								1111111		
	I.1. Consignor				I.2. IMSOC ref	erence	I.2.a. Local reference			
	Name				I.3. Central Competent Auth					
	Address				I.4. Local Competent Authority					
	Country		ISO Code							
-	1.5. 0				10.0					
님	I.5. Consignee				establishmen	conducting assembly of	perations independen	tly of an		
Part I: Description of consignment	Name									
2	Address				Name Address					
50	Country		ISO Code		Approval Nu:	mhor				
ns					Country	linder	ISO Code			
ខ					country		150 Coue			
Ч	I.7. Country of orig	gin		ISO Code	I.9. Country of	f destination		ISO Code		
g										
E I					140 D					
음	I.8. Region of origi			Code	I.10. Region of			Code		
S	I.11. Place of dispa	itch			I.12. Place of o	lestination				
ĕ	Name				Name					
$\overline{\cdot}$	Address				Address					
ť	Approval Number	r			Approval Nu	mber				
Pai	Country		ISO Code		Country		ISO Code			
ŀ	I 12 Dloop of last	nď			114 Data 3	time of department				
	I.13. Place of loadi	пg			1.14. Date and	time of departure				
	Name					<b>A</b>				
	Address									
	Approval Number	r								
	Country		ISO Code							
-	I.15. Means of Trai	nsnort			I.16. Transpor	ter				
		1	T. 1							
	Mode	International transport	Identification		Name					
	document				Address					
					Approval Number					
					Country ISO Code					
					L17. Accompa	nying documents				
					[en] accompanyi ng.documen Date of issue t.document. number Country Place of					
-					Ambient					
	I.18. Transport cor	nditions		_						
	Chilled 📙		Frozen L							
-	140.0	(C 1) .								
	I.19. Container No	/ Seal No								
ŀ	I.20. Certified as									
	Germinal products	. П								
	Germinal producta	, <b>ш</b>								
ļ	I.21. For transit th	rough a third cour	ntry							
	Third country	-			ISO Code					
	Exit point				BCP code					
	Entry point				BCP code					
	I.22. For transit th	rough Member Sta	ite(s)		I.23. For expo	rt				
	Member State	0	ISO Code		Third country		ISO Code			
					Exit point		BCP code			
	I.24. Estimated jou	rney time			I.25. Journey	Log				
	,	-								
╞	1.00 Tetel	f l			107 5-1-1					
	I.26. Total number				I.27. Total qua	unuty				
	I.28. Total gross we	eigin								
ŀ	I.30. Description of	f consignment			1					
		-	26	Idoptification	Number	Quantity	Noture of com	amodity		
	Commodity	Specie	28	Identification	numper	Quantity	Nature of cor	innoany		
			1				I			
	Identification Mar	rk	Package count		Date of collect	tion / production	Plant / Establishmen	t / Centre		
ſ										

## 2021/403 EQUI-SEM-B-INTRA

_	UKOI LAI						2021/ <del>1</del> 03 LQ01-3LM-D-1	ITIGI	
	II. Health i	nformation							
	I, the un	dersigned off	icial veterina	arian, hereb	y certify tha	t:			
	II.1.	stored, for		pproved an	emen described in Part I wa l by the competent authorit 5/EEC(2);				
Part II: Certification		II.1.1.	described	in Part I unt	il the date th	days prior to the date of first collection of the semen ne fresh or chilled semen was dispatched or until the 30 frozen semen elapsed, the semen collection centre:			
			II.1.1.1.	territory(3	) of a Memborse sickness	ritory or in the case of regi er State which was not cons in accordance with Article	sidered to be infected wit	h	
			II.1.1.2.	fulfilled th 2009/156/E		for a holding laid down in	Article 4(5) of Directive		
			II.1.1.3.		only equidae gious equine	e which were free of clinica metritis;	al signs of equine viral art	eritis	
	II.2.		dae satisfyin EC have beer			wn in Articles 4 and 5 or Ar re.	ticles 12 to 16 of Directive	e	
	II.3.	The seme	n described i	n Part I was	s collected fr	om donor stallions, which:			
	-	II.3.1.				n infectious or contagious d nd on the day the semen w		ission	
		II.3.2.		wed any cli		rior to the date of semen co equine viral arteritis or co		re no	
		II.3.3.	semen coll	ection and f	from the dat	luring a period of at least 3 es of the first sample referr ction period;			
		II.3.4.	Manual of laboratory hereinafte	Diagnostic 7 which is re	Fests and Va cognised by n its accredit	t least the requirements of ccines for Terrestrial Anim the competent authority ar ation in accordance with A	als of the OIE, carried out nd has the tests referred t	t in a o	
			II.3.4.1.	Coggins tes	st) or an enz	naemia (EIA), an agar-gel iı yme-linked immunosorben h a negative result;			
			II.3.4.2.	for equine	viral arterit	is (EVA),			
		(3)	□ either	[II.3.4.2.1.	a serum ne of one in fo	utralisation test with a neg ur;]	ative result at a serum di	lution	
		(3)	□ and/or	[II.3.4.2.2.		ation test, polymerase chai negative result on an aliqu ion;]			
			II.3.4.3.	three speci interval of	imens (swab	metritis (CEM), an agent ide s) taken from the donor sta n 7 days at least from the p clandis;	allion on two occasions w	ith an	
				days (local were place medium, b	treatment) and treatment to the treatment of the treatmen	o case taken earlier than 7 after antimicrobial treatme rt medium with activated o ch to the laboratory where t for:	ent of the donor stallion a charcoal, such as Amies	nd	

