EUROPEAN UNION INTRA

	I.1. Consignor Name					I.2. IMSOC reference I.2.a. Local reference				
							I.3. Central Competent Authority			
	Address					I.4. Local Competent Authority				
	Country		ISO Code							
	·									
	I.5. Consignee				I.6. Operator of	onducting assembly o	operations independently of an			
Part I: Description of consignment	-				establishment		- p			
<u>e</u>	Name				Nama					
莒	Address				Name					
ם	Country		ISO Code		Address					
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∺	Address				Address					
ب	Approval Number	r			Approval Number					
ਰ	Country		ISO Code		Country		ISO Code			
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	I.13. Place of loadi	ng			I.14. Date and	time of departure				
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	Name									
	Address									
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	Country		ISO Code							
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	I.15. Means of Trai	nsport			I.16. Transpor	ter				
	Mode	International	Identification		Name					
		transport			Address					
		document			Approval Number					
					Country ISO Code					
					I.17. Accompa	nying documents				
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	Chilled I.19. Container No I.20. Certified as Germinal products	/ Seal No	5		accompanyi ng.documen t.document. number		Place of issue			
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	II. Health information							
	I, the undersigned official veterinarian, hereby certify that:							
	processed and stored, for trade wa				a which the semen described in Part I was collected, as approved and supervised by the competent authority in ad Chapter I(II)(1) of Annex D to Directive 92/65/EEC;			
Part II: Certification		II.1.1.	semen des dispatched	cribed in Par	rt I until the date the fresh or 30 days storage period for fr			
			II.1.1.1.	part of the to be infect		te which was not considered ss in accordance with Article		
			II.1.1.2.		conditions for a holding laid 009/156/EC(3);	l down in Article 4(5) of		
			II.1.1.3		only equidae which were free tis and contagious equine me			
	II.2.		-	-	ons laid down in Articles 4 a admitted onto the centre.	nd 5 or Articles 12 to 16 of		
	II.3.	The semen	described i	in Part I was	collected from donor stallion	ıs, which:		
		II.3.1.			nical sign of an infectious or the centre and on the day th			
		II.3.2.	where no	equine has sl	lays prior to the date of seme nown any clinical sign of equ ritis during that period;			
		II.3.3.	first semen	n collection a		ast 30 days prior to the date of t sample referred to in points ction period;		
		II.3.4.	relevant Cl Animals of	hapter of the f the OIE, car	Manual of Diagnostic Tests a ried out on samples taken in			
	(2)	o either	[II.3.4.1.		immuno-diffusion test (Cogg IA) with negative result;]	gins test) for equine infectious		
	(2)	\circ or	[II.3.4.1.	an ELISA fo	or equine infectious anaemia	(EIA) with negative result;]		
	and (2)	o either	[II.3.4.2.		utralisation test for equine v sult at a serum dilution of on			
	(2)	o or	[II.3.4.2.		ation test for equine viral art sult on an aliquot of the enti	reritis (EVA) carried out with re semen of the donor		
	and		II.3.4.3.	carried out seven days of 7 to 14 da from genita	ays from pre-ejaculatory flui	s taken with an interval of uigenitalis after a cultivation d or a semen sample and he penile sheath, urethra and		
		II.3.5.			ith the results specified in II.: umes(4) detailed in points II.3			

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	II. Health information							
			II.3.5.1.	centre for a during the no equidae	stallion was continuously rest at least 30 days prior to the da period of collection of the sen on the semen collection cent ae of lower health status than	te of the first collection and nen described in Part I and re came into direct contact		
Part II: Certification			taken(5)pri	described in point II.3.4. have been carried out on samples prior to the first semen collection and at least 14 days the date of the commencement of the residence period of 0 days.				
Part II: Ce			II.3.5.2.	least 30 day period of co centre und continuous	stallion was resident on the so ys prior to the date of the first ollection of the semen describ er the responsibility of the ces speriod of less than 14 days, a centre came into direct contactus.	collection and during the red in Part I, but has left the ntre veterinarian for a nd/or other equidae on the		
				taken(5)pri collection p collected a	escribed in point II.3.4. have be for to the first semen collection period in the year the semen of and at least 14 days following t ment of the residence period	n of the breeding season or lescribed in Part I was he date of the		
	and			last carried	scribed in point II.3.4.1. for eq l out on a sample of blood tak semen described in Part I was	en(5) not more than 90 days		
	and	(2)	o either	was last ca	tests described in point II.3.4 rried out on a sample taken(5 semen described above was c) not more than 30 days		
		(2)	o or	negative re stallion tak described i same date(olation test for equine viral ar esult on an aliquot of the entir en(5) not more than six mont n Part I was collected and a b 5) reacted positive in a serum al arteritis at a serum dilution	e semen of the donor hs before the semen lood sample taken on the neutralisation test for		
	and)	last carried	scribed in point II.3.4.3. for co l out on samples taken(5)not r cribed in Part I was collected.			
			II.3.5.3.	taken(5)pri	escribed in point II.3.4. have be for to the first semen collection period in the year the semen of	n of the breeding season or		
	and			taken(5) no	escribed in point II.3.4. were la otless than 14 days and not mo of the semen described in Part	ore than 90 days after the		
		II.3.6.	have unde following	-	sting provided for in point II.S	3.5. on samples taken on the		

