	I.1. Consignor						I.2. IMSOC Re	eference				
	Name	ime						I.2.a. Local Reference				
	Address											
	Country ISO Code											
	I.5. Consignee						I.3. Central co	ompetent au	uthority			
Ŀ	Name						I.4. Local con	npetent autl	hority			
len	Address											
E	Country			ISO Cod	e							
Sig	I.7. Country of orig	7in				ISO Code	I.9. Country o	of destination	n			ISO Code
ü	1.7. country of ong	5111					1.5. Country C	or acountatio	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			
Part I : Details of consignment	IO Denien of onini					Code	I.10. Region of	£				
S 0	I.8. Region of origi											
ail	_							destination				
et	Address						Name Address					
-	Approval Number	r					Approval Nu	umber				
ц I	Country			ISO (	Code		Country ISO Code					
Pai												
	I.13. Place of Load	ing					I.14. Date and	d time of de	parture			
	Name											
	Address	_										
	Approval Number Country	r		ISO (	odo							
	country			130 (	Loue							
	I.15. Means of Tra	nsport					I.16 Entry Po	int				
	Mode	Internatio	nal	Identificatio	n							
		transport document										
	I.18. Transport cor			۸h:: 🗆	6.		I.17. Accomp		uments			
	Frozen 🗆	Chilled 🛛		Ambient 🗆	te	ontrolled mperature 🛛	Accompanyi					
						•	ng Date of issue reference					
										Place of		
							Country			issue		
	I.19. Container No	/ Seal No										
	I.20. Certified as											
	Artificial reproduc	rtion 🗌		Breeding 🗆	1							
	ni unclui reprodute											
	I.21. For transit th	rough a thii	rd coun	try			I.22. For transit through Member State(s)					
	Country			ISO Code								
	EU Exit Authority			BCP code			Country			ISO Code	2	
	EU Entry						iso code					
	Local Control     BCP code       Authority     I.24. Total quantity											
							I.25. Total gross weight					
	I.28. Description of consignment <b>1. 05</b> PRODUCTS OF ANIMAL ORIGIN, NOT ELSEWHERE SPECIFIED OR INC. <b>0511</b> Animal products not elsewhere specified or included; dead animals											
								1   or 3. unfit	t for human	consumpt	tion	
	<b>051199</b> Other			1			1			1		
	05119985	Other										
	Commodity		Specie	es		Identification	number	Identifica	tion mark	N	lature of con	nmodity
	Quantity				Date of o	collection/produ	ıction		Manufactur	ring plant		
										01		

-0	ROPEAN	onion				Holitz		Junineo	GBHC009E (V3	
	II. Health info	ormation								
	I, the unde	ersigned, offi	icial veterina	arian, hereb	y certify tha	t:	•			
	II.1.	the exporti	ing country		(name of e	exporting country)(2)				
ion		II.1.1.	caprine ple collection o	europneumo of the semen	nia and Rift to be expor	te des petits ruminants, sl Valley fever during the 1 ted and until its date of d took place during that pe	12 mo lispat	onths imm	ediately prior to	0
Part II: Certification		II.1.2.	collection o	of the semen	to be expor	n disease during the 12 m ted and until its date of d ok place during that perio	lispat			0
rt II: (	II.2.	The semen and stored		collection centre described in box I.11 and at which the semen to be exported was collect						
Pa		II.2.1.	meets the conditions for the approval of semen collection laid down in Chapter I(I)(1) of Annex D to Directive 92/65/EEC;							
		II.2.2.				rdance with the condition wn in Chapter I(II)(1) of A				
	II.3.	The ovine(	1)/caprine(1	) animals sta	anding at the	e semen collection centre	e:			
		II.3.1.	-		-	e accommodation describ		n point II.3	3.3,	
		(1)(4)eithe r	○ [II.3.1.1.	-		itory described in Box I.8, . melitensis)-free,]	, whic	ch has bee	en recognised as	3
		(1)or	○ [II.3.1.1.			ding which has obtained a is) – free status in accord				,]
	the last 12 months, nor against this disease, sa years ago, and all ovin subjected to at least tw taken on				ng, where in respect of bruve been free from clinical the of the ovine and caprin the those vaccinated with the e and caprine animals over the tasts (3), carried out with (date) and on within 30 days before entities	l or an ne ani Rev. 1 er six th neg (d	ny signs o mals have l vaccine r months o gative resu late) at lea	of this disease fo e been vaccinate more than two of age have beer ults on samples ast six months	ed n	
