EUROPEAN UNION INTRA

	I.1. Consignor					I.2. IMSOC ref	erence	I.2.a. Local reference	
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	Name							I.3. Central Competent Au	
	Address							I.4. Local Competent Auth	iority
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≓ ∣	I.5. Consignee					establishment	conducting assembly of	perations independently of	an
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0	I.7. Country of orig	gin			ISO Code	I.9. Country of	destination	ISC	O Code
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Part I: Description of consignment									
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	I.13. Place of loadi	ng				I.14. Date and	time of departure		
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	I.15. Means of Trai	nsport				I.16. Transpor	ter		
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ΕU	ROPEAN UNION					(2021/403	B) EQUI-SEM-A-INTRA
	II. Health information						
	(2)	o or	period of t	the preceding	emia has been reported og 12 months prior to colle stablishment has remain	ection of th	ne semen and following
Part II: Certification		(2)	∘ either	subjected to method pro (EU) 2020/6 on two occa infected an	emaining equine animale o a test for equine infection ovided for in Part 9 of An 88, carried out, with neg asions with a minimum i imals have been killed a hment was cleaned and	ous anaen nex I to Do ative resul nterval of nd destroy	nia with the diagnostic elegated Regulation lts, on samples taken 3 months after the red or slaughtered and
Part II:		(2)	o or	was either l	30 days after the last eq killed and destroyed or s ed and disinfected;]]		
		II.2.2.4.	no equine	animal has s	riod of 30 days prior to t hown signs of infection ritis (Taylorella equigeni	with equir	
	II.2.3.				signs of transmissible ar		
	II.2.4.		fied as provi n (EU) 2019/2		rticle 58(1), 59(1) or 62(1)	of Commi	ssion Delegated
	II.2.5.		od of at least ion period	30 days prio	r to the date of first colle	ection of th	e semen and during
		II.2.5.1.	the occurr	ence of Afric	ments not situated in a re an horse sickness, infect rging disease relevant fo	ion with B	urkholderia mallei
		II.2.5.2.	encephalo anaemia, i	myelitis, dou infection with	stablishment where Ven rine, surra (Trypanosom n equine arteritis virus, o is), infection with rabies	na evansi), contagious	equine infections equine metritis
		II.2.5.3.	zone due t	to the occurre	h animals from establish ence of diseases referred lo not meet the condition	to in poin	t II.2.5.1. or from
	II.2.6.	first seme	n collection	and between	g during a period of at lea the dates of the first sar e end of the collection pe	nple referi	-
	II.2.7.				ng tests, referred to in po EU) 2020/686, as follows:		Chapter I of Part 4 of
		II.2.7.1.	diffusion t	_	ne infectious anaemia (E Coggins test) or an enzyı e result;	_	_
		II.2.7.2.	for infecti	on with equi	ne arteritis virus (EVA),		
	(2)	□ either	[II.2.7.2.1.	a serum ne of one in fo	utralisation test with a n ur;]	egative re	sult at a serum dilution
	(2)	□ and/or	[II.2.7.2.2.		ation test, polymerase ch negative result on an ali on;]		
		II.2.7.3.	identificat stallion or	ion test carri two occasion	netritis (Taylorella equiged out on three speciments with an interval of nouce), the urethra and the	ns (swabs) t less than	taken from the donor 7 days at least from

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 50	ROPEAN UNION				(202	21/403) EQUI-SEM-A-INTRA
	II. Health information					
			days (local were place medium, k	l treatment) a ed in transpo	fter antimicrobial treatmen rt medium with activated ch ch to the laboratory where th	arcoal, such as Amies
Part II: Certification	(2)	□ either	[II.2.7.3.1.	microaerop within the 2	he 48 hour period where the	of at least 7 days, set up ne specimens from the donor
Part II:	(2)	□ and/or	[II.2.7.3.2.	time PCR, ca	n of genome of Taylorella eq arried out within the 48 hou from the donor animal;]	
	II.2.8.	following t	esting prog	rammes deta	cified in point II.2.7. in each iled respectively in points 1(egulation (EU) 2020/686:	case to at least one of the b)(i), (ii) and (iii) of Chapter I
	(3)	□ [II.2.8.1.	period of a period of of the semen equine an described stallion at the first co fresh, chill commence semen col	at least 30 day collection of to collection ce imals of lowe in point II.2.7 least once a y ollection of se led or frozen ement of the b lection.]	ys prior to the date of the first he semen described in Part I ntre came during that time is repeated in the done. We were carried out on sample year at the beginning of the lamen intended for movement semen and not less than 14 or residence period of at least 3 or seminary.	, and no equine animals in nto direct contact with or stallion. The tests es taken(4) from the donor preeding season or prior to to another Member State as days following the date of the 0 days prior to the first
	(3)	[II.2.8.2.	least 30 da collection under the less than 1 semen coll health stat taken(4) for breeding s movement less than 1 period of a period of a State as fr	of the semen responsibility 4 days during lection centre tus. The tests rom the dono season or price to another Mays follow at least 30 days collection of tesh, chilled on	te date of the first collection described in Part I, but left to yof the centre veterinarian is get the collection period, or other came into direct contact will described in point II.2.7. were stallion at least once a year	the semen collection centre for a continuous period of the equine animals in the th equine animals of a lower the carried out on samples that the beginning of the tection of semen intended for the or frozen semen and not the cement of the residence the cities of the residence the continuous and during the tement to another Member
			(a)	II.2.7.1. was	nfectious anaemia, one of the last carried out on a sample sprior to the collection of the	-
			(b)	•	n with equine arteritis virus,	
		(2)	o either	[in point II.	2.7.2. was last carried out on s prior to the date of the coll	a sample taken(4) not more

