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	I.1. Consignor				I.2. IMSOC reference		I.2.a. Local refer	ence
	Name						I.3. Central Comp	petent Authority
	Address							
	Country ISO Code				I.4. Local Competent Authority			
ut	I.5. Consignee	I.5. Consignee				I.6. Operator conducting assembly operations independently of an		
of consignment	Name				establishment Name Address Approval			
3	Address							
ĭ	Country		ISO Co	de				
51					Number			
ı					Country		ISO Code	
	I.7. Country of orig	σin		ISO Code	I.9. Country of destinati	nn		ISO Code
	in country of only	5111			is country of destinati	011		
<u>1</u>	I.8. Region of origi	in		Code	I.10. Region of destinati	on		Code
Part I: Description								
al E	I.11. Place of dispa	atch			I.12. Place of destination	1		
::	Name				Name Address Approval			
11	Address Approval							
ية الت	Number				Number			
	Country		ISO Co	de	Country		ISO Code	
	I 12 Dlaga of loadi	ng			I 14 Data and time of de	nartura		
	I.13. Place of loading				I.14. Date and time of departure			
	Name Address							
	Approval							
	Number		ISO Co	do				
	Country		150 00	ue				
	I.15. Means of Tra	nsport	1		I.16. Transporter			
	Mode	International transport	Identificati	on	Name			
		document			Address			
					Approval Number			
					Country		ISO Code	
					I.17. Accompanying doc	uments		
					Document Type Accompanying document			
				Date of Issue Country				
					Place of issue			
	I.18. Transport co	nditions				_		
	Chilled ☐ Ambient ☐			Frozen 🗆				
	I.19. Container No	/ Seal No						
	I.20. Certified as							
	Germinal product	s 🗆						
	I.21. For transit th	rough a third coun	try					
	Third country	Third country			ISO Code			
	Exit point				BCP code			
	Entry point			BCP code				
	I.22. For transit th	rough Member Sta	te(s)		I.23. For export			
	Member State		ISO	Code	Third country		ISO Code	
					Exit point		BCP code	
	I.24. Estimated jou	24. Estimated journey time			I.25. Journey Log			
	I.26. Total number	r of packages		I.27. Total quantity		I.28. Total g	ross weight	
	I.30. Description o	f consignment						
	1. 05 PRODUCTS OF ANIMAL ORIGIN, NOT ELSEWHERE SPECIFIED OR INCLUDED							
	0511 Animal products not elsewhere specified or included; dead animals of Chapter 1 or 3, unfit for human consumption							
	051199 Other							
	05119985 (Other						

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	- 1	#1. Commodity	Identification Number	Quantity	Nature of commodity	Identification Mark	
		Species	Package count	Date of collection / production	Plant / Establishment / Centre	Type	
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1.5	31						
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Dart I: Description of consignment							
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UNION 2024/1044 (2021/403) Model POR-OOCYTES								
	II. Health information							
	I the unde	ersigned of	ficial veterina	arian hereby certify that the \Box [oval (1) \Box [embryos] (1) described in Part I:				
	I, the undersigned official veterinarian, hereby certify that the □ [ova] (1) □ [embryos] (1) described in Part I II.1. were □ [produced] (1) □ [collected] (1), processed and stored by an □ [embryo collection team] □ [embryo production team] (1) (2) approved and supervised in accordance with Chapter I(III) of Annex D to Directive 92/65/EEC;							
<u>দ</u>	II.2.	meet the	requirements of Chapter III(II) of Annex D to Directive 92/65/EEC;					
Certification	II.3. come from donor females of the porcine species which meet the requirements of Chapter IV(2) of Anne D to Directive 92/65/EEC;							
Part II: Certif	(1) o either	[II.4.	are in vivo	derived embryos which:				
			II.4.1.	were conceived as a result of artificial insemination with semen meeting the requirements of Directive 90/429/EEC,				
2			II.4.2.	originate from a Member State or region thereof:				
			(1) \circ either	[listed in Annex I to Decision 2008/185/EC and are destined for a Member State or region thereof listed in Annex I to Decision 2008/185/EC;]				
			(1) ○ or	[listed in Annex I to Decision 2008/185/EC and are destined for a Member State or region thereof not listed in Annex I or II to Decision 2008/185/EC;]				
			(1) ○ or	[listed in Annex II to Decision 2008/185/EC and are destined for a Member State or region thereof listed in Annex I to Decision 2008/185/EC and have been washed with trypsin;]				
			(1) ○ or	[listed in Annex II to Decision 2008/185/EC and are destined for a Member State or region thereof listed in Annex II to Decision 2008/185/EC;]				
			(1) ○ or	[not listed in Annex I or II to Decision 2008/185/EC and are destined for a Member State or region thereof listed in Annex I or II to Decision 2008/185/EC and have been washed with trypsin;]				
			(1) ∘ or	[not listed in Annex I or II to Decision 2008/185/EC and are destined for a Member State or region thereof not listed in Annex I or II to Decision 2008/185/EC;];]				
	(1) or	[II.4.	are 🗆 [in	vitro produced] (1) 🛘 [micromanipulated] (1) embryos which:				
			II.4.1.	were conceived as a result of in vitro fertilisation with semen meeting the requirements of Directive 90/429/EEC,				
			II.4.2.	originate from a Member State or region thereof:				
			(1) ○ either	[listed in Annex I to Decision 2008/185/EC and are destined for a Member State or region thereof listed in Annex I to Decision 2008/185/EC;]				
			(1) ∘ or	[listed in Annex I to Decision 2008/185/EC and are destined for a Member State or region thereof not listed in Annex I or II to Decision 2008/185/EC;]				
			(1) ∘ or	[listed in Annex II to Decision 2008/185/EC and are destined for a Member State or region thereof listed in Annex I to Decision 2008/185/EC and the donor females of the ova used for their production comply with the conditions of Article 1 of Decision 2008/185/EC;]				
			(1) ○ or	[listed in Annex II to Decision 2008/185/EC and are destined for a Member State or region thereof listed in Annex II to Decision 2008/185/EC;]				
			(1) ○ or	[not listed in Annex I or II to Decision 2008/185/EC and are destined for a Member State or region thereof listed in Annex I or II to Decision 2008/185/EC and the donor females of the ova used for their production comply with the conditions of Article 1 of Decision 2008/185/EC;]				
			(1) ∘ or	[not listed in Annex I or II to Decision 2008/185/EC and are destined for a Member State or region thereof not listed in Annex I or II to Decision 2008/185/EC;]]				
	(1) ○ or	[II.4.	are in vivo	derived ova which originate from a Member State or region thereof:				
			(1) o either	[listed in Annex I to Decision 2008/185/EC and are destined for a Member State or region thereof listed in Annex I to Decision 2008/185/EC;]				

