	UNITED KINGDOM				ealth certificate to the EU
	I.1 Consignor/Exporter		I.2 Certificate ref	erence	I.2a
	Name				
	Address		I.3 Central Comp	etent Authority	7
)			DEPARTMENT FOF	OR ENVIRONMENT, AFFAIRS	
′			I.4 Local Compet	ent Authority	7
	Country ISO c	ountry code	ANIMAL AND P	LANT HEALTH AGENCY	
•	I.5 Consignee/Importer		I.6 Operator resp	onsible for the consignmen	nt .
	Name		Name		
Ħ	Address		Address		
Part I: Description of consignment	0				
00 (ountry code	Country]	ISO country code
riptic	I.7 Country of origin ISO o	ountry code	I.9 Country of de	stination	ISO country code
: Desc	I.8 Region of origin Code		I.10 Region of des	stination	Code
rt I	I.11 Place of dispatch Reg	istration/Approval No	I.12 Place of desti	ination	Registration/Approval No
Pa					
	Name	\	Name		
	Address		Address		
		(
-		ountry code	Country		ISO country code
	I.13 Place of loading		L14 Date and tim	e of departure	
	I.15 Means of transport		I.16 Entry Borde	r Control Post	
	☐ Aircraft ☐ Vessel		I.17		
	□ Railway □ Road ve	chicle			
	Identification				
	I.18 Transport conditions	ambient	☐ Chilled	☐ Froz	zen
	I.19 Container number/Seal number				
	Container No		Seal No		
	I.20 Certified as or for				
	☐ Germinal products				
	I.21		I.22 🗆 Fo	or internal market	
	Third country ISO	country code	I.23		
•	I.24 Total number of packages	I.25 Total quantity		I.26	

TINIT	TED KINGDOM			II.a	Certificate reference
I.27	Description of consig	ınment			
1	CN code	Species	Subspecies/Category	Identification number	Quantity
	CN code	Species	Subspecies/Category	Identification number	Quantity
2	Туре		umber of plant/establishment/centre	Identification mark	Date of collection/production
	CN code	Species	Subspecies/Category	Identification number	Quantity
3	Туре	Approval or registration nu	umber of plant/establishment/centre	Identification mark	Date of collection/production
	CN code	Species	Subspecies/Category	Identification number	Quantity
4	Туре		amber of plant/establishment/centre	Identification mark	Date of collection/production
	CN code	Species	Subspecies/Category	Identification number	Quantity
5	Туре	Approval or registration nu	umber of plant/establishment/centre	Identification mark	Date of collection/production
3	CN code	Species	Subspecies/Category	Identification number	Quantity
	Туре	Approval or registration nu	umber of plant/establishment/centre	Identification mark	Date of collection/production

II.a Certificate reference

UNITED KINGDOM

II. Health information

I, the undersigned, official veterinarian of the certify that

(exporting country) (1)

- II.1. The embryos to be exported:
- **II**.1.1. were collected 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 collection;
- [II.1.1.2. was free from foot-and-mouth disease and lumpy skin disease during the 12 month period immediately prior to their collection and did not carry out vaccination against foot-and-mouth disease or lumpy skin disease during that period.]
- (II.1.1.2. was not free from foot-and-mouth disease or lumpy skin disease during the 12 months immediately prior to their collection or carried out vaccination against foot-and-mouth disease or lumpy skin disease during that period, and:
 - the embryos were not subjected to penetration of the zona pellucida,
 - the embryos were stored under approved conditions for at least 30 days immediately after their collection,
 - 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 lumpy skin disease during the 30 days prior to, and at least the 30 days after, the embryos were collected.]
- II.1.2. were collected by the embryo collection team (3) which :
 - had been approved in accordance with Chapter I of Annex A to Directive 89/556/EEC;
 - which carried out the collection, processing, storing and transport of the embryos in accordance with Chapter II of Annex A to Directive 89/556/EEC;
 - was subject to inspection by an official veterinarian at least twice a year.
- II.1.3. were collected and processed 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 dispatch to the Union, in the case of fresh embryos, or during the 30 days after collection, in the case of embryos subject to a mandatory storage for at least 30 days in accordance with point II.1.1.2.
- II.1.4. from the time of collection until 30 days thereafter or, in the case of fresh embryos until the day of their dispatch to the Union, they 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.
- II.1.5. were collected from the donor females, which:
 - II.1.5.1. were located, during the 30 days immediately prior to collection, 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, bluetongue, epizootic haemorrhagic disease, vesicular stomatitis, Rift Valley fever, contagious bovine pleuropneumonia or lumpy skin disease;
 - II.1.5.2. showed no clinical signs of disease on the day of collection;
 - II.1.5.3. spent the 6 months immediately prior to collection within the territory of the exporting country in no more than two herds:
 - which, according to official findings, were free from tuberculosis during that time,
 - which, according to official findings, were free from brucellosis during that time,
 - which were free from enzootic bovine leukosis or in which no bovine animal showed clinical signs of enzootic bovine leukosis during the previous 3 years,
 - in which no bovine animal showed clinical signs of infectious bovine rhinotracheitis/infectious pustular vulvo-vaginitis during the previous 12 months.
- II.1.6. The embryos to be exported were conceived by artificial insemination 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 part thereof listed in Annex I to Implementing Decision 2011/630/EU (4) or by the competent authority of a Member State.

Notes

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

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 animal health certificate include the United Kingdom in respect of Northern Ireland.

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

Part I:

Box reference I.11: "Place of dispatch": Indicate the unique approval number and the name and address of the embryo collection or

 $production \ team \ of \ dispatch \ of \ the \ consignment \ of \ embryos. \ Only \ embryo \ collection \ or \ production \ teams \ listed \ in$

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.

Box reference I.12: "Place of destination": Indicate the address and unique registration or approval number of the establishment of

destination of the consignment of embryos.

Box reference I.19: Seal number shall be indicated.

Certificate model BOV-in-vivo-EMB-B-ENTRY II.a Certificate reference

UNITED KINGDOM	
Box reference I.24:	Total number of packages shall correspond to the number of containers.
Box reference I.27:	"Species": Select amongst "Bos taurus", "Bison bison" or "Bubalus bubalis" as appropriate.
	"Type": Select " <i>in vivo</i> derived embryos".
7	"Identification number": Indicate the identification number of each donor animal.
	"Identification mark": Indicate the mark on the straw or other packages where embryos of the consignment are placed.
	"Date of collection/production": Indicate the date on which embryos of the consignment were collected or produced.
	"Approval or registration number of plant/establishment/centre": Indicate the unique approval number of the embryo
	collection team by which embryos of the consignment were collected, 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.
Y	"Quantity": Indicate the number of straws or other packages with the same mark.
	Quantity 1 material and manifest of statute of other parentages 11 miles state manifest
Part II:	
	y or territory, or zone thereof listed in Annex IX to Commission Implementing Regulation (EU) 2021/404 for embryos of
bovine animals. Delete if not ann	
Defete if not appl	
Only Childry Cool	lection teams listed in accordance with Article 8(2) of Directive 89/556/EEC on Commission website: pu/food/animal/semen_ova/bovine/ova_embryos_en.htm.
(4) OJ L 247, 24.9.20	
OJ L 247, 24.9.20	11, 0.52.
Official veterinarian	
Name (in capital letters)	
Date	Qualification and title
Date	Qualification and title
Stamp	Signature
1	
	•
	Y