ANIMAL HEALTH CERTIFICATE FOR THE MOVEMENT BETWEEN MEMBER STATES OF CONSIGNMENTS OF OOCYTES AND EMBRYOS OF PORCINE 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

EUROF	PEAN UNIO	N				INTRA		
	I.1	Consignor		I.2	IMSOC reference			
		Name		I.2a	Local reference			
		Address		1.3	Central Competent Authority	QR CODE		
ent		Country	ISO country code	I.4 I.6	Local Competent Authority			
шш	1.5	Consignee			Operator conducting assembly operations independently o an establishment			
Isig		Name			Name	Registration No		
f cor		Address			Address			
Part I: Description of consignment		Country	ISO country code		Country	ISO country code		
crip	I.7	Country of origin ISO country code		1.9	Country of destination	ISO country code		
Des	I.8	Region of origin Code		I.10	Region of destination	Code		
Ξ	I.11	Place of dispatch		I.12	Place of destination			
art		Name	Registration/Approval No		Name	Registration/Approval No		
H		Address			Address			
		Country	ISO country code		Country	ISO country code		
	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	IND		
	Railway		Road vehicle		Country	ISO country code		
				I.17	Accompanying documents			
		Identification	□ Other		Туре	Code		
		Demment			Country	ISO country code		
		Document			Commercial document reference			
	I.18	Transport conditions	Ambient		Chilled	Frozen		
	I.19	Container number/Se	al number					
		Container No	S	eal No				

## (MODEL 'POR-OOCYTES-EMB-A-INTRA')

I.20	Certified as or	for								
🗆 Furt	her keeping	□ Slaughter			Confined establishment		Germinal products			
🗆 Regi	istered equine animal	□ Travelling cir	Travelling circus/animal act			oition		□ Event or	activity n	ear borders
🗆 Rele	ease into the wild	□ Dispatch cent	□ Dispatch centre		Relaying area/purification centre		<ul> <li>Ornamental aquaculture</li> <li>establishment</li> </ul>		ulture	
□ Furt	her processing	Organic fertilizers and soil			Technical use		Quarantine or similar		lar	
		improvers	improvers					establishment		
🗆 Prod	lucts for human	Pollination	Pollination			aquatic	animals for	$\Box$ Other		
consu	mption				human	consum	ption			
I.21	□ For transit	through a third country	y							
I	Third country				ISC	O count	ry code			
	Exit point				BCP code					
	Entry point				BC	CP code				
I.22	For transit through	gh Member State(s)			I.23 🗆	For e	kport			
	Member State	ISC	country o	code		Third	l country	IS	SO countr	y code
	Member State	ISC	ISO country code			Exit point		BCP code		
Member State ISO country code			code							
I.24	.24 Estimated journey time				I.25	Jour	ney log	□ yes		□ no
I.26	Total number of packages				I.27	Tota	l quantity			
I.28	Total net weight/gross weight (kg)				I.29	Tota	l space foreseer	n for the con	signment	
I.30	Description of consi	gnment								
CN co	de Species	Subspecies/Category	Sex	Ident syster	ification		Identification n	umber	Age	Quantity
				59500						Туре
Region origin	n of	Cold store		Ident	ification n	nark	Type of packag	ing		Net weight
Slaughterhous e		Treatment type			re of Number of page modity		Number of pack	kages		Batch No
		Date of collection/production		Manu plant	ıfacturing		Approval or reg number of plant/establishr	-	Test	