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	VION 2024/1044 (2021/403) WIOUEI FOR-OOCI ILS-LWID-D-IN II							
	II. Health info	rmation						
		(1) ∘ or	[listed in Annex I to Decision 2008/185/EC and are destined for a Member State or region thereof not listed in Annex I or II to Decision 2008/185/EC;]					
noi		(1) ∘ or	[listed in Annex II to Decision 2008/185/EC and are destined for a Member State or region thereof listed in Annex I to Decision 2008/185/EC and which come from donor females complying with the conditions of Article 1 of Decision 2008/185/EC;]					
ificat		(1) ○ or	[listed in Annex II to Decision 2008/185/EC and are destined for a Member State or region thereof listed in Annex II to Decision 2008/185/EC;]					
Part II: Certification		(1) ∘ or	[not listed in Annex I or II to Decision 2008/185/EC and are destined for a Member State or region thereof listed in Annex I or II to Decision 2008/185/EC and which come from donor females complying with the conditions of Article 1 of Decision 2008/185/EC;]					
<u> </u>		(1) ∘ or	[not listed in Annex I or II to Decision 2008/185/EC and are destined for a Member State or region thereof not listed in Annex I or II to Decision 2008/185/EC;]]					
	II.5.	Chapter III(II) of Annex D to Directive 92/65/EEC and bearing the number detailed in box I.23.						
	II.6.	are dispatched:						
	-(1) either	collection team] (1) \square [embryo production team] (1) or from a zone not subject to s affecting porcine animals and established for reasons of listed diseases relevant seases subject to emergency measures relevant for those species, or those ply to these embryos because they were collected before the restrictions were have not been in contact with other germinal products of a lower health status for						
	(1) or	movement restrictions affecting porcine animals and established for (3), but derogations from movement restrictions have been granted, and:						
	(1) \square [they comply with the requirements set out in (4);]]							
	(1)	(1) \square [and in particular, they are (5).]]						
	Notes In accordance with the Agreement on the withdrawal of the United Kingdom of Great Britain and Northern Ireland from the European Union and the European Atomic Energy Community, and in particular Article 5(4) of the Protocol on Ireland/Northern Ireland in conjunction with Annex 2 to that Protocol, references to the Union in this animal health certificate include the United Kingdom in respect of Northern Ireland.							
	provided f	This animal health certificate shall be completed in accordance with 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 shal ova/embryos collectio	l correspond to the embryo collection team or embryo production team of on/production.					
	Box reference I.12:	Place of destination shall correspond to the embryo collection team, embryo production team, germinal product processing establishment, germinal product storage centre or to the establishment of ova/embryos destination.						
	Box reference I.19:	reference						
	Box reference I.26:	reference						
	Box reference	31 1 3 y						

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	II. Health info	rmation						
	I.30:							
		Identification number shall corres	pond to the official identifica	tion of the animal.				
		Date of collection shall be indicated in the following format: dd/mm/yyyy.						
on		Approval number of the team shall correspond to the embryo collection team or embryo production team of ova/embryos collection/production indicated in box I.11.						
cati	Part II:	n:						
tifi	(1)	Delete if not applicable.						
Part II: Certification	(2)	Only embryo collection or production teams approved by the competent authority and listed in accordance with Article 11(4) of Directive 92/65/EEC.						
ırt I	(3)	Insert the name of the disease(s).						
Pa	(4)	Insert the specific reference to the article(s), title, and number of the relevant legal act(s) adopted by the Commission providing for those requirements.						
	(5)	Insert the specific attestation(s) provided for in and required by the relevant legal act(s) adopted by the Commission, as referred to in Article 159(2), points (a), (b) and (c), of Regulation (EU) 2016/429 of the European Parliament and of the Council.						
	Certifying Offi Name (in capi	icer/Official veterinarian ital letters)	Qualification and title					
	Date of declar		Signature					
	Stamp							

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