	II. Health information									
L L	(3)	□ either	er [II.3.4.3.1. the isolation of Taylorella equigenitalis after cultivation un microaerophilic conditions for at least 7 days, set up within hour period after taking the specimens from the donor and the 48 hour period where the specimens are kept cool dur transport;]							
ication	(3)	□ and/or	[II.3.4.3.2.	time PCR, c	tion of genome of Taylorella equigenitalis by PCR or real- , carried out within the 48 hour period after taking the ns from the donor animal;]					
Part II: Certification	II.3.5.	programm	ected with the results specified in point II.3.4. in each case to at least one of the nes detailed in points II.3.5.1., II.3.5.2. and II.3.5.3., as follows:							
	(6)	⊔ [II.3.5.1.	The donor stallion was continuously resident on the semen collection centre for a period of at least 30 days prior to the date of the first collection and during the period of collection of the semen described above and no equidae on the semen collection centre came into direct contact with equidae of lower health status than the donor stallion.							
			donor stall prior to th semen and	lion at least o e first collect l not less tha	point II.3.4. were carried out o once a year at the beginning o tion of semen intended for tra n 14 days following the date o least 30 days prior to the date	f the breeding season or de in fresh, chilled or frozen f the commencement of the				
	(6)	□ [II.3.5.2.	least 30 da collection responsibi days, and/	ys prior to th of the semen lity of the ce or other equ	resident on the semen collect ne date of the first collection a described in Part I, but has le ntre veterinarian for a contin idae on the semen collection of f lower health status.	nd during the period of eft the centre under the uous period of less than 14				
			donor stall prior to th semen and	lion at least o e first collect l not less tha	point II.3.4. were carried out o once a year at the beginning o tion of semen intended for tra n 14 days following the date o least 30 days prior to the date	f the breeding season or de in fresh, chilled or frozen f the commencement of the				
		and		semen the do	llection of the semen intended onor stallion was subjected to					
			(a)	II.3.4.1. was	infectious anaemia, one of the s last carried out on a sample ys prior to the date of the colle n Part I;	of blood taken(7) not more				
			(b)	for equine	viral arteritis:					
		(3)	○ either	sample tak	tests described in point II.3.4. en(7) not more than 30 days p of the semen described in Part	rior to the date of the				
		(3)	∘ or	aliquot of t than six mo described i stallion du a serum ne	tests described in point II.3.4. he entire semen of the donor souths prior to the date of the constant of the constant of the constant of the six months period real utralisation test for equine vir more than one in four;]	stallion taken(7) not more ollection of the semen ken(7) from the donor octed with a positive result in				
			(c)	II.3.4.3. was	ous equine metritis, one of the s last carried out on three spec 60 days prior to the date of th n Part I	cimens (swabs) taken(7) not				
		(3)	$\circ$ either	[on two occ	casions at least 7 days apart;]					
		(3)	$\circ$ or	[on a single	e occasion and subjected to a I	PCR or real-time PCR.]]				
	L			0	,					

	II. Health info								13 LQUI-3L				
	n. neurin nito	ination											
		(6)	□ [II.3.5.3.	I he donor stallion does not meet the conditions set out in hounts 1 bial and (h)									
   g				samples ta	The tests described in points II.3.4.1, II.3.4.2. and II.3.4.3. were carried out on samples taken(7) from the donor stallion at least once a year at the beginning of the breeding season,								
Part II: Certification			and	the tests de taken(7) fr minimum before the	the tests described in points II.3.4.1 and II.3.4.3. were carried out on sample taken(7) from the donor stallion during the storage period of the semen of minimum period of 30 days from the date of the collection of the semen as before the semen is removed from the semen collection centre, not less the days and not more than 90 days after the collection of the semen described Part I,								
P		and	(3)	∘ either	carried out semen of a collection of semen colle	on samples minimum of the semen ection centr	s taken(7) d period of 30 n and befor re or used, 1	s described i luring the sta O days from The the semen not less than of the semen	orage period the date of t is removed 14 days and	l of the he from the l not more			
			(3)	<ul> <li>or [the non-shedder state of a donor stallion seropositive for equine viral arteritis was confirmed by virus isolation test, PCR or real-time PCR carried out with a negative result on samples of an aliquot of the entire semen of the donor stallion taken(7) twice a year at an interval of at least four months and the donor stallion reacted with a positive result at a serum dilution of at least one in four in a serum neutralisation test for equine viral arteritis.]]</li> </ul>									
		II.3.6.	underwent	t the testing	provided fo	r in point II	.3.5. on san	nples taken o	on the follow	ving dates.			
		Identificat		Start		-		nealth tests(		0			
		ion of	programm		, , , , , , , , , , , , , , , , , , , ,	2000 01 001			.,				
		semen	e	Donor residence	Semen collection	EIA II.3.4.1.	EVA II.3.4.2.		CEM II.3.4.	3.			
			C	Y'			Blood sample	Semen sample	1. sample	2. Sample			
			_										
										—			
										_			
										—			

