**INTRA** 

	I.1. Consignor			I.2. IMSOC reference I.2.a. Local reference			
	Name				I.3. Central Competent Authority		
	Address Country		ISO Code				
ı			100 couc		I.4. Local Competent Authority		
ent	I.5. Consignee			I.6. Operator conducting assembly operations independently of an establishment			
nm	Name Address			Name			
igi	Country		ISO Code	Address			
suc				Approval Number			
of consignment				Country	ISO Code		
	I.7. Country of orig	gin	ISO Code	I.9. Country of destination	ISO Code		
Part I: Description	I.8. Region of origi	in	Code	I.10. Region of destination	Code		
Desc	I.11. Place of dispa	atch		I.12. Place of destination			
<b>I</b> :	Name Address			Name Address			
art	Approval						
Ь	Nûmber		100 0-4-	Approval Number	100 0-1-		
	Country		ISO Code	Country	ISO Code		
	I.13. Place of loadi	ng		I.14. Date and time of departure			
	Name						
	Address Approval						
	Number		100 0-4-				
	Country		ISO Code				
	I.15. Means of Trai		- 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	I.16. Transporter			
	Mode	International transport	Identification	Name			
		document		Address			
				Approval Number	100 0 1		
				Country	ISO Code		
				I.17. Accompanying documents			
				Document Type Accompanying document			
				reference			
				Date of Issue			
				Country Place of issue			
	I.18. Transport cor	nditions					
	Chilled ☐ Ambient ☐			Frozen $\square$			
	I.19. Container No	/ Seal No					
	I.20. Certified as	- Tocal Ivo					
	Confined establish	nment 🗆	Exhibition $\square$	Event or activity near borders $\Box$	Further keeping $\square$		
		rough a third coun	try				
	Third country Exit point			ISO Code BCP code			
	Entry point			BCP code			
		rough Member Sta	te(s)	I.23. For export			
	Member State		ISO Code	Third country	ISO Code		
				Exit point	BCP code		
	I.24. Estimated jou	irney time		I.25. Journey Log			
	I.27. Total quantity			I.28. Total gross weight			
	I.30. Description of	f consignment					
	1. 01 LIVE ANIMALS						
	0101 Live horses, asses, mules and hinnies Non-Registered equine animals						
	#1. Commodity		Subcategory	Sex	Identification system		

en 1/6

	Species	Identification Number	Age	Quantity
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Part I: Description of consignment				
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en 2 / 6

01	ION					=0=1,1	044 (2021/403	,		
	II. Health information									
	I, the undersigned official veterinarian, hereby certify tha					<u> </u>				
	II.1.	•			e following re	equirements:				
uc		II.1.1.	The anima	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.						
Part II: Certification	(1)		or Article issued in a	☐ [The single lifetime identification document was issued in accordance with Article 65(2) or Article 67(1) of Delegated Regulation (EU) 2019/2035, or the temporary document was issued in accordance with Article 61(2) of that Regulation, for a registered equine animal as defined in Article 2(30) of that Delegated Regulation.]						
1 II: 0	(1)			☐ [The single lifetime identification document includes a valid validation mark in accordance with Article 65(1), point (i)(i), of Delegated Regulation (EU) 2019/2035.]						
Pa	(1)			☐ [The single lifetime identification document includes a valid license in accordance with Article 65(1), point (i)(ii), of Delegated Regulation (EU) 2019/2035.]						
		II.1.2.	the clinica	The animal has not shown signs or symptoms of diseases listed for equine animals during the clinical examination, which was carried out within the last 48 hours prior to the time of its departure, or on the last working day prior to the date of its departure, from the registered establishment, on (insert date dd/mm/yyyy).						
	II.2.	According requireme		nformation,	, the animal d	escribed in Pa	art I meets the f	ollowing health		
	(1) o either	[II.2.1.	affecting e diseases s	The animal comes from an establishment or a zone not subject to movement restrictions affecting equine animals and established for reasons of diseases listed for those species or diseases subject to emergency measures relevant for those species, and the animal has not been in contact with kept animals of a listed species of a lower health status for an adequate period l						
	(1) or ∘	[II.2.1.	equine an	imals and e	m an establis stablished for n granted, and	•	•	vement restrictions affecti ations from movement		
	(1)	□ [it comp	lies with th	lies with the requirements set out in (3);]]						
	(1)	□ [and in	particular, i	articular, it is (4).]]						
		II.2.2.		The animal comes from an establishment in which surra (Trypanosoma evansi) has not been reported during the last 30 days prior to the date of its departure, and						
		(1) either	$\circ$ [surra has not been reported in the establishment during the last 2 years prior to the date of its departure.]							
		(1) or	its departı	are and follo		e of the last o		2 years prior to the date of ected establishment has		
	(1)			either o	have been s methods pr Regulation samples tak	subjected to a rovided for in (EU) 2020/688 ken at least 6 1	test for surra w Part 3 of Annex , carried out, w nonths followir	inimals in the establishme with one of the diagnostic It to Commission Delegate ith negative results, on ag the date on which the la the establishment.]]		
	(1)			or o	listed speci	es in the estab	lishment was e	n which the last animal of ither killed and destroyed, leaned and disinfected.] ]		
		II.2.3.				hment in whi of its departu		not been reported during		
	(1)		either $\circ$		nas not been i e of its depart	_	e establishment	t during the last 2 years pr		
the date of i				f its departur	e and followi		ring the last 2 years prior to last outbreak the affecte ictions:			

