ANIMAL HEALTH CERTIFICATE FOR THE MOVEMENT BETWEEN MEMBER STATES OF CONSIGNMENTS OF OOCYTES AND EMBRYOS OF EQUINE ANIMALS COLLECTED OR PRODUCED, PROCESSED AND STORED IN ACCORDANCE WITH REGULATION (EU) 2016/429 AND DELEGATED REGULATION (EU) 2020/686 AFTER 20 APRIL 2021, DISPATCHED BY THE EMBRYO COLLECTION OR PRODUCTION TEAM BY WHICH THE OOCYTES OR EMBRYOS WERE COLLECTED OR PRODUCED

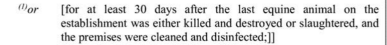
(MODEL 'EQUI-OOCYTES-EMB-A-INTRA')

JRO	OPEAN U	NION				INTR				
	I.1 Consignor				IMSOC reference					
		Name		I.2a	Local reference					
		Address		I.3	Central Competent Authority	QR CODE				
		Country	ISO country code	I.4	Local Competent Authority					
	I.5	Consignee		1.6	Operator conducting assembly operations independently of ar establishment					
0		Name			establishment Name	Registration No				
		Address			Address	Ü				
		Country	ISO country code		Country	ISO country code				
	I.7	Country of origin	ISO country code	1.9	Country of destination	ISO country code				
	I.8	Region of origin	Code	I.10	Region of destination	Code				
	I.11	Place of dispatch		I.12	Place of destination					
		Name	Registration/Approval No		Name	Registration/Approval No				
١		Address			Address					
		Country	ISO country code		Country	ISO country code				
Ì	I.13	Place of loading		I.14	Date and time of departure					
	I.15	Means of transport		I.16	Transporter					
		□ Vessel	□ Aircraft		Name	Registration/Authorisation No				
					Address					
		□ Railway	□ Road vehicle		Country	ISO country code				
				I.17	Accompanying documents					
		Identification	□ Other		Type	Code				
		Document			Country	ISO country code				
					Commercial document reference					
	I.18	Transport condition	ns Ambient		□ Chilled	□ Frozen				
	I.19	Container number/	Seal number							
		Container No	S	eal No						

I.20 Certified as or for										
□ Further keeping	□ Slaughter		□ Confi	ned establishment	□ Germinal	products				
□ Registered equine animal	□ Travelling circus/animal act		□ Exhib	ition	□ Event or activity near borders					
□ Release into the wild	□ Dispatch centre	□ Relaying area/pu centre		ing area/purification	☐ Ornamental aquaculture establishment		ılture			
□ Further processing	□ Organic fertilizers and so	s and soil		ical use	□ Quarantir	lar				
	improvers				establishme	ent				
$\hfill\Box$ Products for human consumption	□ Pollination		□ Live a	quatic animals for	\Box Other					
			human c	consumption						
I.21	gh a third country									
Third country			ISC	country code						
Exit point			BC	P code						
Entry point			ВС	P code						
I.22	mber State(s)	I	.23	For export						
Member State	ISO country cod	de		Third country	ISC	O country	code			
Member State	ISO country cod	de	Exit point BCP code							
Member State	ISO country cod	ie								
I.24 Estimated journey time		I	.25	Journey log	□ yes		□ no			
I.26 Total number of packages		I	.27	Total quantity						
I.28 Total net weight/gross weight	Total net weight/gross weight (kg)				I.29 Total space foreseen for the consignment					
I.30 Description of consignment										
CN code Species Subs		Identifi		Identification	number	Age	Quantity			
		system					Type			
Region of origin Cold	store	Identification mark		nark Type of pack	Type of packaging		Net weight			
Slaughterhouse Treat	3.1	Nature		Number of pa	Number of packages		Batch No			
Date colled		Manufa plant	acturing	Approval or r number of plant/establis		Test				

