	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		
en	Address					* ,		
gnm	Country		ISO Code					
Part I : Details of consignment	I.7. Country of origin			ISO Code	I.9. Country of	f destination		ISO Code
of (I.8. Region of origin			Code	I.10. Region of	f destination		
ls (I.11. Place of Dispatch				I.12. Place of o	lestination		
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					
	-				I 10 Frature Dat			
	I.15. Means of Transpo Mode		Identification		I.16 Entry Poi	nt		
	trai	ernational nsport	Identification					
		rument						
	I.18. Transport condition	0.005			I.17. Accompanying documents			
		lled 🗆	Ambient 🗆	Controlled	Accompanyi	inying documents		
	temperature				ng Date of issue			
					reference			
					Country		Place of issue	
	I.19. Container No / Sea	al No			1			
	I.20. Certified as							
	Artificial reproduction		Breeding 🗆					
	I.21. For transit through a third country					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 cor	nsignment						
	1.05 PRODUCTS OF A	NIMAL ORIGIN	I, NOT ELSEWHER	E SPECIFIED OR IN	CLUDED			
	-	cts not elsewh	ere specified or inc	luded; dead anima	ls of Chapter 1	or 3, unfit for human	consumption	
	051199 Other 05119985 Othe	7						
				T-land: Gaadian		Deta of	Quantita	
	Commodity	Specie	25	Identification	number	Date of collection/production	Quantity	

	II. Health information								
	I, the undersigned, official veterinarian, of the exporting country(2) (name of exporting country) hereby certify that:								
	II.1.	The \circ ova(1) / \circ embryos(1) described above:							
ation		II.1.1.	were \circ collected(1) / \circ produced(1) by the team(3) described in Box I.11, which has been approved and supervised in accordance with Chapter I(III) of Annex D to Directive 92/65/EEC and is subject to inspection by an official veterinarian at least once every calendar year;						
Part II: Certification		II.1.2.		ollected(1) / \circ produced(1) , processed and stored in accordance with rements of Chapter III(II) of Annex D to Directive 92/65/EEC;					
Li Li		II.1.3.	were collected at a place separated from other parts of the premises or holdi which is in good repair and was cleaned and disinfected prior to the collection						
Pa		II.1.4.	were examined, processed and packed in laboratory facilities which are not situated in a zone subject to prohibition or quarantine measures as set out in Box II.1.6, in a section which is separated from the section for storing equipmen and materials used in contact with donor animals and from the area where the donor animals are handled;						
		II.1.5.	come from donor mares which:						
			II.1.5.1.	were continuously resident for three months (or since entry if they were directly imported from Great Britain during the three months period) in the exporting country or, in the case of regionalisation according to Article 13 of Directive 2009/156/EC(8), in that part of the territory of the exporting country which was during that period					
			-	not considered to be infected with African horse sickness in accordance with Article 5(2)(a) and (b) of Directive 2009/156/EC,					
			-	free from Venezuelan equine encephalomyelitis for at least 2 years,					
			-	free from glanders and dourine for at least 6 months;					
	(1)	∘ either	[II.1.5.2.	originated from a country of export which was on the day of collection free of vesicular stomatitis for at least 6 months;]					
	(1)	o or	[II.1.5.2.	were tested by a virus neutralisation test for vesicular stomatitis on a blood sample taken on (4) within 30 days prior to collection, with negative result at a serum dilution of 1 in 12;]					
	(1)	∘ either	[II.1.5.3.	during the past 30 days prior to collection have been located in holdings under veterinary supervision which fulfilled from the day of collection of \circ ova(1) / \circ embryos(1) until the date of their dispatch the conditions for a holding laid down in Article 4(5) of Directive 2009/156/EC, and in particular:]					
	(1)	∘ or	[II.1.5.3.	during the past 30 days prior to collection have been located in holdings under veterinary supervision which fulfilled from the day of collection of \circ ova(1) / \circ embryos(1) until, in the case of frozen \circ ova(1) / \circ embryos(1), the period of 30 days mandatory storage at approved premises elapsed, the conditions for a holding laid down ir Article 4(5) of Directive 2009/156/EC and in particular:]					
		(1)	○ either	[II.1.5.3.1. following a case of a disease mentioned below not all the animals of species susceptible to the disease located on the holding were slaughtered or killed and the holding has been free:					
				- from any type of equine encephalomyelitis for at least 6 months, beginning on the day on which the equidae suffering from the disease are slaughtered,					