en 3/6

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	II. Health information					
Part II: Cartification	(1)		either ∘	establishm subjected to for in Part of carried out following to and destroy	ate on which the remaining e ent, except castrated male eq o a test for dourine with the o 8 of Annex I to Delegated Reg , with negative results, on sar he date on which the infected yed, or slaughtered, or the inf ve been castrated.]]	uine animals, have been liagnostic method provided ulation (EU) 2020/688, nples taken at least 6 months animals have been killed
II. Carti	(1)		or o	the establis	t 30 days after the date on wh chment was either killed and emises were cleaned and disi	destroyed, or slaughtered,
Dart	II.2.4.				shment in which equine infection rior to the date of its departur	
	(1)	[equine infectious anaemia has not been reported in the establishment du the last 12 months prior to the date of its departure.]		- 1		
	(1)	or o	last 12 mo	onths prior to		e establishment during the following the date of the last under movement restrictions:
	(1)		either ∘	establishm anaemia w I to Delegat results, on of 90 days t been killed	ate on which the remaining e ent have been subjected to a s ith the diagnostic method pro ted Regulation (EU) 2020/688, samples taken on two occasion following the date on which the and destroyed, or slaughtered d and disinfected.]]	test for equine infectious ovided for in Part 9 of Annex carried out, with negative ons with a minimum interval he infected animals have
	(1)		or o	animal in t	t 30 days following the date o he establishment was either l d, and the premises were clea	killed and destroyed, or
			shment in which Venezuelan ast 6 months prior to the date			
	(1)	either o	encephalo	omyelitis has	s prior to the date of its depar not been reported in the Men nt is situated.]	_
	(1)	or o	encephalo which the of departi	omyelitis has e establishme ure of the ani	s prior to the date of its depar been reported in the Member nt is situated, and during the mal referred to in point II.1 a mained clinically healthy, and	r State or zone thereof in last 21 days prior to the date ll equine animals in the
	(1)		either o	protected f showed a r with negati encephalor 10, point 10	I referred to in point II.1. was from attacks by insect vectors ise in daily taken body tempe ive result to a diagnostic test f myelitis with the diagnostic m (a), of Annex I to Delegated Re referred to in point II.1 has b	, and any equine animal that erature has been subjected for Venezuelan equine tethod provided for in Part egulation (EU) 2020/688, and
	(1)			either o	[vaccinated against Venezue encephalomyelitis with a cor revaccinated according to m recommendations not less th than 12 months prior to the	mplete primary course and anufacturer's aan 60 days and not more
	(1)			or o	[subjected to a serological te encephalomyelitis with the conforming of Part 10, point 1(b), of Part 10, 2020/688, can	liagnostic method provided Annex I to Delegated