	II. Health informa	tion			II.a	Certificate referen	ice	II.b	IMSOC reference		
	I, the undersigned official veterinarian, hereby certify that:										
	(⁽⁾⁾ [II.1.	The <i>in vivo</i> derived embryos of equine animals described in Part I have been collected, processed and stored, and dispatched by the embryo collection team ⁽²⁾ which									
		II.1.1.	is approve	is approved and kept in a register by the competent authority;							
		II.1.2.	facilities	es with requirements as regards responsibilities, operational procedures, es and equipment set out in Part 2 of Annex I to Commission Delegated ation (EU) 2020/686.]							
	(*)[II.1.	described	e oocytes ⁽¹⁾ / <i>in vitro</i> produced embryos ⁽¹⁾ / micromanipulated embryos ⁽¹⁾ of equine animals scribed in Part I have been collected or produced, processed and stored, and dispatched by the abryo production team ⁽²⁾ which								
		II.1.1.	is approved and kept in a register by the competent authority;								
		II.1.2.	complies with requirements as regards responsibilities, operational procedures, facilities and equipment set out in Parts 2 and 3 of Annex I to Delegated Regulation (EU) 2020/686.]								
Part II: Certification	II.2.	2. The oocytes ⁽¹⁾ / embryos ⁽¹⁾ described in Part I are intended for artificial reproduction obtained from donor animals which							production and were		
: Certil		II.2.1.		een born and remained since birth in the Union, or have entered the Union in lance with the requirements for entry into the Union;							
Part II		II.2.2.		tablishments in a Member State or zone thereof, or from establishments control by the competent authority in a third country or territory, or a							
			II.2.2.1.		ceding	30 days prior to			reported during the // production ⁽¹⁾ of the		
			⁽¹⁾ either						ring the period of the (1) of the oocytes(1)/		
			⁽¹⁾ or	preceding 2 year	rs prio llowir	or to collection ⁽¹⁾ ng the last outbre)/ pro	duction	ng the period of the (1) of the oocytes(1)/shment has remained		
				subjected provided Regulati samples	d to a l for i on (El taken	test for surra win Part 3 of An U) 2020/688, car	ith on nex I ried o s after	to Co out, with	olishment have been e diagnostic methods ommission Delegated in negative results, on ti infected animal has		

	(1) or [for at least 30 days from the date of cleaning and disinfection after the last animal of listed species on the establishment was either killed and destroyed or slaughtered;]]
II.2.2.2.	in which dourine has not been reported during the period of the preceding 6 months prior to $collection^{(1)}$ / production ⁽¹⁾ of the $oocytes^{(1)}$ / embryos ⁽¹⁾ , and
⁽¹⁾ either	[dourine has not been reported in the establishment during the period of the preceding 2 years prior to collection (1)/ production (1) of the oocytes (1)/ embryos (1);]
(l) _{OF}	[dourine has been reported in the establishment during the period of the preceding 2 years prior to collection (1)/ production (1) of the oocytes (1)/ embryos (1) and following the last outbreak, the establishment has remained under movement restrictions
	(1) either [until the remaining equine animals in the establishment, except castrated male equine animals have been subjected to a test for dourine with the diagnostic method provided for in Part 8 of Annex I to Delegated Regulation (EU) 2020/688, carried out, with negative results, on samples taken at least 6 months after the infected animals have been killed and destroyed or slaughtered, or the infected entire male equine animals have been castrated;]]
	(1) or [for at least 30 days after the last equine animal on the establishment was either killed and destroyed or slaughtered, and the premises were cleaned and disinfected;]]
II.2.2.3.	in which equine infectious anaemia has not been reported during the period of the preceding 90 days prior to collection $^{(1)}/$ production $^{(1)}$ of the oocytes $^{(1)}/$ embryos $^{(1)},$ and
⁽¹⁾ either	[equine infectious anaemia has not been reported on the establishment during the period of the preceding 12 months prior to collection ⁽¹⁾ / production ⁽¹⁾ of the oocytes ⁽¹⁾ / embryos ⁽¹⁾ ;]
(1) _{or}	[equine infectious anaemia has been reported on the establishment during the period of the preceding 12 months prior to collection ⁽¹⁾ / production ⁽¹⁾ of the oocytes ⁽¹⁾ / embryos ⁽¹⁾ and following the last outbreak the establishment has remained under movement restrictions
	(1) either [until the remaining equine animals in the establishment have been subjected to a test for equine infectious anaemia with the diagnostic method provided for in Part 9 of Annex I to Delegated Regulation (EU) 2020/688, carried out, with negative results, on samples taken on two occasions with a minimum interval of 3 months after the infected animals have been killed and destroyed or slaughtered and the establishment was cleaned and disinfected;]]



- II.2.3. were examined by the team veterinarian or a team member and did not show symptoms or clinical signs of transmissible animal diseases on the day of collection of the oocytes⁽¹⁾/ embryos⁽¹⁾;
- II.2.4. are identified as provided for in Article 58(1), 59(1) or 62(1) of Commission Delegated Regulation (EU) 2019/2035;
- II.2.5. for a period of at least 30 days prior to the date of first collection of the oocytes⁽¹⁾/ embryos⁽¹⁾ and during the collection period
 - II.2.5.1. were kept on establishments not situated in a restricted zone established due to the occurrence of African horse sickness, infection with Burkholderia mallei (glanders) or of an emerging disease relevant for equine animals:
 - ▶"II.2.5.2. were kept on establishments where Venezuelan equine encephalomyelitis, dourine, surra (*Trypanosoma evansi*), equine infections anaemia, contagious equine metritis (*Taylorella equigenitalis*), infection with rabies virus and anthrax have not been reported; ◀
 - II.2.5.3. were not in contact with animals from establishments situated in a restricted zone due to the occurrence of diseases referred to in point II.2.5.1. or from establishments which do not meet the conditions referred to in point II.2.5.2.;
- II.2.6. were not used for natural breeding during a period of at least 30 days prior to the date of the collection of the oocytes⁽¹⁾/ embryos⁽¹⁾ and between the date of the first samples referred to in points II.2.7.1. and II.2.7.2. and the date of the collection of the oocytes⁽¹⁾/ embryos⁽¹⁾;
- II.2.7. have been subjected to the following tests, referred to in points 2(b) and (c) of Chapter II of Part 4 of Annex II to Delegated Regulation (EU) 2020/686, as follows:

