MODEL ANIMAL HEALTH CERTIFICATE FOR THE MOVEMENT BETWEEN MEMBER STATES OF AN INDIVIDUAL EQUINE ANIMAL NOT INTENDED FOR SLAUGHTER (MODEL 'EQUI-INTRA-IND')

EURC	PEAN UN	IION				INTRA			
	I.1	Consignor		I.2	IMSOC reference				
		Name		I.2a	Local reference				
		Address		I.3	Central Competent Authority	QR CODE			
nt		Country	ISO country code	I.4	Local Competent Authority				
nme	I.5	Consignee	Consignee Name		Operator conducting assembly operations independently establishment				
nsig		Name			Name	Registration No			
oj co		Address			Address				
Part I: Description of consignment		Country	ISO country code		Country	ISO country code			
scrip	I.7	Country of origin	ISO country code	1.9	Country of destination	ISO country code			
De	I.8	Region of origin	Code	I.10	Region of destination	Code			
Ξ:	I.11	Place of dispatch		I.12	Place of destination				
Part		Name	Registration/Approval No		Name	Registration/Approval No			
		Address			Address				
		Country	ISO country code		Country	ISO country code			
	I.13	Place of loading		I.14	Date and time of departure				
	I.15	Means of transport		I.16	Transporter				
		□ Vessel □ Aircraft			Name	Registration/Authorisation No			
					Address				
		□ Railway	□ Road vehicle		Country	ISO country code			
				I.17	Accompanying documents				
		Identification	□ Other		Type	Code			
		Document			Country Commercial document reference	ISO country code			
	I.18	Transport condition	s Ambient	'	□ Chilled □	Frozen			
	I.19	Container number/S	Seal number						
		Container No Seal No							

I.20	Certified as or	for							
□ Furth	er keeping	□ Slaughter	□ Slaughter		□ Confined establishment		□ Germinal products		
□ Regis	stered equine animal	□ Travelling circus/anima	□ Travelling circus/animal act		□ Exhibition		□ Event or activity near borders		
□ Relea	ase into the wild	□ Dispatch centre	□ Dispatch centre		□ Relaying area/purification		□ Ornamental aquaculture		
					centre		establishment		
□ Furth	er processing	□ Organic fertilizers and	□ Organic fertilizers and soil		□ Technical use		□ Quarantine or similar		
		improvers	improvers				establishment		
	ucts for human	□ Pollination		□ Live aquat	ic animals for	□ Other			
consum	nption			human consu	ımption				
I.21	□ For transit	through a third country							
	Third country				ıntry code				
	Exit point		BCP code BCP code						
	Entry point						~		
I.22	□ For transit throu	gh Member State(s)			export				
	Member State	ISO country	ISO country code ISO country code		Third country Exit point		ISO country code BCP code		
	Member State	ISO country							
	Member State	ISO country	code						
I.24	Estimated journey	time		I.25 Jo	urney log	□ yes		□ no	
I.26	Total number of packages I.27 Total quantity								
I.28	Total net weight/gross weight (kg) I.29 Total space foreseen for the consignment						t		
I.30	Description of cons	0							
CN cod	le Species	Subspecies/Category Sex	Ident system	ification m	Identification 1	number	Age	Quantity	
			system					Type	
Region of origin		Cold store	Identification mark		Type of packa	ging		Net weight	
Slaughterhouse		Treatment type	Nature of commodity		Number of pac	Number of packages		Batch No	
		Date of collection/production	Manı plant	ufacturing	Approval or re number of plant/establish		Test		

