EUROPEAN UNION INTRA

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	I.1. Consignor				I.2. IMSOC ref	erence	I.2.a. Local reference	
	Name						I.3. Central Competent Authority	
						I.4. Local Competent Authority		
	Address						1.4. Local Competent Authority	
	Country ISO Code							
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ᆈ	I.5. Consignee				I.6. Operator conducting assembly operations independently of an			
Part I: Description of consignment	Name				establishment			
۳۱	Address				Name			
빔								
<u>છ</u>	Country		ISO Code			Address		
<u>छ</u>					Approval Nu	nber		
히					Country		ISO Code	
ರ					,			
ᇷ	I.7. Country of origi	in		ISO Code	I.9. Country of	destination	ISO Code	
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⊙∣								
밁	I.8. Region of origin	1		Code	I.10. Region of	destination	Code	
	I.11. Place of dispate				I.12. Place of o			
ပ္က	1.11. Place of dispati	.CII			1.12. Place of C	lestifiation		
اق	Name				Name			
7	Address				Address			
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디	Approval Number				Approval Nu	uner		
۵.	Country		ISO Code		Country		ISO Code	
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	I.13. Place of loadin	ıg			I.14. Date and	time of departure		
	Name							
	Address							
	Approval Number							
	Country		ISO Code					
	I.15. Means of Trans	sport			I.16. Transpor	ter		
			-1			102		
	Mode	International	Identification		Name			
		transport document			Address			
		document			Approval Nu	nber		
					Country		ISO Code	
					Country		150 Code	
					I 17 Assamna	nying documents		
					1.17. Accompa	nying documents		
					[en]			
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					accompanyi ng.documen t.document.		Date of issue	
				number				
						Place of		
					Country		issue	
ŀ	I 10. Transport cont	J:L:	one					
	I.18. Transport cond	aitions		_	Chilled □			
	Ambient \square		Frozen [_				
	I.19. Container No /	Seal No						
100 Control or								
	Dogistored equidae	I.20. Certified as						
	Registered equidae \square							
	I.21. For transit thro		ıntry					
			untry		ISO Code			
	I.21. For transit thro		untry					
	I.21. For transit thro Third country Exit point		untry		BCP code			
	I.21. For transit thro Third country Exit point Entry point	ough a third cou			BCP code BCP code	_		
	I.21. For transit thro Third country Exit point	ough a third cou			BCP code BCP code I.23. For expo			
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	I.21. For transit thro Third country Exit point Entry point I.22. For transit thro	ough a third cou	tate(s)		BCP code BCP code I.23. For exportant country	rt	ISO Code	
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	I.21. For transit thro Third country Exit point Entry point I.22. For transit thro Member State I.24. Estimated jour I.28. Total gross wei	ough a third cou ough Member S rney time	tate(s)		BCP code BCP code I.23. For expo Third country Exit point	og	ISO Code	
	I.21. For transit thro Third country Exit point Entry point I.22. For transit thro Member State I.24. Estimated jour I.28. Total gross wei	ough a third cou ough Member S rney time	tate(s) ISO Code	Subcategory	BCP code BCP code I.23. For expo Third country Exit point	rt	ISO Code	
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		UNION		(2021) 100,	MODEL EQUI-INTRA-IND		
	II. Health information						
	I, the und	I, the undersigned official veterinarian, hereby certify that:					
	II.1. The equine animal described in Part I meets the following requirements:						
		II.1.1.	The animal is accompanied by its single lifetime identification document as provided for in Article 65, 67 or 68 of Commission Delegated Regulation (EU) 2019/2035, or by a temporary document issued in accordance with Article 61(2) thereof.				
rt II: Certificatio	(1)(1)(1)) □ [The single lifetime identification document was issued in accordance wor 67(1) of Delegated Regulation (EU) 2019/2035, or the temporary document accordance with Article 61(2) of that Regulation, for a registered equine an Article 2(30) of that Delegated Regulation.]					
	(1)		\Box [The single lifetime identification document includes a valid validation mark in accordance with Article 65(1)(i)(i) of Delegated Regulation (EU) 2019/2035.]				
