EUROPEAN UNION INTRA

	I.1. Consignor						I.2. IMSOC ref	erence		I.2.a. I	ocal reference	
	Name										ntral Competen	t Authority
	Address									cal Competent A		
	Country ISO Code									•	,	
	,											
Ħ	I.5. Consignee						I.6. Operator conducting assembly operations independently of an establishment					
of consignment	Name											
Ħ		Address Country ISO Code						Name Address				
٠Ë	Country							Approval Number				
ns								Country ISO Code				
2		country				150 couc						
	I.7. Country of orig	gin				ISO Code	I.9. Country of	destinatio	n			ISO Code
Part I: Description												
ΡÜ	I.8. Region of origi	in				Code	I.10. Region of	destinatio	n			Code
Ξ	I.11. Place of dispa						I.12. Place of destination					
es	Name						Name					
A	Address						Address					
Ţ	Approval Number	r					Approval Number					
ar	Country			ISO	Code		Country ISO Code					
4							_					
	I.13. Place of loadi	ng					I.14. Date and	time of de	parture			
	Name											
	Address											
	Approval Number	r										
	Country			ISO (Code							
	I.15. Means of Tra		I.16. Transporter									
	Mode	Internatio	nal	Identification	n		Name					
	Mode	transport		lacitilicatio	,,,,		Address Activity ID					
		document										
							Country			IS	O Code	
							I.17. Accompa	nying docu	iments			
							Commercial document			Date o	fissue	
							reference					
						Country			Place of issue	of		
	I 18 Transport co											
	I.18. Transport conditions Frozen ☐ Ambient ☐						Chilled □					
	I.19. Container No	/ Seal No										
	I.20. Certified as											
	Germinal product	s Ц										
	I.21. For transit th	rough a thi	trv									
	Third country	rough a thi	i a couri	.cry			ISO Code					
	Exit point						BCP code					
	Entry point						BCP code I.23. For export					
	I.22. For transit th	rough Mem	ber Sta	te(s)								
	Member State							Third country ISO Code				
							Exit point BCP code					
	I.26. Total number of packages I.27. Total quantity						I.25. Journey Log					
									I.28. Total g	ross we	ight	
	I.30. Description o											
		1 COHSIGIHI		_		T-3+:6:+:	Normalia and	0			N-4 6	
	Commodity		Specie	!S		Identification	Number	Quantity			Nature of com	ιποαπγ
	Identification Mark				Date of collect	rtion / Plant / Establishment / Type Centre						
					1,							
	1										'	

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EUROPEAN UNION 2023/594 (2021/403) POR-OOCYTES-EMB-A-											
	II. Health info	rmation									
	I, the undersigned official veterinarian, hereby certify that:										
Part II: Certification		prize animals described in Part I have been collected, processed and bryo collection team(2) which									
		II.1.1.	in a register by the competent authority;								
		II.1.2.	•	-	ements as regards responsibilities, operational procedures, facilities t in Part 2 of Annex I to Delegated Regulation (EU) 2020/686.]						
	(1) ° [II.1.	described		ve been colle	embryos(1)/ micromanipulated embryos(1) of porcine animals ected or produced, processed and stored, and dispatched by the						
		II.1.1.	is approve	d and kept i	in a register by the competent authority;						
		II.1.2.			ements as regards responsibilities, operational procedures, facilities t in Parts 2 and 3 of Annex I to Delegated Regulation (EU) 2020/686.]						
	II.2.		The oocytes(1)/ embryos(1) described in Part I are intended for artificial reproduction and were obtained from the donor animals which								
		II.2.1.	have been born and remained since birth in the Union, or have entered the Union in accordance with the requirements for entry into the Union;								
	-	(1)(3) □ [II.2.2.	disease vii		State or zone thereof which is free from infection with Aujeszky's e an approved eradication programme for infection with Aujeszky's d out;]						
		II.2.3.	come from establishments in a Member State or zone thereof, or from establishments under official control by the competent authority in a third country or territory, or a zone thereof								
			II.2.3.1.	animals ha	nfection with Brucella abortus, B. melitensis and B. suis in porcine as not been reported during the last 42 days prior to collection of the / embryos(1), and in which during at least the 12 month period prior to of the oocytes(1)/ embryos(1)						
			(1) □ either	[II.2.3.2.1.	biosecurity and risk mitigating measures set out in Article 19(1)(f)(i) of Delegated Regulation 2020/688 have been introduced;						
			(1) □ and/or	[II.2.3.2.2.	surveillance for infection with Brucella abortus, B. melitensis and B. suis has been carried out on the porcine animals kept on the establishments in accordance with Article 19(1)(f)(ii) of Delegated Regulation 2020/688;]						
			II.2.3.2.	with Aujes	clinical, serological, virological or pathological evidence of infection szky's disease virus had been detected during the period of at least 12 rior to collection of the oocytes(1)/ embryos(1).						
		II.2.4.		ns of transr	e team veterinarian or its team member and did not show symptoms or missible animal diseases on the day of collection of the oocytes(1)/						
		II.2.5.	are identif	ied as provi	ided for in Article 52 or 54(2) of Delegated Regulation (EU) 2019/2035;						
	II.2.6. for a period of at least 30 days prio embryos(1) and during the collection				30 days prior to the date of first collection of the oocytes(1)/g the collection period;						
			II.2.6.1.	the occurr	on establishments not situated in a restricted zone established due to rence of foot-and-mouth disease, infection with rinderpest virus, wine fever and African swine fever or of an emerging disease relevant rcine animals;						
			II.2.6.2.	melitensis Aujeszky's	on a single establishment where infection with Brucella abortus, B. and B. suis, infection with rabies virus, anthrax, infection with s disease virus and infection with porcine reproductive and respiratory virus have not been reported;						
zone due to the occi				zone due t	n contact with animals from establishments situated in a restricted to the occurrence of diseases referred to in point II.2.6.1. or from nents which do not meet the conditions referred to in point II.2.6.2.;						
			II.2.6.4.	were not u	used for natural breeding;						

