	I.1. Consignor				I.2. IMSOC Re	ference			
	Name				I.2.a. Local Reference				
	Address								
	Country		ISO Code						
	I.5. Consignee				I.3. Central co	mpetent authority			
	Name					petent authority			
en	Address								
gnm	Country		ISO Code						
Part I : Details of consignment	I.7. Country of origin			ISO Code	I.9. Country of destination ISO			ISO Code	
of (I.8. Region of origin			Code	I.10. Region o	f destination			
ls (I.11. Place of Dispatch				I.12. Place of o	destination			
tai	Name				Name				
Be	Address				Address				
	Approval Number				Approval Nu	mber			
art	Country		ISO Code		Country ISO Code				
Ä	I.13. Place of Loading				I.14. Date and	time of departure			
	Name								
	Address								
	Approval Number Country		ISO Code						
	-		130 couc						
	I.15. Means of Transpo		x 1		I.16 Entry Poi	nt			
	trar	ernational Isport	Identification						
	doc	ument							
						· · ·			
	I.18. Transport condition Frozen Chil	led 🗆	Ambient 🗆	Controlled	I.17. Accompanying documents				
				temperature \Box	Accompanyi ng Date of issue document				
					document part of issue reference				
					Country Place of issue				
	I.19. Container No / Sea	l No			1				
	I.20. Certified as								
	Artificial reproduction		Breeding 🗆						
	I.21. For transit throug	h a third coun			I.22. For transit through Member State(s)				
	Country EU Exit		ISO Code						
	Authority		BCP code		Country		ISO Code		
	EU Entry Authority		BCP code						
	I.24. Total quantity				I.25. Total gross weight				
	I.28. Description of con	signment							
	1.05 PRODUCTS OF AN								
	-	cts not elsewh	ere specified or inc	luded; dead anima	ls of Chapter 1	or 3, unfit for human	consumption		
	051199 Other 05119985 Othe	r							
	Commodity	Specie		Identification	number	Date of	Quantit		
	Commounty	Specie	.5	Identification	Inninger	collection/production	Quantit	у	

	II. Health information										
_	I, the unde hereby cer	-	cial veterin	arian, of the	e exporting c	ountry (2)	(nan	ne of exporting country)			
	II.1	stored for e	The semen collection centre (3), in which the semen described above was collected, processed and stored for export to Great Britain is approved and supervised by the competent authority in accordation with the conditions of the Chapters I(I)(1) and I(II)(1) of Annex D to Directive 92/65/EEC (4);								
	II.2	until the da	-	or chilled s		or to the date of first collection of the semen described above as dispatched or until the 30 days storage period for frozen					
			II.2.1.	to Article 1	-	e 2009/156/EC (5), in th		f regionalisation according tof the territory of the			
			-			fected with African ho f Directive 2009/156/E		ness in accordance with			
			-	free from V	/enezuelan e	equine encephalomyel	itis for	a period of at least 2 years,			
			-	free from glanders and dourine for a period of at least 6 months;							
			II.2.2.		e conditions C and in par	for a holding laid dow ticular:	'n in Ar	ticle 4(5) of Directive			
		(1)	○ either	[II.2.2.1.	species sus	case of a disease ment ceptible to that disease l or killed and the hold	locate	-			
				-	months, beg			tis for a period of at least 6 the equidae suffering from			
				-	to obtain a or Coggins a animals we	negative result in an a test) carried out on sar	gar gel nples ta	r at least the period required immunodiffusion test (AGID aken after the infected ons 3 months apart from			
				-	from vesicu the last reco		a perio	d of at least 6 months from			
				-	from rabies case,	for a period of at leas	t one n	nonth from the last recorded			
				-	from anthra case,]	ax for a period of at lea	ast 15 d	lays from the last recorded			
		(1)	• or	[II.2.2.1.	species sus slaughtered was free for encephalon and rabies which follo	ceptible to that disease l or killed and the prer r a period of at least 30 nyelitis, equine infection or 15 days in the case of	e locate mises d) days f ous ana of anth: of the ar	below all the animals of d in the holding have been isinfected, and the holding from any type of equine hemia, vesicular stomatitis rax, beginning on the day on himals the disinfection of the			
			II.2.3.		only equidae gious equine		inical s	igns of equine viral arteritis			
		II.3.	Prior to en located in t		emen collecti	ion centre the donor st	tallions	and any other equidae			
			II.3.1.	directly im exporting of of Directive	ported from country or, in	Great Britain during t n the case of regionalis C, in that part of the te	the 3 m sation i	(or since entry if they were onths period) in the n accordance with Article 13 of the exporting country			
			-	not considered to be infected with African horse sickness in accordance with Article 5(2)(a) and (b) of Directive 2009/156/EC,							

