	I.1. Consignor					I.2. IMSOC Reference				
	Name					I.2.a. Local Reference				
	Address									
	Country ISO Code									
	I.5. Consignee					I.3. Central co	mpetent authority			
	Name						petent authority			
e l	Address									
Ĩ	Country			ISO Code						
Part I : Details of consignment	I.7. Country of origin ISO Code						fdestination		ISO Code	
ť	I.8. Region of origi	n			Code	I.10. Region of	destination			
S	I.11. Place of Dispa	itch				I.12. Place of d	lestination			
tai	Name					Name				
ลื่	Address					Address				
	Approval Number	ſ				Approval Nu	mber			
art	Country			ISO Code		Country ISO Code				
<u>-</u>	I.13. Place of Loadi	ing				I.14. Date and	time of departure			
	Name									
	Address	-								
	Approval Number Country	Ľ		ISO Code						
	-			130 code						
	I.15. Means of Tran	-				I.16 Entry Poin	nt			
	Mode	Internation transport	al	Identification						
		document								
-	I.18. Transport cor	ditions				I 17 Accompa	nying documents			
		Chilled 🗆		Ambient 🗆	Controlled		itying documents			
					temperature \Box	Accompanyi ng Date of issue document reference				
						Country Place of issue				
-	I.19. Container No	/ Seal No								
	I.20. Certified as									
	Artificial reproduc	tion \Box								
ľ	I.21. For transit thi	rough a third	d coun	try 🛛		I.22. For transit through Member State(s) □ Country ISO Code I.25. Total gross weight □				
	Country			ISO Code						
	EU Exit Authority			BCP code						
	EU Entry Authority			BCP code						
- F	I.24. Total quantity	7								
ŀ	I.28. Description of	f consignme	nt							
				I, NOT ELSEWHER	E SPECIFIED OR IN	CLUDED				
							or 3, unfit for human	consumption		
	051199 Other									
	05119985 (
	Commodity		Specie	s	Identification	number	Date of collection/production	Quantity		
ĺ										

	II. Health info	rmation									
	I, the unde hereby cer	-	icial veterin	arian, of the	e exporting c	ountry (2)	(nam	e of exporting country)			
	II.1	stored for	export to Gr	eat Britain	is approved a	semen described above was collected, processed and and supervised by the competent authority in accordance pter I(II)(1) of Annex D to Directive 92/65/EEC,					
Part II: Certification	II.2.	-	-	-	to the date of first collection of the semen described above nen elapsed, the semen collection centre:						
	II.2.1. was situated in the e to Article 13 of Direc exporting country w					n the exporting country or, in the case of regionalisation according f Directive 2009/156/EC, in that part of the territory of the ntry which was:					
Part II			-			fected with African ho f Directive 2009/156/E		mess in accordance with			
			-	free from '	Venezuelan e	equine encephalomyel	litis for	or 2 years,			
			-	free from g	glanders and	d dourine for 6 months;					
			II.2.2.		e conditions EC (8) and in	for a holding laid dow particular:	vn in Ar	ticle 4(5) of Directive			
		(1)	∘ either	[II.2.2.1.	species sus	case of a disease men ceptible to the disease l or killed and the hole	located	8			
				-	beginning o	vpe of equine encepha on the day on which th slaughtered,		tis for at least 6 months, lae suffering from the			
				-	obtain a ne test) carrie	gative result in an aga d out on samples takei l on two occasions 3 m	n gel im n after t	ast the period required to umunodiffusion test (Coggins he infected animals were upart from each of the			
				-	from vesicu case,	ılar stomatitis for at le	east 6 m	onths from the last recorded			
				-	from rabies	s for at least one mont	h from	the last recorded case,			
				-	from anthr	ax for at least 15 days	from th	e last recorded case,]			
		(1)	∘ or	[II.2.2.1.	species sus slaughtered been free fo encephalor and rabies which follo	ceptible to the disease d or killed and the pre- or at least 30 days fror nyelitis, equine infecti or 15 days in the case	located mises d n any ty ous ana of anth: of the an	below all the animals of on the holding have been isinfected, the holding has ope of equine temia, vesicular stomatitis rax, beginning on the day on himals the disinfection of the			
			II.2.3.		nly equidae gious equine		nical sig	ns of equine viral arteritis			
	II.3.	Prior to entering the semen collection centre the donor stallions and any other equidae located i centre:									
			II.3.1.	imported f country or	from Great B r, in the case EC (8), in that	ritain during the 3 mo of regionalisation acco	onths pe ording t	ntry if they were directly riod) in the exporting o Article 13 of Directive porting country which was			
			_			fected with African ho f Directive 2009/156/E		mess in accordance with			
			-	free from `	Venezuelan e	equine encephalomyel	litis for	at least 2 years,			
			-	free from g	glanders and	dourine for at least 6	months	;;			

