#### ANIMAL HEALTH CERTIFICATE FOR THE MOVEMENT BETWEEN MEMBER STATES OF CONSIGNMENTS OF GERMINAL PRODUCTS LISTED BELOW, DISPATCHED AFTER 20 APRIL 2021 FROM THE GERMINAL PRODUCT PROCESSING ESTABLISHMENT:

ROPEAN	UNION				INTR	
I.1	Consignor		I.2	IMSOC reference		
	Name		I.2a	Local reference		
	Address		1.3	Central Competent Authority	QR CODE	
	Country	ISO country code	I.4	Local Competent Authority		
1.5	Consignee	,		Operator conducting assembly operations independently of an		
	Name			<b>establishment</b> Name	Registration No	
	Address			Address		
I.5 I.7 I.8 I.11	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	
	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		
	🗆 Railway	Road vehicle		Country	ISO country code	
			I.17	Accompanying documents		
	Identification	□ Other		Туре	Code	
Document			Country Commercial document reference	ISO country code		

### (MODEL 'POR-GP-PROCESSING-INTRA')

I.18	Transport co	onditions	Ambier	nt			□ Chill	ed	□ Frozen		
I.19	Container n	umber/Se	al number								
	Container No				Seal N	Ňo					
I.20	Certified as	or for									
□ Furt	her keeping		□ Slaughter			Cont	fined esta	ablishment	Germina	al products	S
🗆 Reg	istered equine animal		Travelling ci	rcus/anima	al act	🗆 Exhi	bition		□ Event or	activity r	near borders
🗆 Rele	ease into the wild		Dispatch centre		Relaying area/purification		Ornamental aquaculture				
					centre		establishment				
🗆 Furt	her processing		Organic fertilizers and soil		Technical use		□ Quarant	Quarantine or similar			
			improvers						establishm	ent	
D Proc	lucts for human consu	mption	Pollination			□ Live	aquatic	animals for	□ Other		
						human	consum	ption			
I.21	□ For trans	it through	n a third countr	у							
	Third countr	У				IS	O count	ry code			
	Exit point					В	CP code				
	Entry point					В	CP code				
I.22	□ For transit thro	ugh Mem	ber State(s)			I.23	□ For e	xport			
	Member State		ISO	country c	ode		Third	country	IS	SO country	y code
	Member State		ISO	country c	ode		Exit p	ooint	В	CP code	
	Member State		ISO	country c	ode						
I.24	Estimated journey	time				I.25	Journ	iey log	□ yes		□ no
I.26	Total number of pa	ickages				I.27	Total	quantity			
I.28	Total net weight/gr	oss weigh	ıt (kg)			I.29	Total	space foreseen	for the cons	signment	
I.30	Description of cons	ignment									
CN cc	de Species	Subspo	ecies/Category	Sex	Iden syste	tification		Identification	number	Age	Quantity
					sysu	2111					Туре
Regio	n of origin	Cold s	tore		Iden	tification	mark	Type of packa	ging		Net weight
T							1		Detal Ma		
Slaughterhouse Tre		Ireatn	51		ure of Numb modity		Number of pac	ckages		Batch No	
						-					
		Date o collect	f ion/production		Man plan	ufacturing t	g	Approval or re number of	gistration	Test	
		0011001	production		Phillip	-		plant/establish	ment/centre		

