, which according to official findings:					
		II.1.1.1	was free from rinderpest during the 12 month period immediately prior to their produced	luction;				
	⁽²⁾ either	[II.1.1.2.	was free from foot-and-mouth disease and lumpy skin disease during the 12 month period immediately prior to their production and did not carry out vaccination against foot-and-mouth disease or lumpy skin disease during that period.]					
	(2)or	[II.1.1.2. was not free from foot-and-mouth disease or lumpy skin disease during the 12 month period immediately prior to their production or car out vaccination against foot-and-mouth disease or lumpy skin disease during that period, and						
		the embryos were produced without penetration of the zona pellucida, the embryos were stored under approved conditions for at least 30 days immediately after their production, the donor females come from holdings on which no animal was vaccinated against foot-and-mouth disease or lumpy skin disease during the 30 days prior to collection and no animal of a susceptible species showed clinical signs of foot-and-mouth disease or lump skin disease during the 30 days prior to, and at least the 30 days after, the oocytes were collected.] Were produced by the embryo production team ⁽³⁾ which:						
			 had been approved in accordance with Chapter I of Annex A to Directive 89/5: carried out the production, processing, storing and transport of the embryos in 89/556/EEC; 					
	II.2.	- was subject to inspection by an official veterinarian at least twice a year. The oocytes used in the production of the embryos to be exported were collected on premises situated in an area of at least 10 km radius centred on them, on which according to official findings there was no occurrence of foot-and-mouth disease, epizootic haemorrhagic disease, vesicular stomatitis, Rift Valley fever, contagious bovine pleuropneumonia or lumpy skin disease in the 30 days immediately prior to their collection and until their dispatch to the Union, in case of fresh embryos, or during the 30 days after collection, in case of embryos subject to a mandatory storage for at least 30 days in accordance with point II.2.2.						
	П.3.	From the time of collection of the oocytes until 30 days thereafter or, in the case of fresh embryos, until the day of dispatch, the embryos to be exported were stored on premises situated in an area of at least 10 km radius centred on them, on which according to official findings there was no occurrence of foot-and-mouth disease, vesicular stomatitis, Rift Valley fever, contagious bovine pleuropneumonia or lumpy skin disease.						
<u>.</u>	II.4.	The donors of oocytes used in the production of the embryos to be exported:						
Part II: Certification		II.4.1.	were located, during the 30 days immediately prior to collection of the oocytes, on possible of ficial findings, there was no occurrence of foot-and-mouth disease, bluetongue, ep Valley fever, contagious bovine pleuropneumonia or lumpy skin disease;					
ၟႄ		II.4.2.	showed no clinical signs of disease on the day of collection;					
<u></u>		II.4.3.	spent the six months immediately prior to collection within the territory of the export	rting cou	intry in no more than two herds:			
Part II			 which, according to official findings, were free from tuberculosis during that ti which, according to official findings, were free from brucellosis during that tin which were free from enzootic bovine leukosis or in which no animal showed previous three years, in which no bovine animal showed clinical signs of infectious bovine rhinotr 	ne, I clinical	_			
	(2)		previous 12 months.					
	(2)either	[II.4.4.	were kept in a bluetongue virus-free country or zone for at least 60 days prior to, an		·			
	(2)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 detect antibodies to the bluetongue virus group, carried out in accordance with the 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.]					
	(2)or	[II.4.4.	underwent a serological test to detect antibodies to the bluetongue virus group, carrie Tests and Vaccines for Terrestrial Animals between 21 and 60 days after collections stored for at least 30 days.]					
	⁽²⁾ or	[II.4.4.	underwent an agent identification test, carried out in accordance with the OIE Ma Animals on a blood sample taken on the day of collection or the day of slaughtering produced, in the latter case, without penetration of the zona pellucida.]					
	II.5.	The embryos	to be exported were conceived by in vitro fertilisation using semen coming from	semen c	collection or storage centres approved for the			

This certificate is intended for entry into the Union of embryos of bovine animals, including when the Union is not the final destination of the embryos.

 $2011/630/EU^{(4)}$ or by the competent authority of a Member State.

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 subject to the conditions laid down in this certificate are excluded from intra-Union trade.

collection, processing and/or storage of semen by the competent authority of a third country or a part thereof listed in Annex I to Implementing Decision

In accordance with the Agreement on the withdrawal of the United Kingdom of Great Britain and Northern Ireland from the European Union and the European Atomic Energy Community, and in particular Article 5(4) of the Protocol on Ireland / Northern Ireland in conjunction with Annex 2 to that Protocol, references to European Union in this certificate include the United Kingdom in respect of Northern Ireland.

This animal health certificate shall be completed according to the notes for the completion of certificates provided for in Chapter 4 of Annex I to Commission Implementing Regulation (EU) 2020/2235.

UNITED KINGDOM

II.a Certificate reference

Part I:

Box I.6: "Operator responsible for the consignment": This box is to be filled in only if it is a certificate for transit commodity.

Box I.11: "Place of dispatch" shall correspond to the embryo production team from which the embryos are dispatch to the Union and listed in accordance with Article 8(2) of Directive 89/556/EEC on the Commission website:

http://ec.europa.eu/food/animal/semen_ova/bovine/ova_embryos_en.htm.

Box I.19: Identification of container and seal number shall be indicated.

Box I.21: Fill in according to whether it is a transit or an import certificate.

Box I.22: Fill in according to whether it is a transit or an import certificate.

Box I.24: Total number of packages shall correspond to the number of containers.

Box I.27: "Species": Select amongst "Bos taurus", "Bison bison" or "Bubalus bubalis" as appropriate.

"Type": Select "in vitro produced embryos".

"Identification number":

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

"Sire identity" shall correspond to the official identification of the animal.

"Approval or registration number of plant/establishment/centre" shall correspond to the embryo production team by which the embryos were produced, processed and stored; and listed in accordance with Article 8(2) of Directive 89/556/EEC on the Commission website: http://ec.europa.eu/food/animal/semen_ova/bovine/ova_embryos_en.htm.

Part II:

- Only third country, territory or zone thereof listed in Annex IX to Commission Implementing Regulation (EU) 2021/404 for embryos of bovine animals.
- (2) Delete as appropriate.

Official veterinarian

- Only embryo production teams listed in accordance with Article 8(2) of Directive 89/556/EEC on the Commission website: http://ec.europa.eu/food/animal/semen_ova/bovine/ova_embryos_en.htm.
- (4) Only third country, territory or zone thereof listed in Annex IX to Implementing Regulation (EU) 2021/404 for semen of bovine animals.

Name (in capital letters)	
Date	Qualification and title
Stamp	Signature