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	II. Health info	rmation								
		Identificat ion of semen	Test programm e	Start date(5)		Date of sar	npling for he	ealth tests(5)	
				Donor residence	Semen collection	EIA II.3.4.1.	EVA II.3.4.2.		CEM II.3.4.	3.
Part II: Certification							Blood sample	Semen sample	1. sample	2. Sample
l: Certi				_	_	_		_	_	_
Part I										_
		_	_	_	_					_
		_	_		_		2			_
		_	_		_					
				(
	(2) o either	[II.4.	No antibiot	ics were ad	ded to the s	— emen;]	_		_	
	(2) ○ or	[II.4.					tibiotics was		roduce a coi	ncentration
		II.5.	The semen II.5.1.		rocessed, st	ored and tr	ansported u			
			II.5.2.		(I) of Annex		aled contain ive 92/65/EE			

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II. Health information		
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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 semen collection centre of origin of the semen.

Box I.12: Place of destination shall correspond to the semen collection or storage centre or to the holding of

semen destination.

Box I.19: Identification of container and seal number shall be indicated.

Box I.30: 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 centre shall correspond to the approval number of the semen centre indicated

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Identificat Test

Part II:

Guidance for the completion of Table in II.3.6:

Abbreviations:

EIA-1 Equine infectious anaemia (EIA) testing first occasion

EIA-2 EIA testing second occasion

EVA-B1 Equine viral arteritis (EVA) testing on blood sample first occasion

EVA-B2 EVA testing on blood sample second occasion

EVA-S1 EVA testing on semen sample first occasion

EVA-S2 EVA testing on semen sample second occasion

CEM-11 Contagious equine metritis (CEM) testing first occasion first sample

CEM-12 CEM testing first occasion second sample taken 7 days after CEM-11

CEM-21 CEM testing second occasion first sample

Start

CEM-22 CEM testing second occasion second sample taken 7 days after CEM-21

Instructions:

For each semen identification in column A in the example below, the test programme (II.3.5.1., II.3.5.2. and/or II.3.5.3.) must be described in column B and columns C and D must be completed with the dates required.

The dates when samples were taken for laboratory testing prior to the first collection of the semen described in Part I as required in II.3.5.1., II.3.5.2. and II.3.5.3., are entered in the upper line of columns 5 to 9 of the table, this being the boxes marked with EIA-1, EVA-B1 or EVA-S1 and CEM-11 and CEM-12 in the example below.

The dates when samples were taken for repeat laboratory testing as required in accordance with II.3.5.2. or II.3.5.3. are entered in the lower line of columns 5 to 9 in table, this being the boxes EIA-2, EVA-B2 or EVA-S2 and CEM-21 and CEM-22 in the example below.

ion of semen	programm e	date(5)						
		Donor residence	Semen collection	EIA II.3.4.1.	EVA II.3.4.2.		CEM II.3.4.	3.
					Blood sample	Semen sample	1. sample	2. sample
A	В	С	D	EIA-1	EVA-B1	EVA-S1	CEM-11	CEM-12
				EIA-2	EVA-B2	EVA-S2	CEM-21	CEM-22

Date of sampling for health tests(5)

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EUROPEAN UNION 2021/403 EQUI-SEM-C-INTRA II. Health information (1) Only semen collection centres approved by the competent authority and listed in accordance with Article 11(4) of Council Directive 92/65/EEC. (2) Delete as appropriate. (3) OJ L 192, 23.7.2010, p. 1. (4) Cross out the programme(s) that do(es) not apply to the consignment. Part II: Certification (5) Insert date in table in point II.3.6 (follow Guidance in Part II of the Notes). (6) Insert names and concentrations. Certifying Officer/Official veterinarian Name (in capital letters) Qualification and title Date of signature Signature Stamp

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