en 4/6

	NION					2024/1044 (2021/403	3) Model EQUI-INTRA-IND
	II. Health info	ormation					
						results, on a sample taken n date of its entry into quarar	not less than 14 days after the ntine.]]]
Part II: Certification	(1) or o		[the body temperature of the animal referred to in point II.1 has been taken daily, either without a rise or the animal has been subjected to a diagnostic test for Venezuelan equine encephalomyelitis with the diagnostic method provided for in Part 10, point 1(a), of Annex I to Delegated Regulation (EU) 2020/688, with negative results, and the animal referred to in point II.1 has been subjected to tests for Venezuelan equine encephalomyelitis with the diagnostic methods provided for in:				
Part II:					-	Part 10, point 1(b), of Annex (EU) 2020/688, without an in carried out on paired sampl with an interval of 21 days, taken during the last 10 day departure, and	ncrease in antibody titre, les taken on two occasions the second of which was
					-	2020/688, with negative rest taken within the last 48 hou departure, and the animal h	rs prior to the time of its
		II.2.6.		l animals h		ishment in which infection w eported during the last 30 day	-
		II.2.7.		al comes froduring the l	ungulates has not been re.		
	II.2.8. The animal comes from an esta inquiry, and as declared by the undetermined cause and the ar species which did not comply we during the last 30 days prior to to in point II.2.7 during the last				ared by the op and the anim ot comply with ys prior to the	perator, there were no abnorn nal has not been in contact wi h the requirements referred t e date of its departure, and w	nal mortalities with an ith kept animals of listed o in points II.2.1 to II.2.6 ith the requirement referred
	II.3.	Arranger	nents are ma	ade to:			
	(1) either o	[transpor	rt the animal	l in accorda	ince with Arti	icle 4 of Delegated Regulation	(EU) 2020/688.]
	(1)		or $\circ$	[move th	e animal on f	coot.]	
	II.4.	This anin	nal health ce	rtificate is	valid for:		
	(1)		either $\circ$	[10 days	from the date	e of issuing, and]	
	(1)		or o	-	from the date d in point II.1	e of issuing, and a valid valida 1, and]	ation mark or a valid license

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 animal 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).

## 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 the Union in this animal health certificate include the United Kingdom in respect of Northern Ireland.

This animal health certificate shall be completed in accordance with the notes for the completion of certificates

UI	UNION 2024/1044 (2021/403) Model EQUI-INTRA-IND								
	II. Health info	rmation							
	provided for	or in Chapter 2 of Annex I to Commission Imple	menting Regulation (EU) 2020	)/2235.					
Certification	Box "Place of dispatch": Indicate a registered establishment of dispatch of the equine animal or, provide reference the animal is transported, an establishment approved for assembly operations in accordance with								
	Box "Place of destination": Indicate a registered establishment of destination or, provided the animal is reference transported, an establishment approved for assembly operations in accordance with Articles 97 at of Regulation (EU) 2016/429, or a veterinary clinic.								
Part II: Ce	Box "For transit through Member State(s)": Indicate each Member State of transit. In the case of an animal reference health certificate that is valid for 30 days, indicate all Member States that the equine animal has transited while returning to the establishment of its departure.								
Pa	Box reference I.30:	"Identification number": Indicate the unique code of the equine animal referred to in Article 65(1), ence point (b), of Delegated Regulation (EU) 2019/2035.							
	Part II:								
	(1)	Delete if not applicable.							
	(2)	Insert the name of the disease(s).							
	(3)	Insert the specific reference to the article(s), tie Commission providing for those requirements		nt legal act(s) adopted by the					
	(4)	Insert the specific attestation(s) provided for in Commission, as referred to in Article 126(1), po							
		icer/Official veterinarian	O						
	Name (in capi Date of declar		Qualification and title Signature						

en 6/6