EUROPEAN UNION INTRA

	I.1. Consignor				I.2. IMSOC ref	erence	I.2.a. Local reference	
	_						I.3. Central Competent Authority	
	Name							
	Address						I.4. Local Competent Authority	
	Country	ountry ISO Code						
	,							
	I.5. Consignee				I.6. Operator conducting assembly operations independently of an			
ᆵ	_	-				establishment		
1 9	Name				Nama			
ਖ਼	Address				Name			
g	Country		ISO Code		Address			
SI	•				Approval Nu	mber		
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—	I.8. Region of origin			Code	I.10. Region of		Code	
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es	Name				Name			
Ă۱								
∷ ∣	Address				Address			
اب	Approval Number	r			Approval Nu	mber		
aı	Country		ISO Code		Country		ISO Code	
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	I.13. Place of loadi	ng			I.14. Date and	time of departure		
		0				or acparture		
	Name							
	Address							
	Approval Number	r						
		L	100 0 1					
_	Country		ISO Code					
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	I.15. Means of Tran	nsport			I.16. Transpor	ter		
	Mode	International	Identification		Name			
		transport			Address			
		document						
					Approval Nu	mber		
					Country		ISO Code	
					I.17. Accompa	nying documents		
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ealth information						
ie undersigned	official veterin	arian, hereby certify that	t:			
	men collection for trade:	centre(1), in which the se	emen described in Part I was	collected, processed and		
II.1.1. was approved and supervised by t Chapter I of Annex D to Directive 9			he competent authority according to the conditions of 02/65/EEC;			
II.1.2.	Member S dispatche frozen ser	tate which was on the da d as fresh/chilled(2) seme nen elapsed(2) not consid	the case of regionalisation in a part of the territory(2) of a y semen was collected until the date the semen was n or until the 30 days mandatory storage period for lered to be infected with African horse sickness in (b) of Directive 2009/156/EC(3);			
II.1.3.	the date th	ne semen was dispatched eriod for frozen semen el	ncing 30 days prior to the dat as fresh/chilled(2) semen or apsed(2), the conditions of A	until the 30 days mandatory		
II.1.4.	the date th storage pe	ne semen was dispatched	as fresh/chilled(2) semen or apsed(2) only equidae which	date of semen collection until runtil the 30 days mandatory nwere free of clinical signs of		
II.2. All equidae have been admitted onto the centre under the provisions of Article 4 and 5 of Directive 2009/156/EC(3);						
The se	men described	in Part I was collected fro	om donor stallions, which:			
II.3.1.	on the day contagiou		l have not shown clinical sig	ns of an infectious or		
II.3.2.	during at service,	least 30 days prior to coll	ection of the semen have no	t been used for natural		
II.3.3.	_	e 30 day period prior to c e showed clinical signs o		been kept on holdings where		
II.3.4.			collection of the semen have f contagious equine metritis,	been kept on holdings where		
 II.3.5. to the best of my knowledge and as equidae suffering from an infectio preceding collection of the semen, II.3.6. have undergone the following anim the competent authority, in according to the semental process. 						
				, , ,		
	II.3.6.1.	an agar-gel immuno-di with negative result;	ffusion test (Coggins test) for	equine infectious anaemia		
(2) o either	[II.3.6.2.	a serum neutralisation serum dilution of one i	test for equine viral arteritisn four; and]	s with negative result at a		
(2) ○ 0	r [II.3.6.2.		r equine viral arteritis carrie semen of the donor stallion	ed out with negative result on ;]		
	II.3.6.3.	occasions on samples c days by isolation of Tay semen sample and fror	ollected from the donor stall ylorella equigenitalis from p n genital swabs taken at leas	lion with an interval of seven re-ejaculatory fluid or a st from the penile sheath,		
II.3.7.	have been	subject to the one of the	following test programmes((4):		
	II.3.7.1.	30 days prior to the sen	nen collection, and during th on centre came during that t	ne collection period, and no ime into direct contact with		
	II.3.7.	II.3.7. have been	occasions on samples of days by isolation of Tay semen sample and from urethra and urethral for II.3.7. have been subject to the one of the II.3.7.1. The donor stallion was 30 days prior to the ser equidae on the collection	occasions on samples collected from the donor stall days by isolation of Taylorella equigenitalis from posemen sample and from genital swabs taken at least urethra and urethral fossa with negative result in EII.3.7. have been subject to the one of the following test programmes (

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d		II.3.7.2.	The tests described in point II.3.6. have been carried out on samples taken on				
		II.3.7.2.	other equidae on the collection centre came into contact with equidae of lower health status than the donor stallion. The tests described in point II.3.6. have been carried out on samples taken on				
			(5) and in the case of contagious equine metritis on a second				
d			the test described in point II.3.6.1. for equine infectious anaemia was last carried out on a sample of blood taken on (5), being not more than 120 days before the semen described in Part I was collected				
	(2)	o either	[one of the tests described in point II.3.6.2. for equine viral arteritis was last carried out on a sample collected on (5), being not more that days before the semen described in Part I was collected,]				
	(2)	o or	[the non-shedder state of the seropositive stallion for equine viral arteritis was confirmed by a virus isolation test which was carried out on an aliquot of the entire semen of the donor stallion collected on (5), being not more than one year before the semen described in Part I was collected;]				
		II.3.7.3.	The tests described in point II.3.6. have been carried out during the 30 days mandatory storage period of frozen semen and not less than 14 days after the collection of the semen on samples taken on				
The semen described in Part I was collected, processed, stored and transported under conditions which comply with the requirements of Chapters II and III of Annex D to Directive 92/65/EEC.							
tos							
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:							
x I.11:	: place of dispatch shall correspond to the semen collection centre of origin of the semen.						
x I.12:	semen destination.						
x I.19:							
x I.30:	0: donor identity shall correspond to the official identification of the animal.						
			ll be indicated in the following format: dd/mm/yyyy.				
	approval number of the centre shall correspond to the approval number of the semen centre indicated in Box I.11. where the semen was collected.						
	art II:						
rt II:	1) Only semen collection centres approved by the competent authority and listed in accordance with Article 11(4) of Council Directive 92/65/EEC.						
rt II:		Delete as appropriate.					
rt II:	Article 1	s appropriate.					
rt II:	Article 1 Delete as	s appropriate. , 23.7.2010, p.					
rt II:	Article 1 Delete as OJ L 192	, 23.7.2010, p.					
	II:	Only sen	Only semen collection				

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	II. Health information		
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	Name (in capital letters)	Qualification and title	
	Date of signature	Signature	
	Stamp		
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Part II: Certification			
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