Export Health Certificate

- 1	I.1. Consignor					I.2. IMSOC Ref	ference			
	Name				I.2.a. Local Reference					
	Address									
	Country ISO Code									
	·									
	I.5. Consignee					I.3. Central competent authority				
nt	Name					I.4. Local competent authority				
E L	Address Country		ISO Code	0						
뎚	Country		130 Cou	е						
Part I : Details of consignment	I.7. Country of origin ISO Code				SO Code	I.9. Country of	f destinatio	n		ISO Code
8										
ot	I.8. Region of origin			Co	ode	I.10. Region of		n		
ils	I.11. Place of Dispatch				I.12. Place of destination					
eta	Name				Name					
Ă	Address					Address				
ii.	Approval Number	•	100.0	2-1-		Approval Number				
Part	Country		ISO C	Loae		Country ISO Code				
_	I.13. Place of Loadi	ng				I.14. Date and	time of dep	parture		
	Name									
	Address									
	Approval Number	•	*00.0							
	Country		ISO (Code						
	I.15. Means of Trar	isport				I.16 Entry Poi	nt			
	Mode	International	Identificatio	n						
		transport document								
		uocument								
	I.18. Transport con		_		_	I.17. Accompa	nying docu	iments		
	Ambient ☐ Controlled Frozen ☐ Chilled ☐ temperature ☐				Commercial					
		temperature 🗀				document reference		Д	ate of issue	
						Country		P.	lace of	
	I 10 Container N-					issue				
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ъс	EUROPEAN UNION (GB) Bovine semen – Section B from EU countries - GBHC004E (
Part II: Certification	II. Health information							
	I, the unde	, the undersigned official veterinarian, hereby certify that :						
	II.1		(name of exporting country) (2) has been free from rinderpest and foot-and-mouth disease the 12 months immediately prior to collection of the semen for export and until its date of ch and no vaccination against these diseases has taken place during the same period.					
	II.2	-	n described above was collected before 31 December 2004 at the semen collection centre					
		II.2.1	meets the conditions laid down in Chapter I of Annex A to Directive 88/407/EEC;					
		II.2.2	is operated and supervised in accordance with the conditions laid down in Chapter II of Annex A to Directive 88/407/EEC.					
	II.3	brucellosis	at which the semen to be exported was collected was free from rabies, tuberculosis, s, anthrax and contagious bovine pleuropneumonia during the period commencing 30 days e date of collection of the semen to be exported and the 30 days after collection.					
	II.4	At the time centre:	semen described above was collected, all bovine animals standing at the semen collection					
		II.4.1	came from herds and/or were born to dams which satisfy the conditions of paragraph 1(b)and (c) of Chapter I of Annex B to Directive 88/407/EEC;					
		II.4.2	had tested negative, within the 30 days preceding the quarantine isolation period, to:					
			the tests referred to in points 1(d)(i), (ii) and (iii) of Chapter I of Annex B to Directive 88/407/EEC, and					
			- a serum neutralisation test or an ELISA test for infectious bovine rhinotracheitis/infectious pustular vulvo-vaginitis, and					
			a virus isolation test (fluorescent antibody test or immunoperoxidase test) for bovine viral diarrhoea, deferred until the animal reached the age of 6 months in the case of younger animals;					
		had undergone the 30-day quarantine isolation period and had tested negative to the following health tests:						
			a serological test for brucellosis carried out in accordance with the procedure described in Annex C to Directive 64/432/EEC,					
			either an immunofluorescent antibody test or a culture test for Campylobacter fetus infection on a sample of preputial material or artificial vagina washings, or, in the case of a female animal, a vaginal mucus agglutination test,					
			a microscopic examination and culture test for Trichomonas foetus on a sample of preputial material or artificial vagina washings, or in the case of a female animal a vaginal mucus agglutination test;					
		II.4.4	had tested negative, at least once a year, to the routine tests referred to in points 1(a), (b) and (c) of Chapter II of Annex B to Directive 88/407/EEC.					
	II.5	At the time the semen described above was collected,						
		II.5.1	all female bovine animals in the centre had tested negative at least once a year to a vaginal mucus agglutination test for Campylobacter fetus infection, and					
		II.5.2	all bulls used for semen production had tested negative either to an immunofluorescent antibody test or to a culture test for Campylobacter fetus infection on a sample of preputial material or artificial vagina washings carried out in 12 months prior to collection.					
			to be exported was obtained from donor bulls which					
	II.6.1	satisfy the	onditions laid down in Annex C to Directive 88/407/EEC;					
	(1)either o	[II.6.2	were resident in the exporting country during the 6 months immediately prior to collection of the semen for export;]					
	(1)or ∘	[II.6.2	were imported from .(2) after spending less than 6 months in the exporting country and at the time of import satisfied the animal health conditions applying to donors of the semen which is intended for export to Great Britain;]					
		II.6.3	stand in a semen collection centre at which:					