(GB) Equine semen – Section A from EU countries GBHC046E (v3.0)

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	II. Health info	ormation										
			-	free from V	Venezuelan	equine encephalomyelitis for a period of at least 2 years,						
	d dourine for a period of at least 6 months;											
	(1)	\circ either	[II.3.2.	originated from the country of export which was on the day of admission into the centre free from vesicular stomatitis (VS) for a period of at least 6 months,]								
Part II: Certification	(1)	∘ or	[II.3.2.	out with a with a neg Diagnostic	were subjected to a virus neutralisation test for vesicular stomatitis (VS) carried out with a negative result at a serum dilution of 1 in 32 or a VS ELISA carried out with a negative result in accordance with the relevant Chapter of the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals of the WOAH (formerly OIE) on a blood sample taken (6) within 14 days prior to entering the centre;]							
rt II: C			II.3.3.	-	from holdir ements of po	ngs which on the day of admission onto the centre fulfilled pint II.2.2;						
Pai		II.4.	The semer	n described a	above was c	ollected from donor stallions which:						
		II.4.1. did not show any clinical sign of an infectious or contagious dis of admission onto the semen collection centre and on the day the collected;										
			II.4.2.	were kept for a period of at least 30 days prior to the date of semen collection in holdings where no equine animal has shown any clinical sign of equine viral arteritis or contagious equine metritis during that period;								
			II.4.3.	date of firs	st semen coll	ural mating during a period of at least 30 days prior to the lection and between the dates of the first sample referred l.5.2 and/or II.4.5.3 and until the end of the collection						
			II.4.4.	II.4.4. underwent the following tests, which meet at least the requirements of the relevant Chapter of the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals of the WOAH (formerly OIE), carried out in a laboratory which is recognised by the competent authority and has the tests referred to hereinafter included in its accreditation equivalent to that provided for in Article 37 of Regulation (EU) No 2017/625 (7), as follows:								
			(8)	∟ [II.4.4.1.	test (AGID	infectious anaemia (EIA), an agar-gel immuno-diffusion or Coggins test) or an enzyme-linked immunosorbent SA) for equine infectious anaemia with a negative result;]						
				II.4.4.2.	for equine	viral arteritis (EVA)						
			(1)	🗆 either	[II.4.4.2.1.	a serum neutralisation test with a negative result at a serum dilution of one in four;]						
			(1)	□ and/or	[II.4.4.2.2.	a virus isolation test, polymerase chain reaction (PCR) or real time PCR with a negative result on an aliquot of the entire semen of the donor stallion;]						
				II.4.4.3.	ous equine metritis (CEM), an agent identification test t on three specimens (swabs) taken from the donor stallion asions with an interval of not less than 7 days at least enile sheath (prepuce), the urethra and the fossa glandis;							
					es were in no case taken earlier than 7 days (systemic or 21 days (local treatment) after antimicrobial treatment or stallion and were placed in transport medium with harcoal, such as Amies medium, before dispatch to the where they were subjected with a negative result to a test							
			(1)	□ either	[II.4.4.3.1.	the isolation of Taylorella equigenitalis after cultivation under microaerophilic conditions for a period of at least 7 days, set up within 24 hours after taking the specimens from the donor animal, or 48 hours where the specimens are kept cool during transport;]						