 - II.2.7.2. for contagious equine metritis (CEM), an agent identification test carried out with a negative result on at least two specimens (swabs) taken during the period referred to in point II.2.6. from at least the mucosal surfaces of the clitoral fossa and the clitoral sinuses of the donor mare

The samples referred to in points II.2.7.2.1. and II.2.7.2.2. were in no case taken earlier than 7 days (systemic treatment) or 21 days (local treatment) after antimicrobial treatment of the donor mare and were placed in transport medium with activated charcoal, such as Amies medium, before dispatch to the laboratory.

- II.3. The oocytes⁽¹⁾/ embryos⁽¹⁾ described in Part I
 - II.3.1. has been collected, processed and stored in accordance with animal health requirements set out in Part 2⁽¹⁾/Part 3⁽¹⁾/Part 4⁽¹⁾/Part 5⁽¹⁾ and Part 6 of Annex III to Delegated Regulation (EU) 2020/686;
 - II.3.2. are placed in straws or other packages on which the mark is applied in accordance with requirements provided for in Article 10 of Delegated Regulation (EU) 2020/686 and that mark is indicated in Box I.30;
 - II.3.3. are transported in a container which:
 - II.3.3.1. was sealed and numbered prior to the dispatch by the embryo collection or production team under responsibility of the team veterinarian, or by an official veterinarian, and the seal bears the number as indicated in Box I.19;
 - II.3.3.2. has been cleaned and either disinfected or sterilised before use, or is single-use container:
 - (1)(4)[II.3.3.3. has been filled in with the cryogenic agent which not have been previously used for other products;]
 - (1)(5)[II.3.4. are placed in straws or other packages which are securely and hermetically sealed;
 - II.3.5. are transported in a container where they are separated from each other by physical compartments or by being placed in secondary protective bags.]

(1)(6)[II.4. The *in vivo* derived embryos⁽¹⁾/ *in vitro* produced embryos⁽¹⁾/ micromanipulated embryos⁽¹⁾ described in Part I were conceived by artificial insemination using semen coming from a semen collection centre, germinal product processing establishment or germinal product storage centre approved for the collection, processing and/or storage of semen by the competent authority of a Member State or by the competent authority of a third country, territory or zone thereof listed in Annex XII to Commission Implementing Regulation (EU) 2021/404.]

Notes

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 reference I.11: "Place of dispatch": Indicate the unique approval number and the name and address of

the embryo collection or production team of dispatch of the consignment of oocytes or

embryos.

Box reference I.12: "Place of destination": Indicate the address and unique registration or approval

number of the establishment of destination of the consignment of oocytes or embryos.

Box reference I.19: Seal number shall be indicated.

Box reference I.26: Total number of packages shall correspond to the number of containers.

Box reference I.30: "Type": Specify if in vivo derived embryos, in vivo derived oocytes, in vitro produced

embryos or micromanipulated embryos.

"Identification number": Indicate identification number of each donor animal.

 ${\it ``Identification mark''}: Indicate mark on the straw or other packages where oocytes or$

embryos of the consignment are placed.

"Date of collection/production": indicate the date on which oocytes or embryos of the

consignment was collected or produced.

"Approval or registration number of plant/establishment/centre": Indicate the unique

approval number of the embryo collection or production team by which the oocytes or embryos were collected or produced.

"Quantity": Indicate number of straws or other packages with the same mark.

Certificate model EQUI-OOCYTES-EMB-A-INTRA

Part II:

- (1) Delete if not applicable.
- Only embryo collection or production teams approved by the competent authority and included in the register referred to in Article 101(1)(b) of Regulation (EU) 2016/429 and Article 7 of Delegated Regulation (EU) 2020/686.
- (3) Insert date in the following format: dd.mm.yyyy.
- (4) Applicable for frozen oocytes or embryos. (5) Applicable for the consignment where in one container oocytes, *in vivo* derived embryos, *in vitro* produced embryos and micromanipulated embryos of equine animals are placed and transported.
- (6) Does not apply to oocytes.
- (7) Mandatory attestation in case antibiotics were added.
- (8) Insert the name(s) of the antibiotic(s) added and its(their) concentration.

` ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' '	
Official veterinarian	
Name (in capital letters)	Qualification and title
Local Control Unit name	Local Control Unit code
Date	
Stamp	Signature