EURO	<b>PEAN UNION</b>			Certificate mode	I POR-C	P-PROCESSING-INTRA				
	II. Health inform	ation	II.a	Certificate reference	II.b	IMSOC reference				
	I, the undersigned official veterinarian, hereby certify that:									
	II.1. The germinal product processing establishment <sup>(1)</sup> described in Box I.11. at which the semen <sup>(2)</sup> / oocytes <sup>(2)</sup> / <i>in vivo</i> derived embryos <sup>(2)</sup> / <i>in vitro</i> produced embryos <sup>(2)</sup> / micromanipulated embryos <sup>(2)</sup> was/were processed and stored:									
	II.1.1.	is approved and kept in a register by the	compe	tent authority;						
	II.1.2.	II.1.2. complies with requirements as regards responsibilities, operational procedures, facilities and equipment set out in Part 4 of Annex I to Commission Delegated Regulation (EU) 2020/686.]								
	II.2. The sembry	$emen^{(2)}$ oocytes <sup>(2)</sup> / <i>in vivo</i> derived emb os <sup>(2)</sup> described in Part I is/are intended for	oryos <sup>(2)</sup> r artific	/ in vitro produced e ial reproduction and	mbryos	(2)/ micromanipulated				
rari II: Ceruncauon	(2)either[11.2.1.	embryo collection team <sup>(2)(3)</sup> / by an emb germinal product processing establish centre <sup>(2)(3)</sup> situated in the Member State requirements as regards responsibilities Part 1 <sup>(2)</sup> /Part 2 <sup>(2)</sup> /Part 3 <sup>(2)</sup> /Part 4 <sup>(2)</sup> /Part 5 <sup>(2)</sup> was/were moved to the germinal produ	d, processed and stored in a semen collection centre <sup>(2)(3)</sup> / by an embryo production team <sup>(2)(3)</sup> , and/or processed and stored in a blishment <sup>(2)(3)</sup> , and/or stored in a germinal product storage State of its/their collection or production and complying with ties, operational procedures, facilities and equipment set out in art $5^{(2)}$ of Annex I to Delegated Regulation (EU) 2020/686, and oduct processing establishment indicated in Box I.11. situated collection or production under animal health certification are provided for int							
Ceru	<sup>(2)</sup> either	[Model POR-SEM-A-INTRA <sup>(4)</sup> ;]								
	<sup>(2)</sup> and/or	[Model POR-SEM-B-INTRA <sup>(4)</sup> ;]								
Part	<sup>(2)</sup> and/or	[Model POR-OOCYTES-EMB-A-INTF	RA <sup>(4)</sup> ;]							
	<sup>(2)</sup> and/or	[Model POR-OOCYTES-EMB-B-INTR	(4);]							
	<sup>(2)</sup> and/or	[Model POR-OOCYTES-EMB-C-INTR	( <sup>4)</sup> ;]							
	<sup>(2)</sup> and/or	[Model POR-GP-PROCESSING-INTR.	A <sup>(4)</sup> ;]							
	<sup>(2)</sup> and/or	[Model POR-GP-STORAGE-INTRA <sup>(4)</sup> ;	]]							
	<sup>(2)</sup> and/or[II.2.1.	embryo collection team <sup>(2)(3)</sup> / by an emb germinal product processing establish centre <sup>(2)(3)</sup> situated in the Member State requirements as regards responsibilities Part 1 <sup>(2)</sup> /Part 2 <sup>(2)</sup> /Part 3 <sup>(2)</sup> /Part 4 <sup>(2)</sup> /Part 5 <sup>(2</sup>	processed and stored in a semen collection centre <sup>(2)(3)</sup> / by an abryo production team <sup>(2)(3)</sup> , and/or processed and stored in a shment <sup>(2)(3)</sup> , and/or stored in a germinal product storage ate of its/their collection or production and complying with es, operational procedures, facilities and equipment set out in $5^{(2)}$ of Annex I to Delegated Regulation (EU) 2020/686, and buct processing establishment indicated in Box I.11. situated by certificate(s) in accordance with:							
	<sup>(2)</sup> either	[Model POR-SEM-A-INTRA <sup>(4)</sup> ;]								
		[Model POR-SEM-B-INTRA <sup>(4)</sup> ;]								
	<sup>(2)</sup> and/or	[Model POR-OOCYTES-EMB-A-INTF	RA <sup>(4)</sup> ;]							