Part II: Certification

IL Health information		(61	, Lquille o						
II. Health information									
	(1)	□ and/or	[II.4.4.3.2.	PCR or rea	-	aylorella equigenital l out within 48 hours ne donor animal;]	-		
	II.4.5.	one of the		4.4 in each case to at points 1.6(a), (b) and ws:	I				
	(9)	□ [II.4.5.1.	The donor stallion was continuously resident on the semen collection centre for a period of at least 30 days prior to the date of the first collection and during the period of collection of the semen described above, and no equidae on the semen collection centre came during that time into direct contact with equidae of lower health status than the donor stallion. The tests described in point II.4.4 were carried out on samples taken (6) from the donor stallion at least once a year at the beginning of the breeding season or prior to the first collection of semen intended for imports into Great Britain of fresh, chilled or frozen semen and not less than 14 days following the date of the commencement of the residence period of at least 30 days prior to the first semen collection.]						
(9) □ The donor stallion was resident on the semen collection ce [II.4.5.2. [II.4.5.2. period of at least 30 days prior to the date of the first collect during the period of collection of the semen described above the semen collection centre under the responsibility of the veterinarian for a continuous period of less than 14 days, at other equidae on the semen collection centre came in to di contact with equidae of a lower health status. contact with equidae of a lower health status.							on and but left ntre l/or		
			The test described in point II.4.4 were carried out on samples taken (6) from the donor stallion at least once a year at the beginning of the breeding season or prior to the date of the first collection of semen intended for imports into Great Britain of fresh, chilled or frozen semen and not less than 14 days following the date of the commencement of the residence period of at least 30 days prior to the first semen collection.						
		and	into Great	Britain of fi	resh, chilled or froz	nen intended for imp zen semen the donor oint II.4.4., as follows	stallion		
			(a)	in point II taken (6) r	.4.4.1 was last carri	a, one of the tests de ied out on a sample c ays prior to the colle	of blood		
			(b)	for equine	e viral arteritis, one	e of the tests describe	d		
			(1) • either [in point II.4.4.2. was last carried out on a sample taken (6) not more than 30 days put to the date of the collection of the semen described above;]						
			(1)	o or	aliquot of the ent stallion taken (6) prior to the date semen described taken (6) from th months period re in a serum neutr	2 was carried out on the semen of the dom not more than 6 more of the collection of the above and a blood s e donor stallion duri eacted with a positive alisation test for equi- um dilution of more t	or nths ne ample ng the 6 e result ine viral		
					semen described taken (6) from th months period re in a serum neutr arteritis at a seru	above and a bloo e donor stallion d eacted with a posi alisation test for e	od sa luri tive equi		

Part II: Certification

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	II. Health information					
				(c)	II.4.4.3. w taken (6) i	gious equine metritis, the test described in point as last carried out on three specimens (swabs) not more than 60 days prior to the date of the of semen described above
				(1)	\circ either	[on two occasions;]
				(1)	\circ or	[on a single occasion and subjected to a PCR or real-time PCR.]]
		(9)	□ [II.4.5.3.	1.6(a) and	(b) of Chap	es not meet the conditions set out in points ter II of Annex D to Directive 92/65/EEC and the r imports into Great Britain of frozen semen.
T (1) T				out on san	nples taken	a points II.4.4.1, II.4.4.2 and II.4.4.3 were carried (6) from the donor stallion at least once a year le breeding season,
			and	samples ta of the sem collection semen col	iken (6) from en of a min of the seme lection cent	points II.4.4.1 and II.4.4.3. were carried out on m the donor stallion during the storage period aimum period of 30 days from the date of the en and before the semen is removed from the tre, not less than 14 days and not more than 90 on of the semen described above,
			and	(1)	○ either	[the tests for equine viral arteritis described in point II.4.4.2 were carried out on samples taken (6) during the storage period of the semen of a minimum period of 30 days from the date of collection of the semen and before the semen is removed from the semen collection centre or used, not less than 14 days and not more than 90 days after the date of the collection of the semen described above.]
				(1)	° or	[the non-shedder state of a donor stallion seropositive for equine viral arteritis was confirmed by a virus isolation test, PCR or real-time PCR carried out with a negative result on samples of an aliquot of the entire semen of the donor stallion taken (6) twice a year at an interval of at least 4 months and the donor stallion has reacted with a positive result at a serum dilution of at least one in four in a serum neutralisation test for equine viral arteritis.]
		II.4.6.		t the testing he followin	-	or in points 🗆 II.3.2.(1) and II.4.5 on samples

EU	JROPEAN U	JNION		(GB) Equine se	emen – S	ection A fro	m EU cou	ntries GBHC	046E (v3.0)
	II. Health info	rmation								
	Identificat ion of semen	Test programm e	Start date(6)		Date of san	pling for	r health tests(6)		
			Donor residence	Semen collection	VS(1) II.3.2	EIA II.4.4.1.	EVA II.4.4.2.		CEM II.4.4	3.
Part II: Certification							Blood sample	Semen sample	1. Sample	2. Sample
art II: Ce										
	(1)	\circ either	[II.5.	No antibio	tics were ad	ded to the	e semen;]			
	(1)	∘ or	[II.5.				bination of ar d semen of n		as added to pi (10):	roduce a ;]
		II.6.	The semen	described a	bove was:					
			II.6.1.						litions which o nnex D to Dire	
			II.6.2.		(I) of Annex	0			rdance with p ring the numb	

