EUROPEAN UNION INTRA

	I.1. Consignor					I.2. IMSOC reference I.2.a. Local reference			
	Name Address						I.3. Centra	al Competent Authority	
	Country ISO Code						I.4. Local	Competent Authority	
or consignment	I.5. Consignee Name Address Country ISO Code				I.6. Operator conducting assembly operations independently of an establishment Name Address Approval Number				
3					Country		ISO C	ode	
	I.7. Country of orig	gin	ISO Code	I.9. Country o	f destination		ISO Code		
	I.8. Region of origi	in		Code	I.10. Region o	f destination		Code	
rarı I: Descripuon	I.11. Place of dispatch Name Address Approval Number Country ISO Code				Name Address	ddress pproval Number			
ŀ	I.13. Place of loadi	ing			I.14. Date and time of departure				
	Name Address Approval Number Country ISO Code					·			
ŀ	I.15. Means of Tra	nsport			I.16. Transporter				
	Mode	International transport	Identification		Name				
ŀ	document				Address Activity ID				
ŀ					Country		ISO C	ode	
ŀ					L17. Accompa	anying documents			
ŀ				Accompanyir reference					
					Date of issue Country Place of issue				
- 1	I.18. Transport coi	nditions							
	Ambient ☐ Chilled ☐			Frozen 🗆					
İ	I.19. Container No / Seal No								
- 1	I.20. Certified as Germinal products								
İ	I.21. For transit th	rough a third coun	try						
	Third country Exit point Entry point				ISO Code BCP code BCP code				
ŀ		rough Member Sta	te(s)		I.23. For export				
	Member State ISO Code			Third country ISO Code Exit point BCP code					
Ī				I.25. Journey Log					
Ī	I.26. Total number of packages I.27. Total quantity				I.28. Total gross weight				
	I.30. Description o	f consignment	,						
						or 3, unfit for human o	consumpti	on	
	05119985 Other					T	1		
Н	#1. Commodity		cation Number	Quantity		Nature of commodity Identification Mark			
ļ	Species	Package	count	Date of collection	on / production	Plant / Establishment / Ce	ntre Ty	<i>r</i> pe	

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	II. Health information									
		I, the undersigned official veterinarian, hereby certify, that:								
tion	II.1.	The semen(1)/ in vivo derived embryos(1)/ oocytes(1)/ in vitro produced embryos(1)/ micromanipulated embryos(1) described in Part I has/have been collected or produced, processed and stored, and dispatched from the confined establishment(2) which								
		II.1.1.	1. is approved, assigned with a unique approval number and kept in a register by the competent authority;							
Part II. Certification		II.1.2.	.1.2. complies with requirements as regards quarantine, isolation and other biosecurity measures, surveillance and control measures, facilities and equipment referred to in Article 16 of Commission Delegated Regulation (EU) 2019/2035.							
mt III	II.2.	The semen(1)/ oocytes(1)/ embryos(1) described in Part I is/are intended for artificial reproduction a was/were obtained from donor animals which								
ثم		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;							
		II.2.2.	have remained in a single confined establishment of origin for a period of at least 30 days prior to the date of collection of the semen(1)/ oocytes(1)/ embryos(1);							
	(1)	□ [II.2.3.	are bovine animals and they are identified as provided for in Article 38 of Delegated Regulation (EU) 2019/2035.]							
	(1)	□ [II.2.3.	are porcine animals and they are identified as provided for in Article 52(1) or 54(2) of Delegated Regulation (EU) 2019/2035.]							
	(1)	□ [II.2.3.								
	(1)	□ [II.2.3.	are equine animals and they are identified as provided for in Article 58(1) or 59(1) or 62(1) of Delegated Regulation (EU) 2019/2035.]							
	(1)	□ [II.2.3.	are terrestrial animals other than bovine, porcine, ovine, caprine and equine animals and they are identified and registered in accordance with the rules of the confined establishment.]							
	II.3.		e semen(1)/ oocytes(1)/ embryos(1) described in Part I comes/come from the confined establishment licated in Box I.11. and is/are destined to another confined establishment.							
	II.4.	According to official information, the semen(1)/ oocytes(1)/ embryos(1) described in Part I was/were obtained from donor animals which								
		II.4.1.	do not come from a confined establishment, nor have been in contact with animals from confined establishment, situated in a restricted zone established due to the occurrence category A disease, referred to in the Annex to Commission Implementing Regulation (E 2018/1882, or of an emerging disease relevant for species in those donor animals;							
		II.4.2.	come from a confined establishment where no category D disease relevant for that species as referred to in the Annex to Implementing Regulation (EU) 2018/1882 has been reported for a period of at least 30 days prior to the date of collection of the semen(1)/ oocytes(1)/ embryos(1).							
	II.5.	To the best of my knowledge and as declared by the operator, the semen(1)/ oocytes(1)/ embryos(1) described in Part I was/were obtained from donor animals which								
		II.5.1.	have been clinically examined by the establishment veterinarian responsible for the activities carried out at the confined establishment and showed no disease symptoms on the day of collection of the semen(1)/ oocytes(1)/ embryos(1);							
		II.5.2.	as much as possible, were not used for natural breeding during a period of at least 30 days prior to the date of collection of the semen(1)/ oocytes(1)/ embryos(1) and during the collection period.							
	II.6.	establishn semen(1)/	st of my knowledge and based on the documentary check of the data submitted by the nent veterinarian responsible for the activities carried out at the confined establishment, the oocytes(1)/ embryos(1) described in Part I is/are placed in straws or other packages on which is applied in accordance with requirements provided for in							
	(1)(2)	☐ [Article 10 of Commission Delegated Regulation (EU) 2020/686 and that mark is indicated in I								
	(1)(3)	\square [Article 11 of Delegated Regulation (EU) 2020/686 and that mark is indicated in Box I.30.]								