Ъ	(1)		☐ [The single lifetime identification Article 65(1)(i)(ii) of Delegated Reg		license in accordance with		
		II.1.2.	The animal has not shown signs or the clinical examination, which wa departure, or on the last working destablishment, on (in	is carried out within the 48 ho lay prior to its departure(2), f	our period prior to its		
	II.2. According to official information, the animal described in Part I meets the following health requirements:				ollowing health		
		II.2.1.	The animal does not come from an situated in a restricted zone establincluding African horse sickness at	ished for reasons of diseases l	isted for equine animals,		
		II.2.2.	The animal comes from an establis been reported during the 30 day pe				
	(1)		either \circ [surra has not been report departure.]	ted in the establishment durin	ng the 2 year period prior to		
	(1)		or \circ [surra has been reported in the departure and following the last our estrictions		, , ,		
	(1)		to a test for surra with Annex I to Commission negative results, on sar	aining animals in the establis one of the diagnostic method Delegated Regulation (EU) 20 nples taken at least 6 months ved from the establishment.]]	s provided for in Part 3 of 020/688, carried out, with		
	(1)			rs from the date of cleaning a e last animal of listed species byed or slaughtered.]]			
		II.2.3.	The animal comes from an establis the last 6 months prior to its depar		not been reported during		
	(1)		either o [dourine has not been rep to its departure.]	orted in the establishment du	ring the 2 year period prior		
	(1)		or \circ [dourine has been reported in departure and following the last or movement restrictions				
	(1)		castrated male equine the diagnostic method (EU) 2020/688, carried of months after the infect	aining equine animals in the canimals, have been subjected provided for in Part 8 of Annobut, with negative results, on ed animals have been killed accted entire male equine anim	to a test for dourine with ex I to Delegated Regulation samples taken at least 6 and destroyed or		
or \circ [for at least 30 days from the date of cleaning and disin equine animal on the establishment was either killed and disingly slaughtered.]]							

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_	_) MODEL EQUITATION
	II. Health inform	ation			
Part II: Certification	II	I.2.4.	The animal comes from an establis reported during the 90 day period	=	tious anaemia has not been
	(1)		either \circ [equine infectious anaemi 12 month period prior to its depart	•	ne establishment during the
	(1)		or \circ [equine infectious anaemia hamonth period prior to its departure remained under movement restrict	e and following the last outbr	
	(1)		subjected to a test for e provided for in Part 9 c out, with negative resu interval of 90 days follo	aining equine animals in the equine infectious anaemia with of Annex I to Delegated Regula lts, on samples taken on two bowing cleaning and disinfections ave been killed and destroyed	th the diagnostic method ation (EU) 2020/688, carried occasions with a minimum on of the establishment after
	(1)			ys from the date of cleaning a e last equine animal on the es r slaughtered.]]	
	II	I.2.5.	The animal comes from an establishas not been reported during the 6		
	(1)		either o [during the 2 year period] encephalomyelitis has not been rej establishment is situated.]		
	(1)		or \circ [during the 2 year period prio has been reported in the Member S situated, and during the 21 day per II.1 all equine animals in the estab	State or zone thereof in which riod prior to departure of the	n the establishment is animal referred to in point
	(1)		insect vectors in a quar rise in daily taken body diagnostic test for Vene method provided for ir	-	equine animal that showed a cted with negative result to a elitis with the diagnostic elegated Regulation (EU)
	(1)		with a commanufactur	accinated against Venezuelan plete primary course and rev rer's recommendations not le 12 months prior to the date o	accinated according to ss than 60 days and not
	(1)		encephalor 10(1)(b) of a out, with no	cted to a serological test for V nyelitis with the diagnostic m Annex I to Delegated Regulati egative results, on a sample to ate of its entry into the quara	ethod provided for in Part on (EU) 2020/688, carried aken not less than 14 days
	(1)		daily, either without a for Venezuelan equine for in Part 10(1)(a) of A negative results, and th	nture of the animal referred to rise or the animal has been so encephalomyelitis with the d annex I to Delegated Regulation he animal referred to in point quine encephalomyelitis with	ubjected to a diagnostic test liagnostic method provided on (EU) 2020/688, with II.1 has been subjected to
				Part 10(1)(b) of Annex I to De 2020/688, without an increas out on paired samples taken interval of 21 days, the secon the 10 day period prior to the	e in antibody titre, carried on two occasions with an nd of which was taken during

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	II. Health information							
			-	Part 10(2) of Annex I to Deleg 2020/688, with negative resultaken within the 48 hour per and the animal has been provectors after sampling until i	It, carried out on a sample iod prior to its departure, tected from attacks by insect			
Part II: Certification	II.2.6. The animal comes from an establishment in which infection with rabies viru terrestrial animals has not been reported during the 30 day period prior to it					_		
		II.2.7.	The animal comes from an establis reported during the 15 day period	shment in which anthrax in ungulates has not been I prior to its departure.				