			and	have not be	en kept pre	viously in a holding of a l	lower	status		
			II.3.1.2.			iously for at least 60 days s (Brucella ovis) has been				
	prior to their stay in the complement fixation te sensitivity and specific than 50 ICFTU/ml;] II.3.1.3. to the best of my know contact with animals of system and according t following diseases has				the ovine species and ha e quarantine accommoda est, or any other test with ity, to detect contagious e	ation an ec	described Juivalent	l in point II.3.3 a documented		
					ledge do not come from h f a holding, in which, base o the written declaration been clinically detected w o their stay in the quarant	ed on mad vithin	the offici e by the o the perio	al notification wner, any of the ods referred to i	n	
					Mycoplasm	agalactia of sheep or goat a capricolum, Mycoplasn ithin the last six months,				rge
					paratuberc months,	ulosis and caseous lymph	naden	itis, withi	in the last 12	
				(c)	pulmonary	adenomatosis, within the	e last	three yea	ırs;	
			(1)either			a for sheep or caprine vir in the last three years;]	ral ar	thritis/end	cephalitis for	

JROPEAN	UNION			Ovine and caprine semen Section A (approved collection centr) from EU countries GBHC009E (v3.					
II. Health inf	ormation								
		(1)or	○ [(d)	Maedi/Visna for sheep or caprine viral arthritis/encephalitis for goats, within the last 12 months, and all the infected animals were slaughtered and remaining animals subsequently reacted negativel to two tests carried out at least six months apart;]					
	II.3.2.	have undergone the following tests carried out on a blood sample collected within the days preceding the commencement of the period of quarantine specified in point II.3.							
		-		sis (B. melitensis), with negative results in each case in accordance with to Directive 91/68/EEC;					
	us epididymitis (Brucella ovis, in the case of sheep only, with negative n each case in accordance with annex D to Directive 91/68/EEC, or any its with an equivalent documented sensitivity and specificity;								
		-		isease in accordance with point 1.4(c) of Chapter II(II) of Annex D to 92/65/EEC;					
	II.3.3.	accommo	-	arantine isolation period of at least 28 days in a quarantine cifically approved for the purpose by the competent authority and					
		II.3.3.1.	only ani accomm	mals of at least the same health status were present in the quarantine odation;					
		II.3.3.2.	approve not earli	als have undergone the following tests, carried out by the laboratory d by the competent authority of the exporting country on samples take er than 21 days after the animals were admitted to the quarantine odation, for:					
	brucellosis (B. melitensis) with negative results in each case in accordance with Annex C to Directive 91/68/EEC;								
		_	results ii	us epididymitis (Brucella ovis), in the case of sheep only, with negative n each case in accordance with Annex D to Directive 91/68/EEC, or any its with an equivalent documented sensitivity and specificity;					
		-		isease in accordance with point 1.6 of Chapter II(II) of Annex D to 92/65/EEC;					
II.3.4	have und	ergone at le	east once a	year the routine tests for:					
-	brucellos 91/68/EEC		(B. melitensis) with negative results in each case in accordance with Annex C to Directive						
-	accordan	is epididymitis (Brucella ovis), in the case of sheep only, with negative results in each case ce with Annex D to Directive 91/68/EEC, or any other tests with an equivalent documented y and specificity;							
-	border di	sease in acc	cordance w	h point 5(c) of Chapter II(II) of Annex D to Directive 92/65/EEC;					
II.4.	The seme	n to be exp	orted was o	btained from donor rams(1)/ bucks(1) which:					
	II.4.1.	were admitted to the approved semen collection centre with the express permission of the centre veterinarian.							
