EUROPEAN UNION INTRA

	I.1. Consignor		I.2. IMSOC reference I.2.a. Local reference												
	Name									I.3. Central Competent Authority					
	Address									.4. Local Competent Authority					
	Country ISO Code					In a Books component munority									
	150 code														
ب	I.5. Consignee						I.6. Operator conducting assembly operations independently of an establishment								
of consignment	Name						establishment								
Ħ	Address						Name								
텯	Country ISO Code						Address								
ß							Approval Number								
8								Country ISO Code							
Ę.	I.7. Country of orig	ISO Code	I 9 Country of	ry of destination ISO Code				ISO Code							
							1.5. Country of	i destination							
Part I: Description															
Ę.	.8. Region of origin Code						I.10. Region of								
Š	I.11. Place of dispatch						I.12. Place of destination								
ĕ	Name	Name													
Ξ	Address						Address								
ㅂ	Approval Number	r					Approval Nu	mber							
Pa	Country			ISO (Code		Country				ISO Code				
	I.13. Place of loadi	ng					I 14 Date and	time of der	artura						
		ng					I.14. Date and time of departure								
	Name														
	Address														
	Approval Number	r		***											
	Country			ISO (Code										
	I.15. Means of Tra	nsport					I.16. Transpor	ter							
	Mode														
	transport					Name Address									
		document					Activity ID Country ISO Code								
							-								
							I.17. Accompa	nying docu	ments						
							Commercial			5 .	c ·				
						document reference		Date of issue							
							Country	Dlana af			of				
						issue									
			I.18. Transport conditions												
		nditions			Frozen Chilled						Ambient □				
		nditions			Chilled \Box	_			Ambient 🗀	ı					
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					Chilled \Box				Ambient 🗀	1					
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	Frozen 🗆	/ Seal No			Chilled				Ambient 🗀						
	Frozen . I.19. Container No I.20. Certified as	/ Seal No	rd count	ry	Chilled C										
	Frozen I.19. Container No I.20. Certified as Germinal products	/ Seal No	rd count	ry	Chilled □		ISO Code								
	Frozen I.19. Container No I.20. Certified as Germinal product I.21. For transit th	/ Seal No	rd count	ry	Chilled C		ISO Code BCP code								
	Frozen I.19. Container No I.20. Certified as Germinal product: I.21. For transit th Third country	/ Seal No	rd count	ry	Chilled □										
	Frozen I.19. Container No I.20. Certified as Germinal product: I.21. For transit th Third country Exit point	/ Seal No			Chilled		BCP code								
	Frozen I.19. Container No I.20. Certified as Germinal product: I.21. For transit th Third country Exit point Entry point	/ Seal No		e(s)			BCP code BCP code	rt			□ ISO Code				
	Frozen I.19. Container No I.20. Certified as Germinal product: I.21. For transit th Third country Exit point Entry point I.22. For transit th	/ Seal No		e(s)			BCP code BCP code I.23. For expo	rt			_				
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	Frozen I.19. Container No I.20. Certified as Germinal products I.21. For transit th Third country Exit point Entry point I.22. For transit th Member State I.26. Total number	/ Seal No s rough a thin rough Mem	iber State	e(s)	Code		BCP code BCP code I.23. For export Third country Exit point I.25. Journey I	rt ,			ISO Code BCP code	modity			
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	II. Health info	ormation							
	I, the unde	ersigned off	icial veterin	arian, hereby certify that	t:				
Part II: Certification	(1) o either	[II.1.	the in vivo derived embryos/in vivo derived ova(1) described in Part I were collected, processed and stored by an embryo collection team(2) approved and supervised in accordance with Chapter I(III)(1) of Annex D to Directive 92/65/EEC;]						
	(1) ○ or	[II.1.	produced,	processed and stored by	cromanipulated embryos(1) described in Part I were an embryo production team(2), approved and supervised 1) and (2) of Annex D to Directive 92/65/EEC;]				
	(1) o either	[II.2.	the in vivo derived embryos described in Part I meet the requirements of Chapter III(II)(1) Annex D to Directive 92/65/EEC;]						
	(1) ∘ or	[II.2.	the in vivo derived ova described in Part I meet the requirements of Chapter III(II)(2) of Annex D to Directive 92/65/EEC;]						
	(1) ∘ or	[II.2.	the in vitro produced embryos described in Part I meet the requirements of Chapter III(II)(3) of Annex D to Directive 92/65/EEC;]						
	(1) ∘ or	[II.2.	the micromanipulated embryos described in Part I meet the requirements of Chapter III(II)(4) of Annex D to Directive 92/65/EEC;]						
		II.3.	the ova or	embryos described in Pa	art I come from donor mares	which:			
	_		II.3.1.	Directive 2009/156/EC(4	fulfilling the conditions laid (4) onto which only equidae s 5 or Articles 12 to 16 of Direc	atisfying the conditions laid			
			II.3.2.	meet the additional rec 92/65/EEC;	quirements of Chapter IV(4) o	of Annex D to Directive			
			II.3.3.	of collection of ova or e		east 30 days prior to the date te of the first sample referred ection of ova and embryos;			
			II.3.4.	(Coggins test) or an ELI samples taken on	on of ova or embryos and th n on (3); being				
			II.3.5.	by isolation of Taylorel out with negative resul prior to the date of the of the clitoral fossa and (3) and o	la equigenitalis after a cultiv ts in each case on samples ta first collection of ova or emk l clitoral sinuses on two cons	nken during the past 30 days oryos from mucosal surfaces secutives oestrus periods on n additional culture			
	(1) o either	[II.4.	the embryos described in Part I were conceived as a result of artificial insemination of donor mares with semen which was collected, processed, stored and transported under conditions which comply with the requirements of Chapters I(I), II(I) and III(I) of Annex Directive 92/65/EEC;]						
	(1) ∘ or	[II.4.	the embryos described in Part I were conceived as a result of in vitro fertilisation of ova complying with the conditions in point 2 of Chapter III(II) of Annex D to Directive 92/65/EEC with semen which was collected, processed, stored and transported under conditions which comply with the requirements of Chapters I(I), II(I) and III(I) of Annex D to Directive 92/65/EEC;]						
	(1) or	[II.4.	the ova ha	ive not been in contact w	ith semen of the equine spec	ries;]			
		II.5.	in accorda	-	art I were sent to the place of loading in a sealed container ter III(II) of Annex D to Directive 92/65/EEC and bearing				

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EUROPEAN UNION

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: Part II: Certification Box I.11: Place of dispatch shall correspond to the embryo collection team or embryo production team of ova/embryos collection/production. Box I.12: Place of destination shall correspond to the embryo collection team, embryo production team or to the holding of ova/embryos destination. Box I.19: Identification of container and Seal number shall be indicated. "Type": Specify if: in vivo derived embryos, in vivo derived oocytes, in vitro produced embryos or Box I.30: micromanipulated embryos. Donor identity shall correspond to the official identification of the animal. 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 or embryo production team of ova/embryos collection/production. Part II: (1)Delete as appropriate. (2) Only embryo collection or production teams approved by the competent authority and listed in accordance with Article 11(4) of Council Directive 92/65/EEC. (3)Insert date. (4) OJ L 192, 23.7.2010, p. 1. Certifying Officer/Official veterinarian Name (in capital letters) Authority name Date of signature Signature Stamp