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	II. Health info	rmation						
		II.2.7.	comply wit	h the follov	، ving conditio	ons as regards foot-and-mouth disease		
			II.2.7.1.	they come	from establis	shments		
Part II: Certification	a 10-km radius centred					ere foot-and-mouth disease has not been reported within on the establishment for a period of at least 30 days the date of collection of the oocytes(1)/ embryos(1);		
			-		nths immedia	th disease has not been repor ately prior to the date of colle		
		(1) \circ either	[II.2.7.2.	they were	not vaccinate	ed against foot-and-mouth di	sease;]	
		(1)(4) ○ or	[II.2.7.2.	-		gainst foot-and-mouth diseas e of collection or production	-	
				II.2.7.2.1.		en vaccinated against foot-ar least 30 days immediately pr ryos;		
				II.2.7.2.2.	complies w	used for fertilisation was colloith the conditions set out in pith the conditions set out in pto Delegated Regulation (EU)	oint 2 of Chapter I of Part 5	
				II.2.7.2.3.		ezing, the embryos have beer rried out in accordance with al(5);		
				II.2.7.2.4.	from the da	s were stored deep frozen for te of collection, and during th wn clinical signs of foot-and-	his period the donor animal	
	(1)(6) were subjected to a serological to [II.2.8. syndrome virus, with negative redays, the second test being perfo				negative resu	ılts, on two occasions, at an iı	nterval of not less than 21	
	II.3.	The oocyte	s(1)/ embryo	os(1) descril	bed in Part I			
		II.3.1.		2(1)/Part 3(stored in accordance with and eart 5(1) and Part 6 of Annex	mal health requirements set III to Delegated Regulation	
		II.3.2.	_	nts provide	d for in Artic	nges on which the mark is apple 10 of Delegated Regulation		
		II.3.3.	are transpo	orted in a co	ontainer whic	ch:		
			II.3.3.1.	production	n team under	red prior to the dispatch by the responsibility of the team ve eal bears the number as indic	eterinarian, or by an official	
			II.3.3.2.	has been container;		ither disinfected or sterilised	before use, or is single-use	
			(1)(7)II.3.3. 3.	has been for other p		the cryogenic agent which no	t have been previously used	
		(1)(8) □ [II.3.4.	are placed	in straws o	r other packa	nges which are securely and h	nermetically sealed;	
		II.3.5.				re they are separated from e n secondary protective bags.		
- 1	(1)(9) □ [II.4.	in Part I we germinal p collection, competent	ere conceive roduct proc processing a	ed by artific essing estal and/or stora f a third cou	ial inseminat olishment or age of semen antry, territor	ced embryos(1)/ micromanip tion using semen coming fron germinal product storage cen by the competent authority or ry or zone thereof listed in An	n a semen collection centre, ntre approved for the of a Member State or by the	

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	II. Health info	rmation								
	(1)(10) □ [II.5.	The following antibiotic or mixture of antibiotics(10) has been added to the collection, processing, washing or storage media:								
	II.6.	This certificate is valid for 10 days from the date of issuing.								
	II.7									
Part II: Certification		(2) 🗆	[Germinal products obtained from porcine animals kept in restricted zones II in compliance with the special disease control measures relating to African swine fever laid down in Commission Implementing Regulation (EU) 2023/594.]							
		(2) 🗆	[Germinal products obtained from porcine animals kept in restricted zone III in compliance with the special disease control measures relating to African swine fever laid down in Commission Implementing Regulation (EU) 2023/594.]							
Par										
	1									

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II. Health information

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:

Certification Box reference

ä

Part]

"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.

I.11:

Box reference

"Place of destination": Indicate the address and unique registration or approval number of the

establishment of destination of the consignment of oocytes or embryos.

I.12:

Box Seal number shall be indicated.

reference

I.19: Box

Total number of packages shall correspond to the number of containers.

reference I.26:

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.

"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.

Part II:

- Delete if not applicable. (1)
- (2)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.
- Not applicable for in vivo derived embryos subject to trypsin treatment. (3)
- (4)Option available only for the consignment of in vivo derived embryos.
- Manual of the International Embryo Transfer Society A procedural guide and general information for (5) the use of embryo transfer technology emphasising sanitary procedures, published by the International Embryo Transfer Society 1 111 North Dunlap Avenue, Savoy, Illinois 61 874 USA (http://www.iets.org/).
- (6)Applicable for in vivo derived embryos.
- (7)Applicable for frozen oocytes or embryos.
- Applicable for the consignment where in one container oocytes, in vivo derived embryos, in vitro (8)produced embryos and micromanipulated embryos of porcine animals are placed and transported.
- (9)Does not apply to oocytes.
- (10)Mandatory attestation in case antibiotics were added.
- Insert the name(s) of the antibiotic(s) added and its(their) concentration.

Certifying Officer/Official veterinarian

Name (in capital letters) Authority name Date of signature Signature

Stamp

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