	II.4.2.			ns of disease on the day of admission to the approved semen collection by the semen was collected;					
(1)either	○ [II.4.3.		been vacci n of the sem	nated against foot-and-mouth disease during the 12 months prior to en;]					
(1)or	○ [II.4.3.	and 5% (	with a mini	d against foot-and-mouth disease at least 30 days prior to the collection mum of five straws) of each collection have been submitted to a virus t-and-mouth disease with negative results;]					
	II.4.4.			approved semen collection centre for a continuous period of at least 3 Fior to collection of the semen, in the case of collections of fresh semen					
	II.4.5.			urally after their entry to the quarantine accommodation described in o and including the day of semen collection;					
II.4.6	II.4.6.	have bee	n kept at ar	approved semen collection centres:					

<ul> <li>with negative results in each case to:]</li> <li>(1)either • [a serological test (6) for the detection of antibody to the EHDV group carr out in an approved laboratory on samples of blood taken on two occasions r more than 12 months apart prior to and not less than 21 days after the final collection for this consignment of semen.]]</li> <li>(1)or • [a serological test (6) for the detection of antibody to the EHDV group, carrout in an approved laboratory on samples of blood taken at intervals of not more than 60 days throughout the collection period and between 21 and 60 after the final collection for this consignment of semen.]</li> <li>(1)or • [an agent identification test(6) carried out in an approved laboratory on samples of blood taken at commencement and conclusion of, and at least ever days (virus isolation test) or at least every 28 days (PCR test) during collection this consignment of semen.]]</li> <li>II.4.10. have been kept continuously since birth in a country where the following conditions are fulfilled:</li> <li>II.4.10.1. classical scrapie is compulsorily notifiable;</li> <li>II.4.10.3. ovine and caprine animals affected with classical scrapie are killed and completely</li> </ul>						Hoin 20 (			
Upport         • [14.6.2]         which are situated in the case of fresh seme, until be date of dispatch, and which are situated in the centre an area of 10 kilometres radius in which there has been no case of forsh seme mouth disease for at least 30 days prior to collection of the semen:           [14.6.2]         which have been free, during the period commencing 30 days prior to collection of the semen or, in the case of fresh semu until the date of dispatch, from brueellosis (B. melltensis), contagious epidelymitis (Brucella oxis), anthrax and rabies;           (1)either         • [11.4.7]         have remained in the exporting country for at least the past six months prior to collection of the semen to: be exported.]           (1)or         • [11.4.7]         during the last six months prior to collection of the semen they complied with the animu hey have been imported into the exporting country at least 30 days prior to, and during, collection of the semen;           (1)or         • [11.4.8]         were kept in a bluetongue virus free country or zone for at least 60 days prior to, and during, collection of the semen;           (1)or         • [11.4.8]         were kept in a obluetongue virus seasonally free zone for at least 60 days prior to, and during, collection of the semen;           (1)or         • [11.4.8]         were kept in a necordance with the WOAH (formerly OIF) Manual of Diagnostic Tests and Vaccines for Terrestrial Animals, with negative results, on blood samples taken at la every 70 days throughout the collection period and between 21 and 60 days after the fine collection for this consignment of semen.]           (1)or         • [11.4.8]         were sus		II. Health info	rmation						
<ul> <li>and ending 30 days after collection of the semen or, in the case of fresh semu with the date of dispatch, from brucellosis (B. melitensis), contagious epididymitis (Brucella ovis), anthrax and rabies;</li> <li>(1)either 0 [11.4.7. have remained in the exporting country for at least the past six months prior to collection the semen to be exported.]</li> <li>(1)or 0 [11.4.7. during the last six months prior to collection of the semen they complied with the anima health conditions applying to donors of the semen which is intended for export to Graet Britian and they have been imported into the exporting country at least 30 days prior to collection of the semen from (2);]</li> <li>(1)either 0 [11.4.8. were kept in a butetongue virus free country or zone for at least 60 days prior to, and during, collection of the semen.]</li> <li>(1)or 0 [11.4.8. were kept in a vector protected establishment for at least 60 days prior to, and during collection of the semen.]</li> <li>(1)or 0 [11.4.8. were kept in a vector protected establishment for at least 60 days prior to, and during collection of the semen.]</li> <li>(1)or 0 [11.4.8. were subjected to a serological test for the detection of antibody to the bluetongue virus group, carried out in accord ance with the WOAH (formerly OID) Manual of Diagnostic T and Vaccines for Terrestrial Animals, with negative results, on blood samples taken at the every 60 days throughout the collection period and between 21 and 60 days (PCR test during collection for this consignment of semen.]