$Model health certificate for imports of in vivo derived embryos of domesticanimals of the bovine species collected in accordance with Council Directive 89/556/EEC (Decision 2006/168) \ GBHC006E$

COUNTRY: Countries subject to Health certificate to Great Britain, transitional import arrangements (*) Channel Islands and Isle of Man

t	I.1. Consignor	I.2. Certifi reference nu		I.2.a. UNN		
en	Name Address	reference nu	mber			
ŭ		T. 2. Control Competent 2 liberthy				
i.g		I.3. Central Competent Authority				
consignment	Tel.	I.4. Local Competent Authority				
ö	I.5. Consignee Name			or the load in		
ק	Address	Britain, Channel Islands and Isle of Man Names				
he		Address				
tς	Postal Code					
dispatched	Tel.	Postal Code Tel.				
. <u>⊣</u>	I.7. Country of ISO I.8. Region of Code	I.9. Country		I.10. Region o	of Code	
_	origin code origin	destination	code	destination		
of						
Ø	I.11. Place of origin Names Approval number	I.12. Place Name	of destination	1		
	Address					
etail		Address				
De	Name Approval number					
· .	Address					
H		Postal Code				
4	Name Approval number Address					
Part	- Auditor					
щ	I.13. Place of loading	I.14. Date o	f departure			
	1.15. Flace of folding	1.11. Bacc 3	I depuiledie			
	I.15. Mean of transport	I.16. Entry	BCP in Great E	Britain, Channe	l Islands	
	Aeroplane Ship Railway wagon	or Isle of Man				
	Road vehicle Other Identification:	I.17.				
	Documentary references:					
	I.18. Description of commodity		I.19. Commod	ity code (HS co	ode)	
		05 11 99 85				
	I.20. Quantity		I.22. Number	of packages		
	I.23. Seal/Container No.	1.24.				
	I.25. Commodity certified for:					
	Artificial reproduction					
	П			П		
	I.26. For transit through Great Britain, Channel Islands and Isle of Man to third country	I.27. For import or admission into Great Britain, Channel Islands and Isle of Man				
	islands and isle of man to third country					
	Third country ISO code					
	I.28. Identification of the commodities					
	Species Breed Category Donor (Scientific identity	Date of collection	Date of freezing	Approval number of	Quantity	
	Name)	20110001011	110021119	the team		

Version 1.0 Nov 2020 1/4

In vivo derived bovine embryos

n	II. Health inf	formation		II.a. Certificate reference no	II.b UNN		
ίο				reference no			
at							
Certification							
iį	I, the undersigned, official veterinarian of the						
)rt	(exporting country) $(^2)$						
ပ	II.1. The embryos to be exported:						
	II.1.1.were collected in the exporting country, which according to official findings:						
H H II.1.1.1. was free from rinderpest during the 12 months immediate collection;			nmediately prior to their				
H (1) either [II.1.1.2. was free from foot-and-mou 12 months immediately prio				mouth disease and lumpy skin disease during the rior to their collection and did not carry out t-and-mouth disease or lumpy skin disease during			
	(¹) or	[II.1.1.2.					
	-The embryos were not subjected to penetration of the zona pellucida,						
	-The embryos were stored under approved conditions for at least 30 daysimmediately 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:					
		-has 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; -is subject to inspection by an official veterinarian at least twice a year.					
	II.1.3.	centred on the foot-and-mouth Valley fever, immediately proof fresh embry	em, on which according a disease, epizootic ha contagious bovine pleurior to their collections, or during the 30 d	nises situated in an area of to official findings there temorrhagic disease, vesicular propreumonia or lumpy skin of an and until dispatch to Greatly alays after collection, in the 30 days in accordance with	was no occurrence of lar stomatitis, Rift disease in the 30 days eat Britain, in the case ne case of embryos subject		
	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 Great Britain, 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	d from the donor female	es, which:			
	II.1.5.1.	I.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 clir	nical signs of disease	on the day of collection;			

en 2/4

In vivo derived bovine embryos

II. Health information	II.a. Certificate reference no	II.b UNN

- - 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 three 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 1 to Implementing Decision 2011/630/EU (4) or by the competent authority of Great Britain.

Notes

(*) Those countries subject to the transitional import arrangements include: an EU member State; Liechtenstein; Norway and Switzerland.

References to European Union legislation within this certificate are references to direct EU legislation which has been retained in Great Britain (retained EU law as defined in the European Union (Withdrawal) Act 2018).

References to Great Britain in this certificate include Channel Islands and Isle of Man.

Part I:

- Box I.6: 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 I.11: Place of origin shall correspond to the embryo collection team from which the embryos are dispatched to Great Britain and which is listed in accordance with Article 8(2) of Directive 89/556/EEC
- Box I.16: Do not use this box until the end of the transitional staging period.
- Box I.22: Number of packages shall correspond to the number of containers.
- Box I.23: Identification of container and seal number shall be indicated.
- Box I.26: Fill in according to whether it is a transit or an import certificate.
- Box I.27: Fill in according to whether it is a transit or an Import certificate.
- Box I.28: Species: select amongst 'Bos taurus', 'Bison bison' or 'Bubalus bubalis' as appropriate.

Category: select 'in vivo derived embryos'.

Donor identity shall correspond to the official Identification of the animal.

Date of collection shall be indicated in the following format: dd.mm.yyyy

Approval number of the team: shall correspond to the embryo collection team by which the embryos were collected, processed and stored; and listed in accordance with Article 8(2) of Directive 89/556/EEC.

en 3/4

COUNTRY: Countries subject to transitional import arrangements (*)

In vivo derived bovine embryos

II. Health information	I.2. Certificate reference no	I.2.a. UNN					
Part II:	Part II:						
(1) Delete as appropriate.							
(2) Only third countries listed in Annex 1 to Decision 2006/168/EC.							
(3) Only embryo collection teams listed in accordance with Article 8(2) of Directive 89/556/EEC							
- the signature and the stamp must be in a different colour to that of the printing.							
Official Veterinarian							
Name (in capital letters):	Qualification and title:						
Date:	Signature:						
Stamp:							

en 4/4