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	II. Health info	II. Health information							
	II.7.	The seme	n(1)/ oocytes((1)/ embryos(1) describe	d in Part I				
		II.7.1.							
Dart II: Cortification	100		II.7.1.1. was sealed and numbered prior to the dispatch by the establishment veterinarian responsible for the activities carried out at the confined establishment, or by an official veterinarian, and the seal bears the nur indicated in Box I.19;						
			II.7.1.2.	has been cleaned and container;	either disinfecte	d or sterilise	d before use, or is single-use		
		(1)(4)	□ [II.7.1.3.	has been filled in with the cryogenic agent which not have been previously us for other products;]					
*	(1)(2)(5)	□ [II.7.2.	is/are place	ed in straws or other pa	ckages which are	e securely an	d hermetically sealed;		
-	24	II.7.3.	is/are transported in a container where they are separated from each other by physical compartments or by being placed in secondary protective bags.]						
	(1)	either	o [II.8 Germinal products (semen, ova and/or embryos, indicate as appropriate) obtained from bovine animals kept in vaccination zone I in relation to emergency protective vaccination against lumpy skin disease, in compliance with Article 13(3) of, and Annex IX, Part 3, point (3.4.1), to Commission Delegated Regulation (EU) 2023/361.]						
	(1)	or © [II.8 Germinal products (semen, ova and/or embryos, indicate as appropriate) obtained from bovine animals kept in vaccination zone II in relation to emergency protective vaccination against lumpy skin disease, in compliance with Article 13(3) of, and Annex IX, Part 3, point (3.4.2), to Commission Delegated Regulation (EU) 2023/361.]							
	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.								
	Box reference I.11:	reference of dispatch of the consignment of semen, oocytes or embryos.							
	Box reference I.12:	"Place of destination": Indicate the address and the unique approval number of the confined establishment of destination of the consignment of semen, oocytes or embryos.							
Box "Type": "Type": Specify if semen, in vivo derived embryos, reference embryos or micromanipulated embryos. I.30:					ed embryos, in v	ivo derived (oocytes, in vitro produced		
"Identification number": Indicate identification number of each donor animal.						nal.			
"Identification mark": Indicate mark on the straw or other packages where semen, oocytes of the consignment are placed.							semen, oocytes or embryos		
"Date of collection/production": Indicate the date on which semen, oocytes consignment were collected or produced.						nen, oocytes	or embryos of the		
			establishmen	ion number of plant/est at of the collection or pro			que approval number of the embryos of the		
"Quantity": Indicate number of straws or other packages with the same						the same ma	rk.		
	Part II:								
(1) Delete if not applicable.									
	(2)	Applicable equine an	able for the consignment of semen, oocytes or embryos of bovine, porcine, ovine, caprine or animals.						
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Applicable for the consignment of semen, oocytes or embryos of terrestrial animals other than bovine,

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	II. Health info	rmation							
		porcine, ovine, caprine or equine animals.							
	(4) Applicable for frozen semen, oocytes or embryos.								
	(5)	Applicable for the consignment where in one	container oocytes, in vivo der	ived embryos, in vitro					
	produced embryos and micromanipulated embryos of bovine, porcine, ovine, caprine or equine								
Z	animals are placed and transported.								
Certifying Officer/Official veterinarian Name (in capital letters) Date of signature Stamp Qualification and title Signature									
္ဌ	Name (in capi		Qualification and title						
ij	Date of signati	ure	Signature						
ē	Stamp								
日									
ar									
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