</li> <li>(1)or 0 [11.4.8. were subjected to an agent identification test or at least every 28 days (PCR test during collection for this consignment of semen.]</li> <li>(1)or 0 [11.4.9. were resident in the exporting country in which according to official findings the follow serotypes of epizootic haemorrhagic disease (EHD)]</li> <li>(1)or 0 [11.4.9. were resident in the exporting country in which according to official findings the follow serotypes of epizo</li></ul>		prior to collection fresh semen, un an area of 10 ki			prior to collection of th fresh semen, until the an area of 10 kilometre	tion of the semen and 30 days after collection or, in the case of intil the date of dispatch, and which are situated in the centre of cilometres radius in which there has been no case of foot-and-			
<ul> <li>(1)either on attribute of the semen which is intended for export to creat Britian and they have been imported into the exporting country at least 30 days prior to collection of the semen from (2);]</li> <li>(1)either o [II.4.8. were kept in a bluetongue virus-free country or zone for at least 60 days prior to, and during, collection of the semen.]</li> <li>(1)or o [II.4.8. were kept during a bluetongue virus seasonally free zone for at least 60 days prior to, and during, collection of the semen.]</li> <li>(1)or o [II.4.8. were kept in a vector protected establishment for at least 60 days prior to, and during collection of the semen.]</li> <li>(1)or o [II.4.8. were subjected to a serological test for the detection of antibody to the bluetongue virus group, carried out in accordance with the WOAH (formerly OIE) Manual of Diagnostic T and Vaccines for Terrestrial Animals, with negative results, on blood samples taken at the every 60 days throughout the collection period and between 21 and 60 days after the fin collection for this consignment of semen.]</li> <li>(1)or o [II.4.8. were subjected to an agent identification test for bluetongue virus, carried out in accord with the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals with negative results on blood samples taken at to every 7 days (virus isolation test) or at least every 28 days (PCR test during collection for this consignment of semen.]</li> <li>(1)(5)eithe o [II.4.9. were resident in the exporting country in which according to official findings is free from epizootic haemorrhagic disease (EHD).]</li> <li>(1)or o [II.4.9. were resident in the exporting country on samples of blood taken on two occasions r more than 12 months apart prior to and not less than 21 days after the final collection for this consignment of semen.]</li> <li>(1)(5)eithe o [II.4.9. were resident in the exporting country on samples of blood taken on two occasions r more than 12 months apart prior to and not less than 21 days after the final collection for this co</li></ul>	tification	and ending 30 days af until the date of dispa				er collection of the semen or, ch, from brucellosis (B. melite	in the case of fresh semen,		
<ul> <li>Ineallin conditions applying to donors of the semen which is intended for export to creat Britain and they have been imported into the exporting country at least 30 days prior to collection of the semen from (2);</li> <li>(1)either • [II.4.8. were kept in a bluetongue virus-free country or zone for at least 60 days prior to, and during, collection of the semen;]</li> <li>(1)or • [II.4.8. were kept in a bluetongue virus seasonally free zone for at least 60 days prior to, and during, collection of the semen;]</li> <li>(1)or • [II.4.8. were kept in a vector protected establishment for at least 60 days prior to, and during collection of the semen;]</li> <li>(1)or • [II.4.8. were subjected to a serological test for the detection of antibody to the bluetongue virus group, carried out in accordance with the WOAH (formerly OIE) Manual of Diagnostic T and Vaccines for Terrestrial Animals, with negative results, on blood samples taken at the every 60 days throughout the collection period and between 21 and 60 days after the fin collection for this consignment of semen.]</li> <li>(1)or • [II.4.8. were subjected to an agent identification test for bluetongue virus, carried out in accord with the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals with negative results on blood samples taken at to every 60 days throughout the collection period and between 21 and 60 days (PCR test during collection for this consignment of semen.]</li> <li>(1)or • [II.4.9. were subjected to an agent identification test for bluetongue virus, carried out in accord with the exporting country which according to official findings is free from epizotic haemorrhagic disease (EHD);]</li> <li>(1)or • [II.4.9. were resident in the exporting country on samples of blood taken on two occasions r more than 12 months apart prior to a and to less than 21 days after the final collection for this consignment of semen.]</li> <li>(1)or • [II.4.9. were resident in the exporting country on samples of blood t</li></ul>	II: Cel	(1)either	○ [II.4.7.			untry for at least the past six	months prior to collection of		
