ANIMAL HEALTH CERTIFICATE FOR THE MOVEMENT BETWEEN MEMBER STATES OF CONSIGNMENTS OF STOCKS OF OOCYTES AND EMBRYOS OF EQUINE ANIMALS COLLECTED OR PRODUCED, PROCESSED AND STORED IN ACCORDANCE WITH DIRECTIVE 92/65/EEC BEFORE 1 SEPTEMBER 2010, DISPATCHED AFTER 20 APRIL 2021 BY THE EMBRYO COLLECTION OR PRODUCTION TEAM BY WHICH THE OOCYTES OR EMBRYOS WERE COLLECTED OR PRODUCED

URO	PEAN UN	ION				INTRA		
	I.1	Consignor		I.2	IMSOC reference			
		Name		I.2a	Local reference			
		Address		I.3	Central Competent Author	ority QR CODE		
I		Country	ISO country code	I.4	Local Competent Authori	ty		
Part I: Description of consignment	I.5	Consignee		I.6	Operator conducting assembly operations independently of an establishment			
		Name			Name	Registration No		
		Address			Address			
		Country	ISO country code		Country	ISO country code		
	I.7	Country of origin	ISO country code	1.9	Country of destination	ISO country code		
222	1.8	Region of origin	Code	I.10	Region of destination	Code		
	I.11	Place of dispatch		I.12	Place of destination			
art		Name	Registration/Approval No		Name	Registration/Approval No		
-		Address			Address			
		Country	ISO country code		Country	ISO country code		
	I.13	Place of loading			Date and time of departur	e		
	I.15	Means of transport		I.16	Transporter			
		Vessel	□ Aircraft		Name	Registration/Authorisation No		
					Address	NO		
		Railway Road vehicle			Country	ISO country code		
					Accompanying documents			
		Identification	□ Other		Туре	Code		
		Document			Country	ISO country code		
		Document		Commercial document reference				
	I.18	Transport condition	s 🗆 Ambient		Chilled	Frozen		
	I.19	Container number/S	Seal number					
		Container No	S	Seal No				

(MODEL 'EQUI-OOCYTES- EMB-D-INTRA')

I.20	Certified as or f	or							
□ Further k	eeping	□ Slaughter	Slaughter		Confined establishment		Germinal products		
□ Registere	ed equine animal	Travelling cir	Travelling circus/animal act		ibition	□ Event or activity near borders			
□ Release i	nto the wild	Dispatch cent	□ Dispatch centre		aying area/purification	Ornamental aquaculture establishment		ulture	
□ Further p	rocessing	 Organic fertilities improvers 	 Organic fertilizers and soil improvers 		hnical use	 Quarantine or similar establishment 			
□ Products	for human consump	tion	□ Pollination		□ Live aquatic animals for		Other		
				humai	n consumption				
I.21	🗆 For transit t	hrough a third country	y						
	Third country			ISO country code					
	Exit point		BCP code						
	Entry point			E	BCP code				
I.22 🗆	For transit throug	h Member State(s)		I.23	□ For export				
Ν	Iember State	ISO	ISO country code		Third country	ISO country code			
Ν	fember State	ISO	ISO country code		Exit point	BCP code			
Ν	Member State ISO country code								
I.24 E	.24 Estimated journey time			I.25	Journey log	□ yes		□ no	
I.26 T	I.26 Total number of packages				Total quantity				
I.28 T	otal net weight/gro	ss weight (kg)		I.29	29 Total space foreseen for the consignment				
I.30 D	escription of consig	gnment							
CN code	Species	Subspecies/Category		tification	Identification	number	Age	Quantity	
			syst					Туре	
Region of origin Cold		Cold store	Ider	tification	mark Type of packa	aging		Net weight	
Slaughterhouse T		Treatment type		Nature of commodity		Number of packages		Batch No	
		Date of collection/production	Mar plan	ufacturin t	g Approval or r number of plant/establish	0	Test		

ROPEAN UN			Certificate model EQUI-OOCYTES-EMB-D-INTRA					
II. Hea	II. Health information			Certificate reference	II.b	IMSOC reference		
I, the	I, the undersigned official veterinarian, hereby certify that:							
II.1.	II.1. Ova/embryos ⁽¹⁾ described in Part I were collected by a collection team ⁽²⁾ approved by the co authority and processed in an appropriate laboratory;							
II.2.	Ova/embryos ⁽¹⁾ were collected from donor mares which:							
	II.2.1.	II.2.1. on the day of collection have been located in premises situated on the territory or in the cas regionalisation in a part of the territory of a Member State which is not considered to be infer with African horse sickness in accordance with Article 5(2)(a) and (b) of Direc 2009/156/EC ⁽³⁾ ,						
II.2.2. have been located in holdings under veterinary supervision which on the day of fulfilled the conditions of Article 4 of Directive 2009/156/EC,								
	II.2.3. have been kept prior to the collection in holdings free from clinical signs of contagious equi metritis for 60 days,							
	II.2.4. have not been used for natural breeding during the period of 30 days prior to the collectio ova/embryos ⁽¹⁾ ,							
	II.2.5.	to the best of my knowledge and as far a suffering from an infectious or contagio collection of ova/embryos ⁽¹⁾ ,				*		
II.2.6. have on the day of collection not shown clinical signs of an infectious or contagious					ntagious disease;			
II.3.	Ova/embryos ⁽¹⁾ were collected, processed, stored and transported und requirements of Annex D of Directive 92/65/EEC;				litions v	which comply with the		
II.4.	The semen used for the artificial insemination of the donor mares complies with the requirements o Directive $92/65/EEC^{(4)(1)}$;							
II.5.	The ova used for the <i>in vitro</i> production of embryos comply with the requirements of Directive $92/65/\text{EEC}^{(1)}$.							
Notes								
for in	This animal health certificate shall be completed according to the notes for the completion of certificates provided for in Chapter 2 of Annex I to Commission Implementing Regulation (EU) 2020/2235.							
Part I Box I.		Place of dispatch shall correspond to the en	brvo	collection team of ove	embry	os collection		
Box I.	12: P	Place of dispatch shall correspond to the embryo collection team of ova/embryos collection. Place of destination shall correspond to the embryo collection team, embryo production team or to the holding of ova/embryos destination.						
Box I.		Identification of container and Seal number shall be indicated.						
Box I.		"Type": Specify if: <i>in vivo</i> derived embryos, <i>in vivo</i> derived oocytes, <i>in vitro</i> produced embryos o micromanipulated embryos.						
	Γ	Donor identity shall correspond to the offic	ial ide	ntification of the anin	nal.			
		Date of collection shall be indicated in the						
		approval number of the team shall correctly ollection.	espon	d to the embryo coll	ection	team of ova/embryo		

EUROPEAN UNION

Certificate model EQUI-OOCYTES-EMB-D-INTRA

Part II:						
⁽¹⁾ Delete as appropriate.						
⁽²⁾ Only embryo collection teams approved by the competent authority and listed in accordance with Article 11(4) of Council Directive 92/65/EEC.						
⁽³⁾ OJ L 192, 23.7.2010, p. 1.						
⁽⁴⁾ Does not apply to ova.						
Official veterinarian						
Name (in capital letters)		Qualification and title				
Local Control Unit name		Local Control Unit code				
Date						
Stamp		Signature				