EURO	PEAN UNION					Certificate model POR-OOCYTES-EMB-A-INTRA				
	II. Health information			II.a	Certificate reference	II.b IN	ASOC reference			
	I, the ur	ndersigned	official veteri	inarian, hereb	by certify t	certify that:				
	<sup>(1)</sup> [II.1.				ine animals described in Part I have been collected, processed nbryo collection team <sup>(2)</sup> which					
		II.1.1.	is approved and kept in a register by the competent authority;							
		II.1.2.	facilities a		nt set out	regards responsibiliti in Part 2 of Annex				
	<sup>(1)</sup> [II.1.	described	The oocytes <sup>(1)</sup> / <i>in vitro</i> produced embryos <sup>(1)</sup> / micromanipulated embryos <sup>(1)</sup> of porcine animals described in Part I have been collected or produced, processed and stored, and dispatched by the embryo production team <sup>(2)</sup> which							
		II.1.1.	is approved	l and kept in	a register l	by the competent author	rity;			
ion		II.1.2.		nd equipment		regards responsibiliti Parts 2 and 3 of Anne				
Part II: Certification	II.2.		es <sup>(1)</sup> / embryc rom donor ar	are intended for artit	ficial reproc	luction and were				
t II: Ce		II.2.1.				e birth in the Union, of for entry into the Union		ered the Union in		
Part	(1)(	<sup>3)</sup> [II.2.2.	Aujeszky's		is or where	zone thereof which e an approved eradicat urried out;]				
		II.2.3.		ial control b		ember State or zone the petent authority in a th				
			П.2.3.1.	porcine and collection of	imals has	ith <i>Brucella abortus</i> , not been reported duri /tes <sup>(1)</sup> / embryos <sup>(1)</sup> , and to collection of the oc	ing the last in which d	42 days prior to uring at least the		
			<sup>(1)</sup> either	1	19(1)(f)(i)	and risk mitigating of Commission D ave been introduced;				
			<sup>(1)</sup> and/or	-	and <i>B. suis</i> on the esta	e for infection with <i>Br</i> has been carried out blishments in accorda ed Regulation (EU) 202	on the port ince with A	ine animals kept		

EUROPEAN UNION			Certificate model POR-OOCYTES-EMB-A-INTRA
		II.2.3.2.	where no clinical, serological, virological or pathological evidence of infection with Aujeszky's disease virus had been detected during the period of at least 12 months prior to collection of the oocytes <sup>(1)</sup> / embryos <sup>(1)</sup> .
	II.2.4.	symptoms	nined by the team veterinarian or a team member and did not show or clinical signs of transmissible animal diseases on the day of collection tes <sup>(1)</sup> / embryos <sup>(1)</sup> ;
	II.2.5.		ied as provided for in Article 52 or 54(2) of Commission Delegated (EU) 2019/2035;
	II.2.6.		d of at least 30 days prior to the date of first collection of the oocytes <sup><math>(1)</math></sup> / and during the collection period;
		II.2.6.1.	were kept on establishments not situated in a restricted zone established due to the occurrence of foot-and-mouth disease, infection with rinderpest virus, classical swine fever or African swine fever, or of an emerging disease relevant for porcine animals;
		II.2.6.2.	were kept on a single establishment where infection with <i>Brucella abortus, B. melitensis</i> and <i>B. suis</i> , infection with rabies virus, anthrax, infection with Aujeszky's disease virus and infection with porcine reproductive and respiratory syndrome virus have not been reported;
		II.2.6.3.	were not in contact with animals from establishments situated in a restricted zone due to the occurrence of diseases referred to in point II.2.6.1. or from establishments which do not meet the conditions referred to in point II.2.6.2.;
		II.2.6.4.	were not used for natural breeding;
	II.2.7.	comply wi	th the following conditions as regards foot-and-mouth disease
		II.2.7.1.	they come from establishments
			<ul> <li>situated in an area where foot-and-mouth disease has not been reported within a 10-km radius centred on the establishment for a period of at least 30 days immediately prior to the date of collection of the oocytes<sup>(1)</sup>/ embryos<sup>(1)</sup>;</li> <li>in which foot-and-mouth disease has not been reported during a period of at least 3 months immediately prior to the date of collection of the oocytes<sup>(1)</sup>/ embryos<sup>(1)</sup>;</li> </ul>
	<sup>(1)</sup> eith	er [II.2.7.2.	they were not vaccinated against foot-and-mouth disease;]
	(1)(4)	or [II.2.7.2.	they were vaccinated against foot-and-mouth disease during the period of 12 months prior to the date of collection or production of the embryos and
			II.2.7.2.1. have not been vaccinated against foot-and-mouth disease within the period of at least 30 days immediately prior to the date of collection of the embryos;