Part II: C	II.3.	To the best of my knowledge, after due inquiry, and as declared by the operator, the animal comes from an establishment where there were no abnormal mortalities with an undetermined cause and the animal has not been in contact with kept animals of listed species which did not comply with the requirements referred to in points II.2.1. to II.2.6. during the 30 day period prior to its departure, and with the requirement referred to in point II.2.7. during the 15 day period prior to its departure.						
	(1) [II.4. According to official information and as declared by the operator, it is a semen donor animal subjected to the testing programme as referred to in point 1(b)(i) of Chapter I of Part 4 of Annex II to Commission Delegated Regulation (EU) 2020/686, and							
		II.4.1.	it comes from a semen collection centre and will be transported directly to another semen collection centre in accordance with Article 19 of Delegated Regulation (EU) 2020/686; and					
		II.4.2.	since the date of its admission, it was continuously resident at the semen collection centre and was subjected, with negative results, to all compulsory routine tests referred to in point 1(a) of Chapter I of Part 4 of Annex II to Delegated Regulation (EU) 2020/686 during the 12 month period prior to the date of its departure; and					
		II.4.3.	the prior consent of the centre vete been obtained by the operator; and		ion centre of destination has			
		II.4.4.	the means of transport used have been cleansed and disinfected before use.]					
	II.5.	Arrangements are made to						
	(1)		either \circ [transport the animal in accordance with Article 4 of Delegated Regulation (EU) 2020/688.]					
	(1)		or ○ [move the animal on foot.]					
	II.6.	This anima	l health certificate is valid for					
	(1)		either 0 [10 days from the date of i	issuing, and]				
	(1)		or \circ [30 days from the date of issuipoint II.1.1, and]	ing, and a valid validation ma	rk or license is attested in			
	in the case of transport by waterway/sea of the animal, the period of validity of the animal health certificate may be extended by the duration of the journey by waterway/sea.							
	Animal welfare attestation							
	At the time of inspection, the animal covered by this health certificate was fit to be transported in accordance with the provisions of Council Regulation (EC) No 1/2005 on the intended journey due to start on (insert date).							

EUROPEAN UNION (2021/403) MODEL EQUI-INTRA-IND II. Health information Notes: In accordance with the Agreement on the withdrawal of the United Kingdom of Great Britain and Northern Ireland from the European Union and the European Atomic Energy Community, and in particular Article 5(4) of the Protocol on Ireland / Northern Ireland in conjunction with Annex 2 to that Protocol, references to European Union Certification in this certificate include the United Kingdom in respect of Northern Ireland. 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: ä Box "Place of dispatch": Indicate a registered establishment of dispatch of the equine animal or, provided refer I.11: reference the animal is transported, an establishment approved for assembly operations in accordance with Articles 97 and 99 of Regulation (EU) 2016/429. "Place of destination": Indicate a registered establishment of destination or, provided the animal is Box reference transported, an establishment approved for assembly operations in accordance with Articles 97 and 99 of Regulation (EU) 2016/429. I.12: "Identification number": Indicate the unique code of the equine animal referred to in Article 65(1)(b) of Box reference Delegated Regulation (EU) 2019/2035, or the code displayed by the means of identification defined in point (a), (c) or (e) of Annex III to Delegated Regulation (EU) 2019/2035, if the animal is unweaned and I.30: accompanies its dam or foster mare. Part II: (1) Delete if not applicable. (2) Option only available in case of equine animals moved in accordance with Article 92(2) of Delegated Regulation (EU) 2020/688. Certifying Officer/Official veterinarian Qualification and title Name (in capital letters) Date of signature Signature Stamp