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ᆮ	ROPEAN UNION			(GB) Bovine semen – Section B from EU countries - GBHC004E (V3.1)				
	II. Health info	rmation						
		(1)	either o [all bovine animals were not vaccinated against infectious bovine rhinotracheitis and tested negative at least once a year to a serum neutralisation test or an ELISA test for infectious bovine rhinotracheitis/infectious pustular vulvo-vaginitis;]					
Part II: Certification		(1)	or o	[bovine animals not vaccinated against infectious bovine rhinotracheitis tested negative, at least once a year, to a serum neutralisation test or an ELISA test for infectious bovine rhinotracheitis/infectious pustular vulvo-vaginitis, at which testing for infectious bovine rhinotracheitis was not carried out on bulls which had received their first vaccination against infectious bovine rhinotracheitis at the insemination centre after they had tested negative to a serum neutralisation test or an ELISA test for infectious bovine rhinotracheitis/infectious pustular vulvo-vaginitis and which had been regularly re-vaccinated at intervals of not more than 6 months since the first vaccination;]				
ļ	(1)either o	[II.6.4	have not be	een vaccinated against infectious bovine rhinotracheitis,]				
	(1)or 0	[II.6.4	have been vaccinated against infectious bovine rhinotracheitis in accordance with point II.6.3.,]					
		II.6.5	fulfil the import conditions for bovine semen laid down in the bluetongue chapter of the Terrestrial Animal Health Code of the World Organisation for Animal Health (WOAH (formerly OIE)), depending on the status of the country or zone of residence; ****					
		II.6.6	were resident in the country of export in which the following serotypes of epizoo haemorrhagic disease (EHD) exist: : and tested negative on two occumore than 12 months apart to an agar gel immunodiffusion test (3) and to a virus neutralisation test for all above-listed serotypes of EHD, carried out in approved lon samples of blood taken prior to and not less than 21 days following collection (semen; ***					
		II.6.7	haemorrha 6-monthly	ent in the country of export in which the following serotypes of epizootic agic disease (EHD) exist: : and tested negative, prior to entry and at intervals, to an agar gel immunodiffusion test (3) and a virus neutralisation test ve-listed serotypes of EHD, carried out in approved laboratory; **				
		II.6.8	test for Aka	ative on two occasions not more than 12 months apart to a serum neutralisation abane virus carried out in approved laboratory on samples of blood taken prior to s than 21 days following collection of the semen. *				
	II.7			ted was collected after the date on which the centre was approved by the thorities of the exporting country.				
	II.8		en to be exported was processed, stored and transported under conditions which satisfy the Directive 88/407/EEC prior to its amendment by Directive 2003/43/EC.					
	Notes							
	Deferences	to Furonce	n IInion logi	elation within this corrificate are references to direct EU legislation which has				

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) and can be viewed on the UK legislation website (legislation.gov.uk).

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

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 reference transit commodity.

I.6:

Box Place of origin shall correspond to the semen collection centre where the semen was collected.

reference

I.11: Box

Place of destination: this box is to be filled in only if it is a certificate for transit commodity.

reference

I.12: Box

Number of packages shall correspond to the number of containers.

reference

I.22:

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EUROPEAN UNION

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EUROPI	AN UNION (GB) Bovine semen – Section B fror	m EU countries - GBHC004E (v3.1)					
II. Hea	h information							
Box refer I.23:		seal number shall be indicated.						
Box reference I.26:		s a transit or an import certificate.						
Box refer I.27:		Fill in according to whether it is a transit or an import certificate.						
Box refer I.27: Box refer I.27: Box refer I.28:	nce prior to 31 December 2004 and	Donor identity shall correspond to the official identification of the animal; date of collection shall be prior to 31 December 2004 and indicated in the following format: dd/mm/YYYY; approval number of the centre shall correspond to the approval number of the approved semen collection centre where the semen was collected.						
Part l	:							
(1)	Delete as necessary.							
(2)	Only third countries listed in a with Commission Decision 201	document relating to 'bovine semen' 1/630/EU.(4)	published on gov.uk, in accordance					
(3)	Standards for EHD virus diagn Diagnostic Tests and Vaccines	ostic tests are described in the blueto for Terrestrial Animals.	ngue chapter of the Manual of					
(4)	A document relating to 'bovine Scottish and Welsh Ministers, I	semen' published by the Secretary of nay be found here:	f State, with the consent of the					
EU ar	d EFTA states approved to export an	mals and animal products to Great B	ritain - data.gov.uk					
****	Non-EU countries approved to	export animals and animal products	to Great Britain - data.gov.uk					
****	To be used only by Australia, C	anada and the USA.						
***	To be used only by Australia as	nd the USA.						
**	To be used only by Canada.							
*	To be used only by Australia.							
	gnature and the stamp must be in a gray officer	lifferent colour to that of the printing	<u>,</u>					
	(in capital letters) f signature	Qualification and title Signature						

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