	II. Health inform	ation		II.a	Certificate reference	II.b	IMSOC reference		
	I, the t	ındersigned	official veterinarian, hereby ce	rtify t	hat:				
	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.						
			(1) [The single lifetime identification document was issued in accordance with Article 65(2) or 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) [The single lifetime identification document includes a valid validation mark in accordance with Article 65(1)(i)(i) of Delegated Regulation (EU) 2019/2035.]						
			(1) [The single lifetime identification document includes a valid license in accordance with Article 65(1)(i)(ii) of Delegated Regulation (EU) 2019/2035.]						
Part II: Certification		II.1.2.	The animal has not shown signs or symptoms of diseases listed for equine animals during the clinical examination, which was carried out within the 48 hour period prior to its departure, or on the last working day prior to its departure ⁽²⁾ , from the registered establishment, on(insert date dd/mm/yyyy).						
art II: (II.2.	Accordin requireme	ng to official information, the animal described in Part I meets the following health ents:						
Pa		II.2.1.	The animal does not come fi situated in a restricted zon animals, including African (glanders).	e esta	ablished for reasons of	of disea	ses listed for equine		
		II.2.2.	The animal comes from an not been reported during the						
		⁽¹⁾ either	[surra has not been reported departure.]	d in t	he establishment durin	ng the 2	year period prior to		
		⁽¹⁾ or	[surra has been reported in departure and following th movement restrictions						
			test for surra with Annex I to Comm with negative res	one nission sults,	imals in the establishm of the diagnostic meth in Delegated Regulation on samples taken at 1 in removed from the es	ods pro n (EU) 2 east 6	vided for in Part 3 of 2020/688, carried out, months after the last		

	⁽¹⁾ or	[for at least 30 days from the date of cleaning and disinfection of the establishment, after the last animal of listed species on the establishment was either killed and destroyed or slaughtered.]]		
II.2.3.		al comes from an establishment in which dourine has not been reported last 6 months prior to its departure, and		
⁽¹⁾ either	[dourine hadeparture.]	as not been reported in the establishment during the 2 year period prior to its		
$^{(l)}or$	departure	as been reported in the establishment during the 2 year period prior to its and following the last outbreak, the establishment has remained under restrictions		
	⁽¹⁾ either	[until the remaining equine animals in the establishment, except castrated male equine animals, have been subjected to a test for dourine with the diagnostic method provided for in Part 8 of Annex I to Delegated Regulation (EU) 2020/688, carried out, with negative results, on samples taken at least 6 months after the infected animals have been killed and destroyed or slaughtered, or the infected entire male equine animals have been castrated.]]		
	⁽¹⁾ or	[for at least 30 days from the date of cleaning and disinfection after the last equine animal on the establishment was either killed and destroyed or slaughtered.]]		
II.2.4.		al comes from an establishment in which equine infectious anaemia has not red during the 90 day period prior to its departure, and		
⁽¹⁾ either		fectious anaemia has not been reported on the establishment during the 12 iod prior to its departure.]		
⁽¹⁾ or	[equine infectious anaemia has been reported on the establishment during month period prior to its departure and following the last outbreak the establishment during the last outbreak the establishment restrictions			
	⁽¹⁾ either	[until the remaining equine animals in the establishment have been subjected to a test for equine infectious anaemia with the diagnostic method provided for in Part 9 of Annex I to Delegated Regulation (EU) 2020/688, carried out, with negative results, on samples taken on two occasions with a minimum interval of 90 days following cleaning and disinfection of the establishment after the infected animals have been killed and destroyed or slaughtered.]]		
	$^{(l)}or$	[for at least 30 days from the date of cleaning and disinfection of the establishment, after the last equine animal on the establishment was either killed and destroyed or slaughtered.]]		
II.2.5.		nal comes from an establishment in which Venezuelan equine myelitis has not been reported during the 6 month period prior to its and		
⁽¹⁾ either		e 2 year period prior to its departure, Venezuelan equine encephalomyelitis en reported in the Member State or zone thereof in which the establishment		
	(1) either (1) or II.2.4. (1) either (1) or	II.2.3. The anima during the (*)either [dourine hadeparture.] (*)or [dourine hadeparture movement (*)either II.2.4. The anima been report (*)either [equine in month perhas remain (*)either (*)or II.2.5. The anima has remain (*)either (*)either [during the has not be		

