INTRA

	I.1. Consignor				I.2. IMSOC reference I.2.a. Local reference			ence	
	Name Address					I.3. Central Comp	petent Authority		
	Country ISO Code					I.4. Local Compe	tent Authority		
ent	I.5. Consignee				I.6. Operator conducting assembly operations independently of an establishment				
m	Name Address			Name					
\mathbf{sig}	Country ISO Code			Address Approval Number Country ISO Code					
ons									
$^{ m of}$									
tion (I.7. Country of origin ISO Code				I.9. Country of destination	on		ISO Code	
Part I: Description of consignment	I.8. Region of origin Code				I.10. Region of destination	on		Code	
	I.11. Place of dispa	atch			I.12. Place of destination	າ			
I: I	Name				Name				
ırt	Address Approval				Address				
Ρĉ	Number				Approval Number				
	Country		ISO Code		Country		ISO Code		
	I.13. Place of loadi	ing			I.14. Date and time of departure				
	Name								
_	Approval								
		Approval Number							
	Country	Country ISO Code							
	I.15. Means of Tra		- 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		I.16. Transporter				
	Mode	International transport	Identification		Name				
		document			Address Approval				
					Number		100 0 1		
					Country		ISO Code		
					I.17. Accompanying documents				
					Document Type Accompanying document				
			reference Date of Issue Country						
					Place of issue				
	I.18. Transport conditions								
	Chilled ☐ Ambient ☐			Frozen 🗆					
	I.19. Container No / Seal No I.20. Certified as								
	Germinal products								
	I.21. For transit th	. For transit through a third country							
	Third country			ISO Code BCP code BCP code					
	Exit point Entry point								
		22. For transit through Member State(s)		I.23. For export					
	Member State			Third country	aird country ISO Code				
	L24. Estimated ion				Exit point BCP code I.25. Journey Log				
	I.26. Total number		I.27. Total quan	ntity	I.28. Total gross weight				
	I.30. Description o		1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	,		1	- 0 .		
	_	_	, NOT ELSEWHERE SPECIFIE	ED OR IN	CLUDED				
		oducts not elsewhe	re specified or included; dea			t for human (consumption		
	05119985 Other								

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_	ION	1	T	T		INIK
	#1. Commodity	Identification Number	Quantity	Nature of commodity	Identification Mark	
	Species	Package count	Date of collection / production	Plant / Establishment / Centre	Туре	
5						
tar transferred to conditions transferred						
b						
1						
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I						
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2024/1044 (2021/403) MODEL "OV/CAP-OOCYTES-EMB-C-INTRA"

	II. Health info	rmation						
uo	_ , ,							
	I, the undersigned official veterinarian, hereby certify that the \square [ova] (1) \square [embryos] (1) described in Part I:							
	II.1. were collected, processed and stored under conditions which meet the requirements of Directive 92/65/EEC;							
	II.2.	come from female donors of the \square [ovine] (1) \square [caprine] (1) species which meet the requirements of Chapter IV of Annex D to Directive 92/65/EEC;						
ficati	II.3.	are embryos of the ovine or caprine species which comply with the following conditions as regards classical scrapie:						
Certi	(1) o either	[II.3.1.	they meet the requirements of Chapter III of Annex D to Directive 92/65/EEC and of Chapter A(I) of Annex VIII to Regulation (EC) No 999/2001, as applicable on 31 August 2010;					
Part II: Certification	(1) ∘ or	[II.3.1.	they meet the requirements of Chapter III of Annex D to Directive 92/65/EEC and of Chapter A(I) of Annex VIII to Regulation (EC) No 999/2001, as applicable on 31 August 2010, and are destined for a Member State which benefits, for all or part of its territory, from the provisions laid down in point (b) or (c) of Chapter A(I) of Annex VIII to Regulation (EC) No 999/2001, as applicable on 31 August 2010 and the donor animals comply regarding scrapie with the guarantees provided for by the programmes referred to in that point and with the guarantees (2) requested by the Member State of destination;]					
	(1) o either	[II.3.2.	the semen used for fertilisation me requirements of Chapter A(I) of An 31 August 2010;]					
	(1) ∘ or	[II.3.2.	the semen used for fertilisation merequirements of Chapter A(I) of An 31 August 2010, and is destined for territory, from the provisions laid Regulation (EC) No 999/2001, as appregarding scrapie with the guarantees (2) regulation with the guarantees (2) results of the semental scrapic with the guarantees (2) results of the semental scrapic with the guarantees (2) results of the semental scrapic with the guarantees (2) results of the semental scrapic with the guarantees (2) results of the semental scrapic with the guarantees (2) results of the semental scrapic with the guarantees (2) results of the semental scrapic with the semental sc	nex VIII to Regulation (EC) No e a Member State which benef down in point (b) or (c) of Cha plicable on 31 August 2010, an tees provided for by the progr	o 999/2001, as applicable on its, for all or part of its apter A(I) of Annex VIII to and the donor animals comply cammes referred to in that			
	II.3.3.	I.3.3. are dispatched:						
	(1) ○ either	[by an □ [embryo collection team] (1) □ [embryo production team] (1) or from a zone not subject to movement restrictions affecting ovine and caprine animals and established for reasons of listed diseases relevant for those species or diseases subject to emergency measures relevant for those species, or those restrictions do not apply to these embryos because they were collected before the restrictions were established, and they have not been in contact with other ova or embryos of a lower health status for an adequate period.]						
	(1) ∘ or	movement	[embryo collection team] (1) [er restrictions affecting ovine and caps from movement restrictions have	rine animals and established	, ,			
	(1) 🗆	[they comp	oly with the requirements set out in	(4);]]				
	(1) 🗆	[and, in pa	rticular, they are (5).]]				
	Notes							
	from the Eu Protocol on	uropean Un Ireland/No	ne Agreement on the withdrawal of the United Kingdom of Great Britain and Northern Ireland nion and the European Atomic Energy Community, and in particular Article 5(4) of the orthern Ireland in conjunction with Annex 2 to that Protocol, references to the Union in this ate include the United Kingdom in respect of Northern Ireland.					
	This anima provided fo	ompletion of certificates 0/2235.						
	Part I: Box reference I.11:	Box Place of dispatch shall correspond to the embryo collection team of ova/embryos collection. reference						
	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.						

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UNION 2024/1044 (2021/403) MODEL "OV/CAP-OOCYTES-EMB-C-INTR								
	II. Health info	ormation						
	Box Seal number shall be indicated. reference I.19:							
Certification	Box reference I.26:	Total number of packages shall correspond to the number of containers.						
	Box reference I.30:	"Type": specify if: in vivo derived embryos, in vivo derived oocytes, in vitro produced embryos or micromanipulated embryos.						
I:C		Identification number shall correspond to the	official identification of the a	nimal.				
rtI		Date of collection shall be indicated in the following format: dd/mm/yyyy.						
Date of collection shall be indicated in the following format: dd/mm/yyyy. Approval number of the team shall correspond to the embryo collection team of ova/embryos indicated in box I.11.								
	Part II:							
	(1)	Delete if not applicable.						
	(2)	Additional guarantees as laid down in Article	2 of Regulation (EC) No 546/20	006 [OJ L 94, 1.4.2006, p. 28].				
	(3)	Insert the name of the disease(s).						
	(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 Officer/Official veterinarian							
	Name (in capital letters) Date of declaration Qualification and title Signature							
	Stamp							

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