(GB) Equine semen – Section B from EU countries GBHC047E (v3.0)

	ROPEAN U			(0)	B) Equine semen – Section B from EU countries GBHC047E (V3.0)							
	II. Health info	rmation										
	(1)	∘ either	[II.3.2.		from the country of export which was on the day of admission into free of vesicular stomatitis (VS) for at least 6 months,]							
	(1)	\circ or	[II.3.2.	out with n	were subjected to a virus neutralisation test for vesicular stomatitis (VS) carried out with negative result at a serum dilution of 1 in 12 on a blood sample taken (4) within 14 days prior to entering the centre;]							
Part II: Certification			II.3.3.	originated from holdings which on the day of admission onto the centre the requirements of point II.2.2;								
üĥ	II.4.	The semer	described a	above was c	ollected from donor stallions, which:							
II: Cer			II.4.1.		hown any clinical sign of an infectious or contagious disease at the mission onto the centre and on the day the semen was collected;							
Part			II.4.2.	have been kept for 30 days prior to the date of semen collection on he where no equine animal has shown any clinical sign of equine viral a contagious equine metritis during that period;								
			II.4.3.	first seme	een used for natural mating during at least 30 days prior to the date of n collection and between the dates of the first sample referred to in .5.1, II.4.5.2 and/or II.4.5.3 and until the end of the collection period;							
			II.4.4.	relevant C Animals of accordanc	ergone the following tests, which meet at least the requirements of the hapter of the Manual of Diagnostic Tests and Vaccines for Terrestrial f the WOAH (formerly OIE), carried out on samples taken in e with one of the programmes specified in point II.4.5 in a laboratory d by the competent authority:							
		(1)(5)	\circ either	[II.4.4.1.	an agar-gel immuno-diffusion test (Coggins test) for equine infectious anaemia (EIA) with negative result;]							
		(1)(5)	\circ or	[II.4.4.1.	an ELISA for equine infectious anaemia (EIA) with negative result;]							
	and (1)		\circ either	[II.4.4.2.	a serum neutralisation test for equine viral arteritis (EVA) with negative result at a serum dilution of one in four;]							
		(1)	∘ or	[II.4.4.2.	virus isolation test for equine viral arteritis (EVA) carried out with negative result on an aliquot of the entire semen of the donor stallion;]							
	and			II.4.4.3.	an agent identification test for contagious equine metritis (CEM) carried out on two occasions on samples collected with an interval of 7 days by isolation of Taylorella equigemtalis after a cultivation of 7 to 14 days from pre-ejaculatory fluid or a semen sample and from genital swabs taken at least from the penile sheath, urethra and urethral fossa with negative result in each case;							
			II.4.5.	one of the follows:	subjected with the results specified in II.4.4. in each case to at least test programmes (6) detailed in points II.4.5.1, II.4.5.2 and II.4.5.3 as							
				□ II.4.5.1.	The donor stallion was continuously resident on the semen collection centre for 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. have been carried out on samples taken(4) prior to the first semen collection and at least 14 days following the date of the commencement of the residence period of at least 30 days.							

Part II: Certification

_	ROPEAN UNION (GB	J Lyunie se	men – Se		countries GBHC047E (v3.0)		
	II. Health information						
11011	□ II.4.5.2.	The donor stallion was resident on the semen collection centre for at least 30 days prior to the date of the first collection and during the period of collection of the semen described above, but has left the centre under the responsibility of the centre veterinarian for a continuous period of less than 14 days, or other equidae on the collection centre came into direct contact with equidae of lower health status.					
		taken (4) pr breeding se above was o	ior to the o ason or co collected a	date of the first sen ellection period in t	been carried out on samples nen collection of the he year the semen described following the date of the of at least 30 days,		
7	and	last carried	out on a s	-	uine infectious anaemia was en (4) not more than 90 days ollected;		
	and	(1)	∘ either	equine viral arte sample taken (4)	described in point II.4.4.2. for ritis was last carried out on a not more than 30 days n described above was		
		(1)	° or	was carried out w aliquot of the ent stallion taken (4) before the semen collected and a b same date (4) rea neutralisation tes	test for equine viral arteritis with negative result on an tire semen of the donor not more than 6 months a described above was lood sample taken on the acted positive in a serum st for equine viral arteritis at of more than one in four,]		
	and	last carried	out on sar		ntagious equine metritis was t more than 60 days before d.		
	□ II.4.5.3.	taken (4) pr breeding se above was o carried out	ior to the o ason or co collected, a on sample	date of the first sen ellection period in t and the tests descri	been carried out on samples nen collection of the he year the semen described bed in point II.4.4 have been n 14 and 90 days after the		
	II.4.6. have undergone the testing provid following dates:	ed for in poi	nts 🗆 II.3	3.2.(1) and II.4.5. on	samples taken on the		