 $^{(1)}or$

[during the 2 year period prior to its departure, Venezuelan equine encephalomyelitis has been reported in the Member State or zone thereof in which the establishment is situated, and during the 21 day period prior to departure of the animal referred to in point II.1 all equine animals in the establishment have remained clinically healthy, and

(1) either

[the animal referred to in point II.1 was kept protected from attacks by insect vectors in a quarantine station, in which any equine animal that showed a rise in daily taken body temperature has been subjected with negative result to a diagnostic test for Venezuelan equine encephalomyelitis with the diagnostic method provided for in Part 10(1)(a) of Annex I to Delegated Regulation (EU) 2020/688, and the animal referred to in point II.1 has been

(1) either [vaccinated against Venezuelan equine encephalomyelitis with a complete primary course and revaccinated according to manufacturer's recommendations not less than 60 days and not more than 12 months prior to the date of its departure.]]]

(1) or [subjected to a serological test for Venezuelan equine encephalomyelitis with the diagnostic method provided for in Part 10(1)(b) of Annex I to Delegated Regulation (EU) 2020/688, carried out, with negative results, on a sample taken not less than 14 days after the date of its entry into the quarantine station.]

⁽¹⁾or

[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(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 10(1)(b) of Annex I to Delegated Regulation (EU) 2020/688, without an increase in antibody titre, carried out on paired samples taken on two occasions with an interval of 21 days, the second of which was taken during the 10 day period prior to the date of its departure, and
- Part 10(2) of Annex I to Delegated Regulation (EU) 2020/688, with negative result, carried out on a sample taken within the 48 hour period prior to its departure, and the animal has been protected from attacks by insect vectors after sampling until its departure.]]
- II.2.6. The animal comes from an establishment in which infection with rabies virus in kept terrestrial animals has not been reported during the 30 day period prior to its departure.
- II.2.7. The animal comes from an establishment in which anthrax in ungulates has not been reported during the 15 day period prior to its departure.

- 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.
- (I)[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 veterinarian of the semen collection centre of destination has been obtained by the operator; and
 - II.4.4. the means of transport used have been cleansed and disinfected before use.]
 - II.5. Arrangements are made to
 - (1) either [transport the animal in accordance with Article 4 of Delegated Regulation (EU) 2020/688.]
 - (1) or [move the animal on foot.]
 - II.6. This animal health certificate is valid for
 - (1) either [10 days from the date of issuing, and]
 - (1) or [30 days from the date of issuing, and a valid validation mark or license is attested in point II.1.1, and]

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

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 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:

Box reference I.11: "Place of dispatch": Indicate a registered establishment of dispatch of the equine

animal or, provided the animal is transported, an establishment approved for assembly operations in accordance with Articles 97 and 99 of Regulation (EU) 2016/429.

Box reference I.12: "Place of destination": Indicate a registered establishment of destination or, provided

the animal is transported, an establishment approved for assembly operations in

accordance with Articles 97 and 99 of Regulation (EU) 2016/429.

Box reference I.30: "Identification number": Indicate the unique code of the equine animal referred to in

Article 65(1)(b) of 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 accompanies its dam or

foster mare.

Part II:

Delete if not applicable.

 \triangleright ⁽¹⁾(2) Option only available in the case of either:

- (a) an equine animal accompanied by its single lifetime identification document as provided for in Article 114(1), point (c), of Regulation (EU) 2016/429, which includes a valid validation mark referred to in Article 92(2), point (a), of Delegated Regulation (EU) 2020/688; or
- (b) a registered equine animal accompanied by its single lifetime identification document as provided for in Article 114(1), point (c), of Regulation (EU) 2016/429 which includes a valid license referred to in Article 92(2), point (b), of Delegated Regulation (EU) 2020/688, or by its single lifetime identification document accompanied by the FEI Recognition Card together with the validation sticker. ◀

Official veterinarian

Name (in capital letters) Qualification and title

Local Control Unit name Local Control Unit code

Date

Stamp Signature