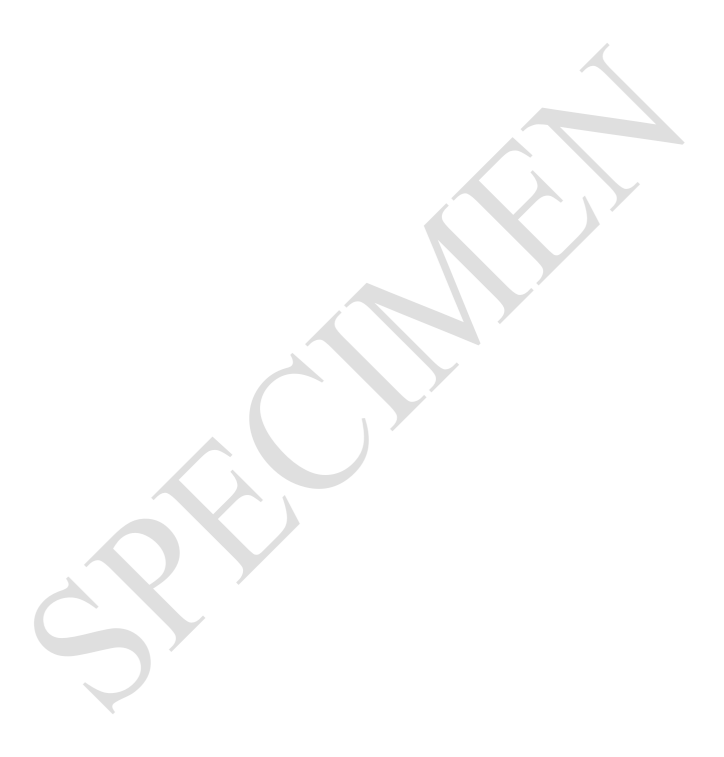
Model health certificate applicable to imports and transits of semen of domestic animals of the bovine species collected, processed and stored in accordance with Council Directive 88/407/EEC, dispatched from a semen collection centre where the semen was collected (Decision 2011/630) GBHC003E

COUNTRY: Countries subject to transition import arrangements (*)

Health certificate to Great Britain, Channel Islands and Isle of Man

nent	I.1. Consignor Name Address			I.2. Certifice reference no		I.2.a. UNN	
Part I: Details of dispatched consignment	Tel.			I.3. Central Competent Authority			
				I.4. Local Competent Authority			
	I.5. Consignee Name Address Postal Code Tel.			I.6. Person responsible for the load in Great Britain, Channel Islands and Isle of Man Name Address Postal Code			
	I.7. Country of origin	ISO code	I.8. Region Code of origin	Tel. I.9. Country destination	y of ISO code	I.10. Region of destination	Code
	I.11. Place of origin Name Approval number Address		I.12. Place of destination Name Address				
	Postal Code			Postal Code			
	I.13. Place of loading			I.14. Date of departure			
	I.15. Mean of transport Aeroplane Ship Railway wagon			I.16. Entry BCP in Great Britain, Channel Islands or Isle of Man			
	Road vehicle Other Identification:			I.17.			
	Documentary references: I.18. Description of commodity			I.19. Commodity code (HS code)			
				05.11.10			
	1.21.			I.20. I.22. Number of packages Quantity			
	I.23. Seal/Container No.			I.24. Type of packaging			
	I.25. Commodity certified for: Artificial reproduction						
	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			
	Third country ISO code						
	I.28. Identific			1			
	Species (Scientific	Donor identity	Identification of straw/s	Date of collection	Quantity	Information	
	Name)					BT(6)	EHD(6)
				<u> </u>			

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ű	II. Health information		II.a. Certificate reference no	II.b. UNN			
tic							
cat							
Certification	I, the undersigned official veterinarian, hereby certify that:						
rti	II.1.						
Ceı		(name of exporting country or part thereof)(2)					
II:	Was free from rinderpest and foot-and-mouth disease during the 12 months immediately princed to collection of the semen for export and until its date of dispatch to Great Britain and respectively.						
under the centre(3) described in box I.11. at which the semen to be					exported was collected:		
		II.2.1.	meets the conditions laid 88/407/EEC;	down in Chapter I(1) of A	nnex A to Directive		
		II.2.2.		d in accordance with the c A to Directive 88/407/EEC			
	II.3.	Brucellosis , anthrax and contagious bovine pleuropneumonia during the 30 days prior to the date of collection of the semen to be exported and the 30 days after collection (in the conformal of fresh semen until the day of dispatch to Great Britain).					
	II.3.						
	II.4.	II.4. The bovine animals standing at the semen collection centre:					
		(⁸) II.4.1.	come from herds which sat Annex B to Directive 88/4		ragraph 1(b) of Chapter I of		
	paragraph 1(c) of at the age of at 1 of Annex B to that II.4.3. underwent the test Annex B to Directi isolation period II.4.4. have satisfied the			months in accordance with	with the conditions of 88/407/EEC, or were tested paragraph 1(c) of Chapter II		
				red in accordance with par 07/EEC in the 28 days prec	ragraph 1(d) of Chapter I of eding the quarantine		
				tine isolation period and Chapter I of Annex B to D			
II.4.5. have undergone, at least once a year, the routine tests referred to in II of Annex B to Directive 88/407/EEC.							
	II.5. The semen to be exported was obtained from donor bulls which: II.5.1. satisfy the conditions laid down in Annex C of Directive 88/407/EEC;						
	(¹)eith	ner [II.5.		d in the exporting country for at least the last six to collection of the semen to be exported;			
	(¹) or	[II.5.	to the collection fromprior to the colle health conditions	of the semen since entry of the semen since entry of the semen since entry of the semen and sa applying to donors of the ct to Great Britain;]	and they were imported ess than six months tisfied the animal		
	II.5.3. comply with at least one of the following conditions as regards bluetongue, detailed in the table in point I.28.:						