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Ε	UROPEAN UNION		(2021/403) EQUI-SEM-A-INTRA
	II. Health information		
2.00	(2)	or	[in point II.2.7.2.2., in case the non-shedder state of a donor stallion seropositive for infection with equine arteritis virus is confirmed, was carried out on an aliquot of the entire semen of the donor stallion taken(4) not more than 6 months prior to the date of the collection of the semen described in Part I and a blood sample taken(4) from the donor stallion during the 6 months period reacted with a positive result in a serum neutralisation test for infection with equine arteritis virus at a serum dilution of more than one in four;]
11. O. 11.		(c)	for contagious equine metritis, the test described in point II.2.7.3. was last carried out on three specimens (swabs) taken(4) not more than 60 days prior to the date of the collection of semen described in Part I
	(2)	o either	[on two occasions;]
	(2)	\circ or	[on a single occasion and subjected to a PCR or real-time PCR.]]
	(3)	Chapter I	r stallion does not meet the conditions set out in points 1(b)(i) and (ii) of of Part 4 of Annex II to Delegated Regulation (EU) 2020/686 and the collected for movement to another Member State as frozen semen.
		samples to the breed carried or period of collection collection	described in points II.2.7.1, II.2.7.2 and II.2.7.3 were carried out on aken(4) from the donor stallion at least once a year at the beginning of ing season, and the tests described in points II.2.7.1 and II.2.7.3. were at on samples taken(4) from the donor stallion during the storage the semen of a minimum period of 30 days from the date of the of the semen and before the semen is removed from the semen centre, not less than 14 days and not more than 90 days after the of the semen described in Part I, and
	(2) ○ either	carried ou minimum before the than 14 da	for infection with equine arteritis virus described in point II.2.7.2. were at on samples taken(4) during the storage period of the semen of a period of 30 days from the date of the collection of the semen and e semen is removed from the semen collection centre or used, not less ays and not more than 90 days after the date of the collection of the scribed in Part I.]
	(2) or	arteritis v out with a donor stal donor stal	shedder state of a donor stallion seropositive for infection with equine cirus was confirmed by virus isolation test, PCR or real-time PCR carried a negative result on samples of an aliquot of the entire semen of the llion taken(4) twice a year at an interval of at least 4 months and the llion has reacted with a positive result at a serum dilution of at least ar in a serum neutralisation test for infection with equine arteritis
	II.2.9. underwen	t the testing	g provided for in point II.2.8. on samples taken on the following dates:

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EUROPEAN UNION			(2021/403) EQUI-SEM-A-INTRA							
	II. Health info	rmation								
		Identificat ion of semen	Test programm e	Start date(4)		Date of sar	npling for h	ealth tests((4)	
				Donor residence	Semen collection	EIA II.2.7.1.	EVA II.2.7.2.		CEM II.2.7.	3.
ication							Blood sample	Semen sample	1. sample	2. sample
Part II: Certification										
Part I										
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E	UROPEA	N UNION	(2021/403) EQUI-SEM-A-INTRA
	II. Health	information	
	II.3.	The semer	described in Part I
		II.3.1.	has been collected, processed and stored in accordance with animal health requirements set out in points 1 and 2 of Part 1 of Annex III to Delegated Regulation (EU) 2020/686;
		II.3.2.	are placed in straws or other packages on which the mark is applied in accordance with requirements provided for in Article 10 of Delegated Regulation (EU) 2020/686 and that mark is indicated in Box I.30;
1	[2	II.3.3.	is transported in a container which:
T. Coutification			II.3.3.1. was sealed and numbered prior to the dispatch from the semen collection centre under responsibility of the centre veterinarian, or by an official veterinarian, and the seal bears the number as indicated in Box I.19;
1	rait		II.3.3.2. has been cleaned and either disinfected or sterilised before use, or is single-use container;
		(2)(5)	III.3.3.3. has been filled in with the cryogenic agent which not have been previously used for other products.]
	(2)(6) □ [II.4.	The semer	n is preserved by the addition of antibiotics as follows:
	[111.4.	II.4.1.	The following antibiotic or mixture of antibiotics has been added to the semen after final dilution, or is contained in the used semen diluents, to reach the indicated concentration per ml of semen:
	(2)	o either	[a mixture of gentamicin (250 μ g), tylosin (50 μ g) and lincomycin-spectinomycin (150/300 μ g);]
	(2)	o or	[a mixture of lincomycin-spectinomycin (150/300 μg), penicillin (500 IU) and streptomycin (500 μg);]
	(2)	\circ or	[a mixture of amikacin (75 μg) and divekacin (25 μg);]
	(2)	o or	[an antibiotic or a mixture of antibiotics(7), with a bactericidal activity at least equivalent to one of the following mixtures:
			- gentamicin (250 μg), tylosin (50 μg) and lincomycin-spectinomycin (150/300 μg);
			- lincomycin-spectinomycin (150/300 μg), penicillin (500 IU) and streptomycin (500 μg);
		II 4 2	- amikacin (75 μg) and divekacin (25 μg).]
		II.4.2.	Immediately after the addition of the antibiotics, and before any possible freezing, the diluted semen was kept at a temperature of at least 5°C for a period of not less than 45 minutes, or under a time-temperature regime with a documented equivalent bactericidal activity.]

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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:

Certification Box reference

"Place of dispatch": Indicate the unique approval number and the name and address of the semen

collection centre of dispatch of the consignment of semen.

I.11:

Box reference

"Place of destination": Indicate the address and unique registration or approval number of the

establishment of destination of the consignment of semen.

I.12:

Part]

Box

Seal number shall be indicated.

"Type": semen.

reference I.19:

Box

Total number of packages shall correspond to the number of containers.

reference

I.26: Box

reference

I.30:

"Identification number": Indicate identification number of each donor animal.

"Identification mark": Indicate mark on the straw or other packages where semen of the consignment is placed.

"Date of collection/production": Indicate the date on which semen of the consignment was collected.

"Approval or registration number of plant/establishment/centre": Indicate the unique approval number of the semen collection centre where the semen was collected.

"Quantity": Indicate number of straws or other packages with the same mark.

Part II:

Guidance for the completion of the table in point II.2.9.

Abbreviations:

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

EIA-2 EIA testing second occasion

EVA-B1 Equine arteritis virus (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 testing second occasion first sample CEM-21

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

Instructions:

For each semen identified in column A in correspondence with Box I.30, the test programme (points II.2.8.1., II.2.8.2. and/or II.2.8.3.) shall be specified in column B, and columns C and D shall 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 points II.2.8.1., II.2.8.2. and II.2.8.3., shall be 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.

	II. Health inf	formation									
		II.2.8.2. or	II.2.8.3. shal	l be entered		r line of co	lumns 5 to 9		accordance is being the		
น		Identificat ion of semen		Start date		_	mpling for	health tests			
псапо				Donor residence	Semen collection	EIA II.2.7.1.	EVA II.2.7.2.		CEM II.2.7	.3.	
Part II: Certification							Blood sample	Semen sample	1. sample	2. sample	
IT I		A	В	С	D	EIA-1	EVA-B1	EVA-S1	CEM-11	CEM-12	
Ьa						EIA-2	EVA-B2	EVA-S2	CEM-21	CEM-22	
	(1)								ed in the regi gated Regula		
	(2)	Delete if no	ot applicable).							
	(3)	Cross out t	he program	mes that do	not apply to	the consig	gnment.				
	(4)	Insert date in table in point II.2.9. (follow Guidance in Part II of the Notes).									
	(5)	Applicable for frozen semen.									
	(6)	Mandatory attestation in case antibiotics were added.									
	(7)	semen dilu	ent contain			nd its(their) concentra	tion or the c	commercial r	name of the	
		fficer/Official vet apital letters) nature	erinarian			Qualificatio Signature	n and title				