during, collection of the semen;]         (1)or       ○ [II.4.8.         were kept during a bluetongue virus seasonally free zone for at least 60 days prior to, and during, collection of the semen;]         (1)or       ○ [II.4.8.         were kept in a vector protected establishment for at least 60 days prior to, and during collection of the semen;]         (1)or       ○ [II.4.8.         were subjected to a serological test for the detection of antibody to the bluetongue virus group, carried out in accordance with the WOAH (formerly OIE) Manual of Diagnostic T and Vaccines for Terrestrial Animals, with negative results, on blood samples taken at le every 60 days throughout the collection period and between 21 and 60 days after the fin collection for this consignment of semen;]         (1)or       ○ [II.4.8.       were subjected to an agent identification test for bluetongue virus, carried out in accord with the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals with negative results on blood samples taken at torumencement and final collection for this consignment of semen;]         (1)or       ○ [II.4.9.       were resident in the exporting country which according to official findings is free from epizootic haemorrhagic disease (EHD);]         (1)or       ○ [II.4.9.       were resident in the exporting country in which according to official findings the follow serotypes of epizootic haemorrhagic disease (EHD) exist: and were subject with negative results in each case to:]         (1)(or       ○ [II.4.9.       were resident in the exporting country in which according to official findings the follow serotypes of epiz	Part	(1)or	○ [II.4.7.	health con Britain an	ditions applying to donc d they have been import	ors of the semen which is inte ed into the exporting country	nded for export to Great		
(1)or       • [II.4.8.       were kept in a vector protected establishment for at least 60 days prior to, and during collection of the semen;]         (1)or       • [II.4.8.       were subjected to a serological test for the detection of antibody to the bluetongue virus group, carried out in accordance with the WOAH (formerly OIE) Manual of Diagnostic T and Vaccines for Terrestrial Animals, with negative results, on blood samples taken at le every 60 days throughout the collection period and between 21 and 60 days after the fin collection for this consignment of semen;]         (1)or       • [II.4.8.       were subjected to an agent identification test for bluetongue virus, carried out in accord with the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals with negative results on blood samples taken at commencement and final collection for this consignment of semen;]         (1)or       • [II.4.8.       were resident in the exporting country which according to official findings is free from epizootic haemorrhagic disease (EHD);]         (1)or       • [II.4.9.       were resident in the exporting country in which according to official findings the follow serotypes of epizootic haemorrhagic disease (EHD);         (1)or       • [II.4.9.       were resident in the exporting country in which according to official findings the follow serotypes of epizootic haemorrhagic disease (EHD);         (1)or       • [II.4.9.       were resident in the exporting country in which according to official findings the follow serotypes of epizootic haemorrhagic disease (EHD);         (1)or       • [II.4.9.       were resident in the exporting country in which according to		(1)either	○ [II.4.8.			ee country or zone for at leas	t 60 days prior to, and		
<ul> <li>collection of the semen;]</li> <li>(1)or • [II.4.8. were subjected to a serological test for the detection of antibody to the bluetongue virus group, carried out in accordance with the WOAH (formerly OIE) Manual of Diagnostic T and Vaccines for Terrestrial Animals, with negative results, on blood samples taken at le every 60 days throughout the collection period and between 21 and 60 days after the fin collection for this consignment of semen.]</li> <li>(1)or • [II.4.8. were subjected to an agent identification test for bluetongue virus, carried out in accord with the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals with negative results on blood samples taken at commencement and final collection for this consignment of semen.]</li> <li>(1)(5)eithe • [II.4.9. were resident in the exporting country which according to official findings is free from epizootic haemorrhagic disease (EHD).]</li> <li>(1)or • [II.4.9. were resident in the exporting country in which according to official findings the follow serotypes of epizootic haemorrhagic disease (EHD) exist: and were subject with negative results in each case to:]</li> <li>(1)either • [a serological test (6) for the detection of antibody to the EHDV group, car out in an approved laboratory on samples of blood taken on two occasions r more than 12 months apart prior to and not less than 21 days after the final collection for this consignment of semen.]</li> <li>(1)or • [a agent identification test(6) carried out in an approved laboratory on samples of blood taken at intervals of not more than 60 days throughout the collection period and between 21 and 60 after the final collection for this consignment of semen.]</li> <li>(1)or • [a nagent identification test(6) carried out in an approved laboratory on samples of blood taken at intervals of not more than 60 days throughout the collection period and between 21 and 60 after the final collection for this consignment of semen.]</li> <li>(1)or • [a nagent identification test(6) carried out in an ap</li></ul>		(1)or	○ [II.4.8.			us seasonally free zone for at	least 60 days prior to, and		