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Bovine semen - Section A

. Health information		II.a. Certificate reference no	II.b. UNN
	L		
$(^1)$ either	[II.5.3.1. w		ongue virus-free country or 0 days prior to, and during, emen;]
(1) and/or	[II.5.3.2.	free period in a se	bluetongue virus seasonally asonally free zone for at leas nd during, collection of the
(¹) and/or	[II.5.3.3.		or-protected establishment for ior to, and during, collection
(¹) and/or	[II.5.3.4.	were subjected to a serological test for the detection of antibody to the bluetongue virus serogroup, carried out in accordance with the O Manual of Diagnostic Tests and Vaccines for Terrestrial Animals, with negative results, at least every 60 days throughout the collection period and between 21 and 60 days after the fin collection for this consignment of semen;]	
(¹) and/or	[II.5.3.5.	bluetongue virus, c the OIE Manual of D for Terrestrial Ani blood samples taken collection for this least every 7 days least every 28 days	n agent identification test for arried out in accordance with iagnostic Tests and Vaccines mals, with negative results, at commencement and final consignment of semen and at (virus isolation test) or at , if carried out as polymerase, , during collection for this n;]
11.5.4.			llowing conditions as regards D), as detailed in the table :
(¹) either	[II.5.4.1.		e exporting country which al findings is free from gic disease (EHD);]
(¹) (⁵) and/or	[11.5.4.2.	according to offici serotypes of epizoo exist: negative results in	e exporting country in which al findings the following tic haemorrhagic disease (EHD) and were subjected with each case to the following n an approved laboratory:
(1) either	[11.5.4.2.1.	antibody to the EHD on samples of blood more than 12 months	(4) for the detection of virus serogroup, carried out taken on two occasions not apart prior to and not less ing collection for this n;]
(¹) and/or	[11.5.4.2.2.	antibody to the EHD on samples taken at days throughout the	4) for the detection of virus serogroup, carried out intervals of not more than 60 collection period and between the final collection for the n 1

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COUNTRY: Countries subject to transition import arrangements (*)

Bovine semen - Section A

	ction import arrangements (*)				
II. Hea	lth information II.a. Certificate reference no II.b. UNN				
	(1) and/or [II.5.4.2.3. an agent identification test(4) carried out blood samples collected at commencement and conclusion of, and at least every 7 days (visolation test) or at least every 28 days, carried out as PCR, during collection for the consignment of semen.]	irus if			
	The semen to be exported was collected after the date on which the centre was approved by competent national authorities of the exporting country.	y the			
	The semen to be exported was processed, stored and transported under conditions which sat the terms of Directive $88/407/\text{EEC}$.	isfy			
Notes					
	se countries subject to the transitional import arrangements include: an EU member State;				
	nstein; 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).					
Referenc	ces 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 onl it is a certificate for transit commodity.	yif			
Box I.11	Box I.11: Place of origin shall correspond to the semen collection centre listed inaccordance with Article 9(2) of Directive 88/407/EEC and where the semen was collected.				
Box I.16	6: Do not use this box until the end of the transitional staging period.				
Box I.22	2: Number of packages shall correspond to the number of containers.				
Box I.23	3: Identification of container and seal number shall be indicated.				
Box I.26	6: Fill in according to whether it is a transit or an import certificate.				
Box I.2	7: Fill in according to whether it is a transit or an import certificate.				
Box I.28	8: Species: select amongst 'Bos taurus', 'Bison bison' or 'Bubalus bubalis' as appropriate.				
	Donor identity shall correspond to the official identification of the animal.				
	Date of collection shall be indicated in the following format: dd/mm/yyyy.				
	Quantity shall correspond to the number of straws of semen collected on a particulate from an identified donor bull complying with particular conditions for bluet and EHD.				

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COUNTRY: Countries subject to transition import arrangements (*)

Bovine semen - Section A

II. He	alth information	II.a. Certificate reference no	II.b. UNN			
Part I	I:					
(1)	Delete as necessary.					
(2)	Only third countries or parts thereof listed in Annex 1 to Implementing Decision 2011/630/EU.					
(3)	Only semen collection centres listed in accordance with Article 9(2) of Directive 88/407/EEC					
(4)	Standards for EHD virus diagnostic tests are described in the Bluetongue Chapter (2.1.3) of the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals.					
(5)	Compulsory for Australia, Canada and the United States.					
(6)	Referring to each straw or batch of straws indicate applicable condition (for example II.5.3.1).					
(7)	Referring to each straw or batch of straw (for example II.5.4.1 or II.5.4.2.1).	s indicate applicable cond	dition			
(8)	(8) For New Zealand, appearing with the entry 'XII' in column 6 of Part 1 of Annex 1 to Commission Regulation (EU) No 206/2010, officially tuberculosis-free bovine herds shall be considered equivalent to officially tuberculosis-free bovine herds in Great Britain recognised based on the conditions laid down in paragraphs 1 and 2 of Annex A.I to Council Directive 64/432/EEC.					
The si	gnature and the stamp must be in a differe	nt colour to that of the p	printing.			
Official Veterinarian						
Name (in capital letters):	Qualification and	Qualification and title:			
Date:		Signature:				
Stamp:						

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