EUROPEAN UNION (GB) Equine semen – Section B from EU countries GBHC047E (v3.0) II. Health information Identificat Test Start Date of sampling for health tests(4) ion of programm date(4) semen е Donor Semen VS(1) II.3.2 EIA EVA CEM II.4.4.3. residence collection II.4.4.1. II.4.4.2. Part II: Certification Blood Semen 1. Sample 2. Sample sample sample (1) [II.5. No antibiotics were added to the semen;] \circ either (1) [II.5. The following antibiotic or combination of antibiotics was added to produce a \circ or concentration in the final diluted semen of not less than(7): ;] II.6 The semen described above was: II.6.1. collected, processed, stored and transported under conditions which comply with the requirements of Chapters II(I)(1) and III(I) of Annex D to Directive 92/65/EEC; II.6.2. sent to the place of loading in a sealed container in accordance with point 1.4 of Chapter III(I) of Annex D to Directive 92/65/EEC and bearing the number indicated in Box I.23.

	II. Health info	rmation								
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.								
Part	Box reference I.22:	The number of packages shall correspond to the number of containers.								
	Box reference I.23:	Dx The identification of container and seal number shall be indicated.								
	Box reference I.28:	The donor identity shall correspond to the official identification of the animal. ce								
	The date of	f collection shall be indicated in the following fo	ormat dd/mm/yyyy.							
	Part II:									
	Guidance f	Guidance for the completion of the table in point II.4.6.								
	Abbreviatio	ons:								
	VS	Vesicular stomatitis (VS) testing if required in accordance with point II.3.2								
	EIA-1	Equine infectious anaemia (EIA) testing first of	ccasion							
	EIA-2	EIA testing second occasion								
	EVA-B1	Equine viral arteritis (EVA) testing on blood sample first occasion								
	EVA-B2	EVA testing on blood sample second occasion								
	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 taken 7 days after CEM-21								
	Instruction	Instructions:								
	For each semen identified in column A in correspondence with Box I.28, the test programme (II.4.5.1., II.4.5.2. and/or II.4.5.3.) must be specified in column B, and columns C and D must 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 II.4.5.1., II.4.5.2. and II.4.5.3., are 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 II.4.5.2. or II.4.5.3. are entered in the lower line of columns 5 to 9 in table, this being the boxes ElA-2, EVA-B2 or EVA-S2 and CEM-21 and CEM-22 in the example below.									

	KOI LAN C			(61) Equine of				unities oblice	<u> </u>	
	II. Health infor	rmation									
	Identificat ion of semen	Test programm e	Start date		Date of sampling for health tests						
			Donor residence	Semen collection	VS 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	
rtifi	А	В	С	D	VS	EIA-1	EVA-B1	EVA-S1	CEM-11	CEM-12	
I: Ce	(1)	Delete ee n	00000771			EIA-2	EVA-B2	EVA-S2	CEM-21	CEM-22	
rt II	(1)	Delete as n	-	an ana auth	aniand frame	a third cour	ature liata din		2 ag agt gut in		
Pa	(2)	Imports of equine semen are authorised from a third country listed in column 2 as set out in a document relating to 'equidae' published on gov.uk, in accordance with Commission Implementing Regulation (EU) 2018/659 (8) provided the semen was collected in the part of the territory of the third country detailed in column 4 of that document from a donor stallion of the category of equidae indicated in columns 11, 12 or 13 of that document.									
	(3)	Only approved semen collection centres listed in accordance with Article 17(3)(b) of Council Directive 92/65/EEC.									
	(4)	Insert date	in table in J	ooint II.4.6 (follow Guida	ance in Part	II of the No	tes).			
(5) The agar gel immunodiffusion test (Coggins test) or the ELIS required for donor equidae which have continuously reside Iceland has remained officially free of equine infectious ana and embryos have been introduced into Iceland from outsid was collected.						ded in Icelai naemia and	nd since l no equio	birth, provided lae and their s	that emen, ova		
	(6)	Cross out th	he program	mes that do	not apply to	the consign	nment.				
	(7)	Insert names and concentrations.									
	(8)		nt relating to the Scottish	-		-	-	the Secr	etary of State, v	with the	
	EU and EFT	A states ap	proved to ex	xport anima	ls and anim	al products	to Great Bri	tain - dat	a.gov.uk		
		The signature and the stamp must be in a different colour to that of the printing. ertifying Officer									
	Name (in cap					Qualification	and title				
	Date of signa					Signature					
	Stamp										