group, carried out in accordance with the WOAH (formerly OIE) Manual of Diagnostic T and Vaccines for Terrestrial Animals, with negative results, on blood samples taken at le every 60 days throughout the collection period and between 21 and 60 days after the fin collection for this consignment of semen.]         (1)or       • [II.4.8.       were subjected to an agent identification test for bluetongue virus, carried out in accord with the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals with negative results on blood samples taken at the mean and at least every 7 days (virus isolation test) or at least every 28 days (PCR test during collection for this consignment of semen.]         (1)(5)eithe       • [II.4.9.       were resident in the exporting country which according to official findings is free from epizootic haemorrhagic disease (EHD);]         (1)or       • [II.4.9.       were resident in the exporting country in which according to official findings the follow serotypes of epizootic haemorrhagic disease (EHD);]         (1)or       • [II.4.9.       were resident in the exporting country in which according to official findings the follow serotypes of epizootic haemorrhagic disease (EHD);]         (1)or       • [II.4.9.       were resident in a approved laboratory on samples of blood taken on two occasions r more than 12 months apart prior to and not less than 21 days after the final collection for this consignment of semen.]]         (1)or       • [ a serological test (6) for the detection of antibody to the EHDV group, car out in an approved laboratory on samples of blood taken at intervals of not more than 60 days throughout the collection period and between 21 and 60 after the final collection for this cons		(1)or	○ [II.4.8.			ablishment for at least 60 day	rs prior to, and during		
<ul> <li>with the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals with negative results on blood samples taken at commencement and final collection for this consignm of semen and at least every 7 days (virus isolation test) or at least every 28 days (PCR test during collection for this consignment of semen;]</li> <li>(1)(5)eithe • [II.4.9. were resident in the exporting country which according to official findings is free from epizootic haemorrhagic disease (EHD);]</li> <li>(1)or • [II.4.9. were resident in the exporting country in which according to official findings the follow serotypes of epizootic haemorrhagic disease (EHD) exist: and were subject with negative results in each case to:]</li> <li>(1)either • [a serological test (6) for the detection of antibody to the EHDV group carr out in an approved laboratory on samples of blood taken on two occasions r more than 12 months apart prior to and not less than 21 days after the final collection for this consignment of semen.]]</li> <li>(1)or • [a serological test (6) for the detection of antibody to the EHDV group, car out in an approved laboratory on samples of blood taken at intervals of not more than 60 days throughout the collection period and between 21 and 60 after the final collection for this consignment of semen.]</li> <li>(1)or • [an agent identification test(6) carried out in an approved laboratory on samples of blood taken at intervals of not more than 60 days throughout the collection of, and at least ever days (virus isolation test) or at least every 28 days (PCR test) during collectio this consignment of semen.]]</li> <li>II.4.10. have been kept continuously since birth in a country where the following conditions are fulfilled:</li> <li>II.4.10.1. classical scrapie is compulsorily notifiable;</li> <li>II.4.10.3. ovine and caprine animals affected with classical scrapie are killed and completely</li> </ul>		group, carried out in accordance w and Vaccines for Terrestrial Anima every 60 days throughout the colle				with the WOAH (formerly OIE) Manual of Diagnostic Tests als, with negative results, on blood samples taken at least ection period and between 21 and 60 days after the final			