	II. Health info	rmation								
Part II: Certification										
	Notes									
	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 reference I.11:	The place of origin shall correspond to the semen collection centre of the semen origin.								
	Box reference I.22:	The number of packages shall correspond to the number of containers.								
	Box reference I.23:	eference								
	Box reference I.28:	The donor identity shall correspond to the offi	cial identification of the anim	al.						
	The date of	e date of collection shall be indicated in the following format dd/mm/yyyy.								
	Part II:									
	Guidance for the completion of the table in point II.4.6.									
	Abbreviati	Abbreviations:								
	VS	esicular stomatitis (VS) testing if required in accordance with point II.3.2								
	EIA-1	Equine infectious anaemia (EIA) testing first o	occasion							
	EIA-2	EIA testing second occasion								
	EVA-B1	Equine viral arteritis (EVA) testing on blood sa	ample first occasion							
	EVA-B2	EVA testing on blood sample second occasion	l							
	EVA-S1	EVA testing on semen sample first occasion								
	EVA-S2	EVA testing on semen sample second occasion								
	CEM-11	Contagious equine metritis (CEM) testing first	occasion first sample							
	CEM-12	CEM testing first occasion second sample taken	n 7 days after CEM-11							
	CEM-21	CEM testing second occasion first sample								
	CEM-22	CEM testing second occasion second sample ta	ken 7 days after CEM-21							
	Instructions:									
	For each semen identified in column A in correspondence with Box I.28, the test programme (points II.4.5.1, II.4.5.2 and/or II.4.5.3) shall be specified in column B, and columns C and D shall be completed with the dates required.									
	The dates when samples were taken for laboratory testing prior to the first collection of the semen described above as required in points II.4.5.1, II.4.5.2 and II.4.5.3, shall be entered in the upper line of columns 5 to 9 of the table, this being the boxes marked with EIA-1, EVA-B1 or EVA-S1 and CEM-11 and CEM-12 in the example below.									
	The dates when samples were taken for repeat laboratory testing as required in accordance with point II.4.5.2. or II.4.5.3. shall be entered in the lower line of columns 5 to 9 in table, this being the boxes EIA-2, EVA-B2 or EVA-S2 and CEM-21 and CEM-22 in the example below.									

II. Health info	rmation										
	Ination										
Identificat ion of semen	Test Start date Date of sampling for health tests programm										
	-	Donor residence	Semen collection	VS II.3.2.	EIA II.4.4.1	EVA II.4.4.2		CEM II.4.4	.3		
А						Blood sample	Semen sample	-	2. Sample		
А	В	С	D	VS	EIA-1	EVA-B1	EVA-S1	L CEM-11	CEM-12		
					EIA-2	EVA-B2	EVA-S2	2 CEM-21	CEM-22		
(1)	Delete as n	ecessary.									
(2)	document Regulation third count	apports of equine semen are authorised from a third country listed in column 2 as set out in a boument relating to 'equidae' published on gov.uk, in accordance with Commission Implementing egulation (EU) 2018/659 (11) provided that the semen was collected in the part of the territory of the ird country detailed in column 4 of that document from a donor stallion of the category of Equidae dicated in columns 11, 12 or 13 of that document.									
(3)	Only appro	oved semen	collection c	entres listed	l in accorda	nce with Ar	ticle 17(3	3)(b) of Directiv	e 92/65/EEC		
(4)	and import	ts into the C	ommunity o	of animals, s	semen, ova a	and embryo	s not sul	ements govern bject to animal I I) to Directive 9	health		
(5)					r 2009 on ar ries of Equid		n conditio	ons governing t	he		
(6)	Insert date in the table in point II.4.6 (follow guidance in Part II of the Notes).										
(7)	controls an	Regulation (EU) No 2017/625 of the European Parliament and of the Council of 15 March 2017 on official controls and other official activities performed to ensure the application of food and feed law, rules on animal health and animal welfare, plant health and plant protection products (Official Controls Regulation).									
(8)	not require Iceland has	The agar gel immunodiffusion test (AGID or Coggins test) or the ELISA for equine infectious anaemia are not required for donor equidae which have continuously resided in Iceland since birth, provided that Iceland has remained officially free of equine infectious anaemia and no equidae and their semen, ova and embryos have been introduced into Iceland from outside prior to and during the period the semen was collected									
(9)	Cross out th	he program	mes that do	not apply to	o the consig	nment.					
(10)	Insert nam	es and conc	entrations.								
 A the consent of the Scottish and Welsh Ministers, may be found here: document relating to 'equidae' for EU and EFTA states published by the Secretary of State, with 											
EU and EFT	A states ap	proved to ex	xport anima	ls and anim	al products	to Great Br	itain - da	ata.gov.uk			
		stamp must	be in a diff	erent coloui	r to that of tl	ne printing.					
Certifying Offi Name (in cap					Qualification	and title					