	II.2.7.2	.2. the semen used for fertilisation was collected from a male donor that complies with the conditions set out in point 1(b) or the semen complies with the conditions set out in point 2 of Chapter I of Part 5 of Annex II to Delegated Regulation (EU) 2020/686;
	II.2.7.2	3. prior to freezing, the embryos have been subjected to trypsin washing carried out in accordance with the recommendations of the IETS Manual <sup>(5)</sup> ;
	П.2.7.2	4. the embryos were stored deep frozen for a period of at least 30 days from the date of collection, and during this period the donor animal has not shown clinical signs of foot-and-mouth disease;]
<sup>(1)(6)</sup> [II.2.8.	respiratory syndrom	a serological test for infection with porcine reproductive and e virus, with negative results, on two occasions, at an interval of ys, the second test being performed within a period of 15 days ection.]
II.3. The oocyte	es <sup>(1)</sup> / embryos <sup>(1)</sup> descri	bed in Part I
II.3.1.		I, processed and stored in accordance with animal health t in Part $2^{(1)}$ /Part $3^{(1)}$ /Part $4^{(1)}$ /Part $5^{(1)}$ and Part 6 of Annex III to n (EU) 2020/686;
П.3.2.		or other packages on which the mark is applied in accordance rovided for in Article 10 of Delegated Regulation (EU) 2020/686 icated in Box I.30;
II.3.3.	are transported in a	container which:
	or prod	led and numbered prior to the dispatch by the embryo collection uction team under responsibility of the team veterinarian, or by cial veterinarian, and the seal bears the number as indicated in 9;
		en cleaned and either disinfected or sterilised before use, or is use container;
		en filled in with the cryogenic agent which not have been sly used for other products;]
<sup>(1)(8)</sup> [II.3.4.	are placed in straws	or other packages which are securely and hermetically sealed;
II.3.5.		container where they are separated from each other by physical being placed in secondary protective bags.]

<sup>(1)(9)</sup> [II.4.	The <i>in vivo</i> derived embryos <sup>(1)</sup> / <i>in vitro</i> produced embryos <sup>(1)</sup> / micromanipulated embry
[	described in Part I were conceived by artificial insemination using semen coming from a se collection centre, germinal product processing establishment or germinal product storage co
	approved for the collection, processing and/or storage of semen by the competent authority Member State or by the competent authority of a third country, territory or zone thereof liste Annex XI to Commission Implementing Regulation (EU) 2021/404.]
<sup>(1)(10)</sup> [II.5.	The following antibiotic or mixture of antibiotics <sup>(11)</sup> has been added to the collection, proces washing or storage media:
Notes	
	alth certificate shall be completed according to the notes for the completion of certificates prove of Annex I to Commission Implementing Regulation (EU) 2020/2235.
Part I:	
Box reference I	.11: <i>"Place of dispatch":</i> Indicate the unique approval number and the name and addre the embryo collection or production team of dispatch of the consignment of oocyte embryos.
Box reference I	number of the establishment of destination of the consignment of oocytes or embry
Box reference I	
Box reference I	1 0 1
Box reference I	.30: <i>"Type":</i> specify if <i>in vivo</i> derived embryos, <i>in vivo</i> derived oocytes, <i>in vitro</i> prod embryos or micromanipulated embryos.
	"Identification number": Indicate identification number of each donor animal.
	" <i>Identification mark</i> ": indicate mark on the straw or other packages where oocyte embryos of the consignment are placed.
	"Date of collection/production": indicate the date on which oocytes or embryos of consignment was collected or produced.
	"Approval or registration number of plant/establishment/centre": Indicate the un approval number of the embryo collection or production team by which the oocyt embryos were collected or produced.
	"Quantity": Indicate number of straws or other packages with the same mark.
Part II:	
	not applicable.
(2) Only emb	ryo collection or production teams approved by the competent authority and included in the reg o in Article 101(1)(b) of Regulation (EU) 2016/429 and Article 7 of Delegated Regulation
(3) Not applie	able for in vivo derived embryos subject to trypsin treatment.
(4) Option av	ailable only for the consignment of in vivo derived embryos.
use of em	f the International Embryo Transfer Society — A procedural guide and general information fo bryo transfer technology emphasising sanitary procedures, published by the International Em Society, 1 111 North Dunlap Avenue, Savoy, Illinois 61 874 USA (http://www.iets.org/).
(6) Applicabl	e for <i>in vivo</i> derived embryos.
oocytes, i	e for frozen oocytes or embryos. <sup>(8)</sup> Applicable for the consignment where in one contra <i>n vivo</i> derived embryos, <i>in vitro</i> produced embryos and micromanipulated embryos of por the placed and transported
(0)	re placed and transported.
Does not	apply to oocytes. y attestation in case antibiotics were added.
mandator	name(s) of the antibiotic(s) added and its(their) concentration.
(insert the	

## EUROPEAN UNION

Certificate model POR-OOCYTES-EMB-A-INTRA

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