r       epizootic haemorrhagic disease (EHD);]         (1)or       • [II.4.9.         were resident in the exporting country in which according to official findings the follow serotypes of epizootic haemorrhagic disease (EHD) exist: and were subject with negative results in each case to:]         (1)either       • [a serological test (6) for the detection of antibody to the EHDV group carr out in an approved laboratory on samples of blood taken on two occasions r more than 12 months apart prior to and not less than 21 days after the final collection for this consignment of semen.]]         (1)or       • [a serological test (6) for the detection of antibody to the EHDV group, carr out in an approved laboratory on samples of blood taken at intervals of not more than 60 days throughout the collection period and between 21 and 60 after the final collection for this consignment of semen.]]         (1)or       • [an agent identification test(6) carried out in an approved laboratory on samples of blood taken at intervals of not more than 60 days throughout the collection period and between 21 and 60 after the final collection for this consignment of semen.]]         (1)or       • [an agent identification test(6) carried out in an approved laboratory on samples of blood taken at commencement and conclusion of, and at least ev days (virus isolation test) or at least every 28 days (PCR test) during collectio this consignment of semen.]]         II.4.10.       have been kept continuously since birth in a country where the following conditions are fulfilled:         II.4.10.       classical scrapie is compulsorily notifiable;         II.4.10.2       an awareness, surveillance and mon		(1)or	○ [II.4.8.	with the M results on of semen a	fanual of Diagnostic Test blood samples taken at c and at least every 7 days	ts and Vaccines for Terrestrial Animals with negative commencement and final collection for this consignment (virus isolation test) or at least every 28 days (PCR test)			
<ul> <li>serotypes of epizootic haemorrhagic disease (EHD) exist: and were subject with negative results in each case to:]</li> <li>(1)either • [a serological test (6) for the detection of antibody to the EHDV group carr out in an approved laboratory on samples of blood taken on two occasions r more than 12 months apart prior to and not less than 21 days after the final collection for this consignment of semen.]]</li> <li>(1)or • [a serological test (6) for the detection of antibody to the EHDV group, carrout in an approved laboratory on samples of blood taken at intervals of not more than 60 days throughout the collection period and between 21 and 60 after the final collection for this consignment of semen.]]</li> <li>(1)or • [an agent identification test(6) carried out in an approved laboratory on samples of blood taken at commencement and conclusion of, and at least ever days (virus isolation test) or at least every 28 days (PCR test) during collection this consignment of semen.]]</li> <li>II.4.10. have been kept continuously since birth in a country where the following conditions are fulfilled:</li> <li>II.4.10.1. classical scrapie is compulsorily notifiable;</li> <li>II.4.10.2. an awareness, surveillance and monitoring system is in place;</li> <li>II.4.10.3. ovine and caprine animals affected with classical scrapie are killed and completely</li> </ul>		(1)(5)eithe $\circ$ [II.4.9. were resident in the exporting cou					ial findings is free from		
<ul> <li>out in an approved laboratory on samples of blood taken on two occasions r more than 12 months apart prior to and not less than 21 days after the final collection for this consignment of semen.]]</li> <li>(1)or         <ul> <li>[a serological test (6) for the detection of antibody to the EHDV group, car out in an approved laboratory on samples of blood taken at intervals of not more than 60 days throughout the collection period and between 21 and 60 after the final collection for this consignment of semen.]</li> <li>(1)or                 <ul></ul></li></ul></li></ul>		serotypes of epizootic haemorrhag				ic disease (EHD) exist:	ficial findings the following and were subjected		
<ul> <li>out in an approved laboratory on samples of blood taken at intervals of not more than 60 days throughout the collection period and between 21 and 60 after the final collection for this consignment of semen.]</li> <li>(1)or • [an agent identification test(6) carried out in an approved laboratory on samples of blood taken at commencement and conclusion of, and at least ever days (virus isolation test) or at least every 28 days (PCR test) during collectio this consignment of semen.]]</li> <li>II.4.10. have been kept continuously since birth in a country where the following conditions are fulfilled:</li> <li>II.4.10.1. classical scrapie is compulsorily notifiable;</li> <li>II.4.10.2. an awareness, surveillance and monitoring system is in place;</li> <li>II.4.10.3. ovine and caprine animals affected with classical scrapie are killed and completely</li> </ul>				(1)either	out in an approved lab more than 12 months a	oratory on samples of blood t apart prior to and not less tha	aken on two occasions not		
samples of blood taken at commencement and conclusion of, and at least eved days (virus isolation test) or at least every 28 days (PCR test) during collection this consignment of semen.]] II.4.10. have been kept continuously since birth in a country where the following conditions are fulfilled: II.4.10.1. classical scrapie is compulsorily notifiable; II.4.10.2. an awareness, surveillance and monitoring system is in place; II.4.10.3. ovine and caprine animals affected with classical scrapie are killed and completely				(1)or	out in an approved lab more than 60 days thro	oratory on samples of blood t oughout the collection period	aken at intervals of not and between 21 and 60 days		