	IROPEAN UNION				(v3.0)	
	II. Health information					
ц				-	from equine infectious anaemia for at least the period required to obtain a negative result in an agar gel immunodiffusion test (Coggins tests) carried out on samples taken after the infected animals were slaughtered on two occasions three months apart from each of the remaining equidae;	
Part II: Certification		-		-	from vesicular stomatitis for at least 6 months from the last recorded case,	
Certif				-	from rabies for at least one month from the last recorded case,	
art II:				-	from anthrax for at least 15 days from the last recorded case,]	
4		(1)	° or	[II.1.5.3.1.	following a case of a disease mentioned below all the animals of species susceptible to the disease located in the holding have been slaughtered or killed and the premises disinfected, the holding has been free for at least 30 days from any type of equine encephalomyelitis, equine infectious anaemia, vesicular stomatitis and rabies or 15 days in the case of anthrax, beginning on the day on which following the destruction of the animals the disinfection of the premises was satisfactorily completed;]	
			II.1.5.4.	each of the	e past 30 days prior to collection have been kept in holdings em having been free from clinical signs of contagious etritis for at least 60 days;	
			II.1.5.5.	the date of first samp	used for natural breeding during at least 30 days prior to f collection of ova or embryos and between the date of the les referred to in points II.1.6.6 and II.1.6.7 and the date of ion of ova and embryos;	
			II.1.5.6.	diffusion t anaemia c being duri of ova or e blood take	subjected with negative result to an agar-gel immuno- est (Coggins test) or an ELISA for equine infectious arried out on a blood sample taken on (4), ing the past 30 days prior to the date of the first collection embryos and the test was last carried out on a sample of en on (4), being not more than 90 days before embryos were collected(5);	
			II.1.5.7.	equine me cultivatior case on sa first collec clitoral fos on culture sp	subjected to an agent identification test for contagious etritis by isolation of Taylorella equigenitalis after a n of 7 to 14 days carried out with negative results in each mples taken during the past 30 days prior to the date of the etion of ova or embryos from mucosal surfaces of the sea and clitoral sinuses on two consecutives oestrus periods (4) and on (4), and on an additional ecimen taken during one of the oestrus periods from the tal cervix on (4);	
			II.1.5.8.	been in co	of my knowledge and as far as I can ascertain, have not ntact with equidae suffering from an infectious or s disease during the 15 days immediately preceding the	
			II.1.5.9.		he day of collection of \circ ova(1) / \circ embryos(1) not shown gns of an infectious or contagious disease;	
		II.1.6. were \circ collected(1) / \circ produced(1) after the date on which the embryo collection(1) / \circ production(1) team described in Box I.11 was approved competent authority of the exporting country;		action(1) team described in Box I.11 was approved by the		

II. Health information				
	II.1.7.	were processed and sto	bred under approved conditio	ns for at least 30 days
		immediately after their	$r \circ collection(1) / \circ production(1)$	on(1) , and were transported
		under conditions whic	h satisfy the terms laid down :	in Chapter III(II) of Annex D

to Directive 92/65/EEC;

- (9)II.2. The embryos described above were conceived by artificial insemination(1) / as a result of in vitro fertilisation(1) using semen meeting the requirements of Directive 92/65/EEC and coming from semen collection centres approved in accordance with Article 11(2) or 17(3)(b) of Directive 92/65/EEC and located respectively in Great Britain or in a third country or parts of the territory of third country listed in columns 2 and 4 in a document relating to 'equidae' published on gov.uk, in accordance with Commission Implementing Regulation (EU) 2018/659 (9) from which the import of equine semen collected from registered horses, registered equidae or equidae for breeding and production is authorised in accordance with Article 4 of Commission Implementing Regulation (EU) 2018/659 and indicated in columns 11,12 and 13 of that document.(6)(7);
- □ II.3. The ova ova used for in vitro production of the embryos described above comply with the requirements of Annex D to Directive 92/65/EEC and in particular the requirements set up in points II.1.1 to II.1.8 of this certificate (1).

Part II: Certification

	II. Health info	rmation								
-										
	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).									
catior	References Part I:	to Great Bri	itain in this certificate include Chan	nel Islands and Isle of Man.						
<u>Part II: Certification</u>	rarri.	Box reference I.11:	accordance with Article 17(3)(b) of Council Directive 92/65/EEC.							
Part		Box reference I.19:	The identification of container and seal number shall be indicated.							
		Box reference I.28:	The number of packages shall correspond to the number of containers.							
		Box reference I.28:	The category: specify if in vivo der embryos or micromanipulated em		ova, in vitro produced					
		The donor	identity shall correspond to the offi	cial identification of the animal.						
		The date of	collection shall be indicated in the							
	Part II:				-					
	(1)	Delete as a	ppropriate.							
 Only third countries or parts of the territory of third countries listed in columns 2 and 4 ir relating to 'equidae' published on gov.uk, in accordance with Commission Implementing I (EU) 2018/659(9) respectively from which imports of registered equidae and equidae for b production are also authorised and as indicated in column 14 of that document. 										
	 (3) Only approved embryo collection teams and embryo production teams listed in accordance with Au 17(3)(b) of Council Directive 92/65/EEC. 									
	(4)	Insert date								
	(5) The agar gel immunodiffusion test (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, or and embryos have been introduced into Iceland from outside prior to and during the period the sem was collected.									
	(6)		ved semen collection centres listed rective 92/65/EEC.	in accordance with Article 11(4) or Article 17(3)(b) of						
	(7)	Does not ap	oply to ova.							
(8) Council Directive 2009/156/EC of 30 November 2009 on animal health conditions governing the movement and importation from third countries of equidae.										
	(9)	Athe consent of the Scottish and Welsh Ministers, may be found here:documentrelating to'equidae'for EU andEFTAstatespublishedby theSecretaryof State,with								

	II. Health information						
	EU and EFTA states approved to export animals and anima	al products to Great Britain - d	lata.gov.uk				
	The signature and the stamp must be in a different colour to that of the printing.						
	Certifying Officer						
lon	Name (in capital letters) Date of signature Stamp	Qualification and title Signature					
cati							
Į							
G							
Part II: Certification							
Par							
_							