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<sup>(2)</sup> and/or	[Model POR-OOCYTES-EMB-B-INTRA <sup>(4)</sup> ;]
<sup>(2)</sup> and/or	[Model POR-OOCYTES-EMB-C-INTRA <sup>(4)</sup> ;]
<sup>(2)</sup> and/or	[Model POR-GP-PROCESSING-INTRA <sup>(4)</sup> ;]
<sup>(2)</sup> and/or	[Model POR-GP-STORAGE-INTRA <sup>(4)</sup> ;]]
<sup>2)</sup> and/or [II.2.1.	has/have been collected or produced, processed and stored in a semen collection centre <sup>(2)(3)</sup> / by an embryo collection team <sup>(2)(3)</sup> / by an embryo production team <sup>(2)(3)</sup> , and/or processed and stored in a germinal product processing establishment <sup>(2)(3)</sup> , and/or stored in a germinal product storage centre <sup>(2)(3)</sup> situated in a third country, territory or zone thereof listed in Annex XI to Commission Implementing Regulation (EU) 2021/404 and complying with requirements as regards responsibilities, operational procedures, facilities and equipment set out in Part 1 <sup>(2)</sup> /Part 2 <sup>(2)</sup> /Part 3 <sup>(2)</sup> /Part 4 <sup>(2)</sup> /Part 5 <sup>(2)</sup> of Annex I to Delegated Regulation (EU) 2020/686, and entered the Union accompanied by certificate(s) in accordance with:
<sup>(2)</sup> either	[Model POR-SEM-A-ENTRY <sup>(4)</sup> ;]
<sup>(2)</sup> and/or	[Model POR-SEM-B-ENTRY <sup>(4)</sup> ;]
<sup>(2)</sup> and/or	[Model POR-OOCYTES-EMB-ENTRY <sup>(4)</sup> ;]
<sup>(2)</sup> and/or	[Model POR-GP-PROCESSING-ENTRY <sup>(4)</sup> ;]
<sup>(2)</sup> and/or	[Model POR-GP-STORAGE-ENTRY <sup>(4)</sup> ;]]
II.2.2.	has/have been collected, processed and stored in accordance with animal health requirements set out in Annex III to Delegated Regulation (EU) 2020/686;
П.2.3.	is/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/or Article 83(a) of Commission Delegated Regulation (EU) 2020/692 and that mark is indicated in Box I.30;
II.2.4.	is/are transported in a container which:
	II.2.4.1. was sealed and numbered prior to the dispatch from the germinal product processing establishment under responsibility of the centre veterinarian, or by an official veterinarian, and the seal bears the number as indicated in Box I.19;
	II.2.4.2. has been cleaned and either disinfected or sterilised before use, or is single-use container;
(2)(5	[II.2.4.3. has been filled in with the cryogenic agent which not have been previously used for other products;]
<sup>(2)(6)</sup> [II.2.5.	is/are placed in straws or other packages which are securely and hermetically sealed;
П.2.6.	is/are transported in a container where they are separated from each other by physical compartments or by being placed in secondary protective bags.]
Notes	
	alth certificate shall be completed according to the notes for the completion of certificates provided 2 of Annex I to Commission Implementing Regulation (EU) 2020/2235.

EUROPEAN UNION	Certificate model POR-GP-PROCESSING-INTRA						
Part I:							
Box reference I.11:	<i>"Place of dispatch":</i> Indicate the unique approval number and the name and address of the germinal product processing establishment of dispatch of the consignment of semen, oocytes, and/or embryos. Only germinal product processing establishments 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.						
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 semen, oocytes, and/or embryos.						
Box reference I.17:	"Accompanying documents": Number(s) of related original certificate(s) shall correspond to the serial number of the individual official document(s) or health certificate(s) that accompanied the semen, oocytes and/or embryos described in Part I from the semen collection centre where the semen was collected, and/or the embryo collection and/or production team by which the oocytes and/or embryos were collected or produced, and/or the germinal product processing establishment where the semen, oocytes or embryos were processed and stored, and/or the germinal product storage centre where the semen, oocytes or embryos were stored to the germinal product processing establishment described in Box I.11. The original(s) of those document(s) or those certificate(s) or the officially endorsed copies thereof must be attached to this certificate.						
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:	<i>"Type":</i> Specify if semen, <i>in vivo</i> derived embryos, <i>in vivo</i> derived oocytes, <i>in vitro</i> produced 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 semen, oocytes and/or embryos of the consignment are placed.						
	"Date of collection/production": Indicate the date on which semen, oocytes and/or embryos of the consignment was/were collected or produced.						
	"Approval or registration number of plant/establishment/centre": Indicate the unique approval number of the semen collection centre where the semen was collected, and/or of the embryo collection and/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.						
Part II:							
	duct processing establishments approved by the competent authority and included in the in Article 101(1)(b) of Regulation (EU) 2016/429 and Article 7 of Delegated Regulation						
<sup>(2)</sup> Delete if not applic	able.						
	duct establishments approved by the competent authority and included in the register le 101(1)(b) of Regulation (EU) 2016/429 and Article 7 of Delegated Regulation (EU)						

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# Certificate model POR-GP-PROCESSING-INTRA

<ul> <li><sup>(4)</sup> The original(s) of the document(s) or the health certificate(s) or the officially endorsed copies of thereof that accompanied the semen, oocytes or embryos described in Part I from the semen collection centre where the semen was collected, and/or the embryo collection or production team by which the oocytes and/or embryos were collected or produced, and/or the germinal product processing establishment where the semen, oocytes or embryos were processed and stored, and/or the germinal product storage centre where the semen, oocytes or embryos dispatch described in Box I.11 must be attached to this certificate.</li> <li><sup>(5)</sup> Applicable for frozen semen, oocytes or embryos.</li> <li><sup>(6)</sup> Applicable for the consignment where in one container semen, oocytes, <i>in vivo</i> derived embryos, <i>in vitro</i> produced embryos and micromanipulated embryos of porcine animals are placed and transported.</li> </ul>						
Offi	cial veterinarian					
Nam	ne (in capital letters)	Qualification and title				
Loca	al Control Unit name	Local Control Unit code				
Date	2					
Starr	q	Signature				