fulfilled: II.4.10.1. classical scrapie is compulsorily notifiable; II.4.10.2. an awareness, surveillance and monitoring system is in place; II.4.10.3. ovine and caprine animals affected with classical scrapie are killed and completely				(1)or	samples of blood taken days (virus isolation te	at commencement and conc st) or at least every 28 days (P	lusion of, and at least every 7		
<ul><li>II.4.10.2. an awareness, surveillance and monitoring system is in place;</li><li>II.4.10.3. ovine and caprine animals affected with classical scrapie are killed and completely</li></ul>		II.4.10. have been kept continuously since				birth in a country where the	following conditions are		
<ul><li>II.4.10.2. an awareness, surveillance and monitoring system is in place;</li><li>II.4.10.3. ovine and caprine animals affected with classical scrapie are killed and completely</li></ul>			II.4.10.1.	classical so	crapie is compulsorily no	otifiable;			
II.4.10.3. ovine and caprine animals affected with classical scrapie are killed and completely									
destroyed:					caprine animals affected	• • •	lled and completely		

II. Health i	nformation								
	II.4.10.4. the feeding to ovine and caprine animals of meat-and-bone meal, or greaves of ruminar origin, as defined in the WOAH (formerly OIE) Terrestrial Animal Health Code, has beer banned and effectively enforced in the whole country for a period of at least the last sev years;								
(1)either (1)or II.5.	• • [II.4.11.	have been kept continuously for a period of the last three years preceding the date of the collection of the semen to be exported in a holding or holdings which has/have fulfilled during that period all the requirements set out in points 1.3(a) to (f) of Section A of Chapt of Annex 8 to Regulation (EC) No 999/2001, except during the period when they were kep a semen collection centre that complied during that period with the conditions set out in four indents of point 1.3(c)(iv) of that Section;]							
(1)or	○ [II.4.11.	are ovine animals of ARR/ARR prion protein genotype.]							
II.5.	The semer	to be exported:							
	II.5.1.	was collected after the date on whi competent authority of the exporti		e was approved by the					
	II.5.2.		was collected, processed, preserved, stored and transported in accordance with the requirements applicable to semen laid down in Chapter III(I) of Annex D to Directive 92/65/EEC;						
-	II.5.3.	was sent to the place of loading in a for semen to be subject to trade lai Directive 92/65/EEC and bearing th	d down in point 1.4 of Chapte	r III(I) of Annex D to					
(1)either	• [II.6.	No antibiotics were added to the se	emen.]						
(1)or	○ [II.6.	The following antibiotic or combin in the final diluted semen of not lea	nation of antibiotics was added to produce a concentration ess than(7): ]						
Notes									
been ret	ained in Grea	an Union legislation within this certi t Britain (retained EU law as defined egislation website (legislation.gov.uk	in the European Union (With						
		ritain in this certificate include Chan							
Part I:									
Box referenc 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 referenc I.11:		rigin shall correspond to the approve and listed in accordance with Article	ved semen collection centre in which the semen was e 17(3)(b)of Directive 92/65/EEC.						
Box referenc I.22:		f packages shall correspond to the n	umber of containers.						
Box referenc I.23:		tion of container and seal number sh	ll be indicated.						
BoxFill in according to whether it is a transit or an import certificate.referenceI.26:BoxFill in according to whether it is a transit or an import certificate.referenceI.27:									
						Box Species: Select amongst 'Ovis aries' or 'Capra hircus' as appropriate. reference			
I.28:									

EU	IROPEAN U	JNION	from EU d	Countries GBHC009E (v3.0)
	II. Health info	rmation		
	Approval n Box I.11	umber of the centre shall correspond to the ap	proval number of the semen of	collection centre indicated in
	Part II:			
	(1)	Delete as necessary.		
tion	(2)	Only third countries as set out in a document r in accordance with Commission Decision 2010,		semen' published on gov.uk,
icat	(3)	Tests shall be carried out in accordance with A	nnex C to Directive 91/68/EE0	2.
Part II: Certification	(4)	Only for the territory appearing with the entry published on gov.uk, in accordance with Comm		
Part II:	(5)	See remarks for exporting country as set out ir published on gov.uk, in accordance with Comm	•	-
	(6)	Standards for EHD virus diagnostic tests are de Manual of Diagnostic Tests and Vaccines for Te	-	ne WOAH (formerly OIE)
	(7)	Insert names and concentrations.		
	(8)	Documents relating to 'ovine and caprine seme by the Secretary of State, with the consent of th	8	-
	EU and EFT	TA states approved to export animals and anima	al products to Great Britain - o	lata.gov.uk
		are and the stamp must be in a different colour	to that of the printing.	
	Certifying Offi Name (in cap Date of signa Stamp	ital letters)	Qualification and title Signature	
	1			