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	II. Health info	rmation									
	(3) 0 either	[II.4.	No antibio	tics were added to the se	emen;]						
	(3) 0 or	[II.4.	The following antibiotic or combination of antibiotics was added to produce a concentration in the final diluted semen of not less than(8):;]								
		II.5.	The semen	described in Part I was:							
Part II: Certification			II.5.1. collected, processed, stored and transported under conditions which comply with the requirements of Chapters II(I)(1) and III(I) of Annex D to Directive 92/65/EEC;								
II: Cert			II.5.2.	eriod of 30 days from the							
Part ]		II.5.3. sent to the place of loading in a sealed container in accordance with Chapter III(I) of Annex D to Directive 92/65/EEC and bearing the num indicated in Box I.19.									
	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:										
	Box I.11:	-	-	-	emen collection centre of orig						
	Box I.12: The place of destination shall correspond to the semen collection or storage centre or to the holding of semen destination.										
	Box I.19:	The identif	ication of co	ontainer and seal numbe	er shall be indicated.						
	Box I.30:	The donor	identity sha	ll correspond to the offi	cial identification of the anim	al.					
	The date of collection shall be indicated in the following format: dd/mm/yyyy.										
	Part II:										
	Guidance f	or the comp	letion of the	e table in point II.3.6.:							
	Abbreviati	ons:									
		EIA-1	Equine inf	ectious anaemia (EIA) te	sting first occasion						
		EIA-2	EIA testing	second occasion							
		EVA-B1	Equine viral arteritis (EVA) testing on blood sample first occasion								
		EVA-B2	EVA testing	g on blood sample secon	d occasion						
		EVA-S1	EVA testing	g on semen sample first	occasion						
		EVA-S2	EVA testing	g on semen sample seco	nd occasion						
		CEM-11	Contagious	equine metritis (CEM) t	esting first occasion first sam	ple					
		CEM-12	CEM testin	g first occasion second s	ample taken 7 days after CEM	I-11					
		CEM-21	CEM testin	g second occasion first s	ample						
		CEM-22	CEM testin	g second occasion secon	d sample taken 7 days after C	EM-21					
	Instruction	.s:									
					below, the test programme (p l D shall be completed with th						
	I, as requir	ed in points	II.3.5.1., II.3	3.5.2. and II.3.5.3., shall b	prior to the first collection of e entered in the upper line of and CEM-11 and CEM-12 in t						
	II.3.5.3. sha	ll be entere	d in the low		testing as required in accord in table, this being the boxes						

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#### **EUROPEAN UNION**

ЕU	ROPEAN (	JNION						2021/4	03 EQUI-SE	M-B-INIK		
	II. Health info	rmation										
		Identificat ion of semen	Test programm e	Start date(7)		Date of sa	mpling for 1	health tests(	77)			
				Donor residence	Semen collection	EIA II.3.4.1.	EVA II.3.4.2.		CEM II.3.4.	3.		
ation	(1)						Blood sample	Semen sample	1. sample	2. sample		
I		А	В	С	D	EIA-1	EVA-B1	EVA-S1	CEM-11	CEM-12		
J						EIA-2	EVA-B2	EVA-S2	CEM-21	CEM-22		
מורחי	(1)	Only semen collection centres approved by the competent authority and listed in accordance with Article 11(4) of Directive 92/65/EEC.										
4	(2)	OJ L 268, 14.9.1992, p. 54.										
	(3)	Delete as appropriate.										
	(4)	OJ L 192, 2	3.7.2010, p. 1	Ι.								
	(5)	OJ L 165, 3	0.4.2004, p. 1	l.								
	(6)	Cross out t	he program	me(s) that d	o(es) not ap	ply to the c	consignmen	t.				
	(7)	Insert date	in table in j	point II.3.6 (	follow Guid	ance in Par	ct II of the N	otes).				
	(8)		les and conc	entrations.								
	Name (in car	icer/Official vet	erinarian			Qualificatio	n and title					
	Date of signa					Signature	ii anu tite					
	Stamp											
			Ĉ									