Imports of consignments of stocks of semen of equidae collected, processed and stored in accordance with Directive 92/65 after 31 August 2010 and before 1 October 2014 and dispatched after 31 August 2010 from an approved semen collection centre of origin of the semen

GBHC811 v1.1 Apr-24

I.1 Consignor I.2 Certificate									
Name:									
Address:			10 11			1.4.1			
			1.2.a N	ot in use		I.4 Loca	I competent auti	nority	
Tel:									
I.5 Consignee					I.6 Person respo	nsible fo	or the load in Gre	at	
Name:							() `		
Address:					Name:				
					Address:				
Tel:					Tel:				
I.7 Country of	ISO	I.8 Regi		Code	I.9 Country of	ISO	I.10 Region of	Code	
origin	code	origi	n		destination	code	destination		
I.11 Place of or	igin				I.12 Place of des	stination			
☐ Semen centre	Э				☐ Semen centre				
Name:					Holding				
Approval number	er:			2	Name:				
Address:					Approval number	:			
					Address:				
I.13 Place of loa	ading				I.14 Date of depa	arture			
I.15 Means of tr	ansport				I.16 Entry BCP				
☐ Aeroplane									
Ship									
Railway wago	on								
☐ Road vehicle									
Other					I.17 Not in use				
Identification:									
Documentation i	references	s:							

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Equine	semen	Model	2
GBHC8	11		

II.a. Certificate reference no.	II.b.

I.18 Description of commodity						
I.19 Commodity code (HS code)	I.21 N	lot in use		I.23 Seal / Containe	r No.	
05 11 99 85						
I.20 Quantity	1.22 N	Number of packages I.24 Not in use				
I.25 Commodity certified for						
Artificial reproduction	Artificial reproduction					
I.26 For transit through Great I country	Britain	to third	I.27 For impo	ort or admission into	Great Britain	
Third country	ISO (Code				
I.28 Identification of the commod	ities					
Species (Scientific name)		Don	or identity	Date of collection	Quantity	
			/\U'			
		2-				

Part II. Certification

Animal Health

AH/E351G Establishment requirement (Collection centre)

The semen collection centre, in which the semen described above was collected, processed and stored for export to Great Britain is approved and supervised by the competent authority as per GB requirements;

AH/E605B Establishment requirements (freedom from disease)

during the period commencing 30 days prior to the date of first collection of the semen described above until the 30 days storage period for frozen semen elapsed, the semen collection centre:

- (a) It is situated in the exporting country or, in the case of regionalisation according to GB requirements, in that part of the territory of the exporting country which was:
 - (i) not considered to be infected with African horse sickness as per GB requirements,
 - (ii) free from Venezuelan equine encephalomyelitis for a period of at least 2 years,
 - (iii) free from glanders and dourine for a period of at least 6 months
- (b) fulfilled the conditions for a holding as per GB requirements and in particular:

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II.a. Certificate reference no.	II.b.

(*)EITHER

(i)

following a case of a disease mentioned below not all the animals of species susceptible to that disease located in the holding were slaughtered or killed and the holding has been free from any type of equine encephalomyelitis, equine infectious anaemia (EIA), vesicular stomatitis (VS), rabies, and anthrax as per GB requirements]

(*)**OR**

- (ii) [following a case of a disease mentioned below all the animals of species susceptible to that disease located in the holding have been slaughtered or killed and the premises disinfected, and the holding was free for a period of at least 30 days from any type of equine encephalomyelitis, equine infectious anaemia, vesicular stomatitis and rabies or 15 days in the case of anthrax, beginning on the day on which following the destruction of the animals the disinfection of the premises was satisfactorily completed;]
- (c) contained only equidae which were free of clinical signs of equine viral arteritis and contagious equine metritis,

AH/A725B Animal requirements

The semen described above was collected from donor stallions which:

- (a) have not shown any clinical sign of an infectious or contagious disease at the time of admission onto the semen collection centre and on the day the semen was collected:
- (b) have been kept for 30 days prior to the date of semen collection on holdings where no equine animal has shown any clinical sign of equine viral arteritis or contagious equine metritis during that period;
- (c) have not been used for natural mating during at least 30 days prior to the date of first semen collection and between the dates of the first sample referred to in points (e)(ii), (e)(ii) and/or (e)(iii) of this attestation and until the end of the collection period;
- (d) have undergone the following tests, which meet at least the requirements of the relevant Chapter of the Manual of Diagnostic Tests and Vaccines for Terrestrial Animals of the WOAH, carried out on samples taken in accordance with one of the programmes specified in point (e) in a laboratory recognised by the competent authority:

(*) **EITHER**

(EIA) considering GB requirements, with negative result;

(*)**OR**

(ii) an ELISA for equine infectious anaemia (EIA), considering GB requirements, with negative result;

and

(*)EITHER

[a serum neutralisation test for equine viral arteritis (EVA) with negative result at a serum dilution of one in four;]

(*)**OR**

[virus isolation test for equine viral arteritis (EVA) carried out with negative result on an aliquot of the entire semen of the donor stallion;]

and

- (iii) an agent identification test for contagious equine metritis (CEM) carried out as per GB requirements;
- (e) have been subjected with the results specified in **point** (d) in each case to at least one of the test programmes:

(*)[(i)],

(*)[(ii option 1)] or (*)[(ii option 2)]

(*)[(iii)]

(f) have undergone the testing provided for in (*)AH/A781B point (b) and point (d) on samples taken on the following dates:

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II.a. Certificate reference no.	II.b.

Identific ation of semen	Test progra mme	Start date: Donor residen ce	Start date: Semen collecti on	Date of samplin g for health tests: (*)VS point (b) of AH/A78 1B	Date of samplin g for health tests: EIA point (d)(i) of AH/A72 5B	Date of samplin g for health tests: EVA point (d)(ii) of AH/A72 5B Blood sample	Date of samplin g for health tests: EVA point (d)(ii) of AH/A72 5B Semen sample	Date of samplin g for health tests: CEM point (d)(iii) of AH/A72 5B 1. sample	Date of samplin g for health tests: CEM point (d)(iii) of AH/A72 5B 2. sample

AH/A781B Animal requirements (freedom from disease)

Prior to entering the semen collection centre the donor stallions and any other equidae located in the centre:

- (a) were continuously resident for a period of 3 months (or since entry if they were directly imported from Great Britain during the 3 months period) in the exporting country or, in the case of regionalisation as per GB requirements, in that part of the territory of the exporting country which was during that period:
 - (i) not considered to be infected with African horse sickness as per GB requirements,
 - (ii) free from Venezuelan equine encephalomyelitis for a period of at least 2 years,
 - (iii) free from glanders and dourine for a period of at least 6 months;
- (b) Vesicular Stomatitis:
 - originated from the country of export which was on the day of admission into the centre free from vesicular stomatitis (VS) for a period of at least 6 months,]
 - (*) **OR** [(ii) were subjected to a virus neutralisation test for vesicular stomatitis (VS) carried out with negative result as per GB requirements]
- (c) originated from holdings which on the day of admission onto the centre fulfilled the requirements of point (b) of AH/E605B Establishment requirements;

AH/P463 Product requirements

(*)EITHER	(a)	[No antibiotics were added to the semen.]
(*) OR	(b)	[The following antibiotic or combination of antibiotics was added to produce a concentration in the final diluted semen of not less than:
]

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Equ	ine	semen	Model	2
GB H	IC8	11		

II.a. Certificate reference no.	II.b.

AH/P551D Product requirements (storage and transport)

the semen described above was:

- (a) collected, processed, stored and transported under conditions which comply with GB requirements;
- (b) sent to the place of loading in a sealed container in accordance with GB requirements.
- (*) Keep as appropriate.

Official Veterinarian						
By signing this certificate, I certify that the requirements laid out above and in the accompanying notes for completion have been met.						
Name (in capital letters):	Qualification and title:					
Date:	Signature:					
Stamp:						

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Part III. Notes for completion

These notes for completion must be read and understood by the certifying officer before signing the certificate. Notes are set out in sections that correspond to the sections in the certificate. By signing this certificate, certifiers are verifying that the consignment meets the requirements set out in the certificate and any relevant corresponding notes for completion.

These notes do not need to be printed as part of a paper certificate that accompanies the consignment or in any electronic copy of the certificate.

References to European Union legislation within this certificate are references to direct EU legislation which has been retained in Great Britain (retained EU law as defined in the European Union (Withdrawal) Act 2018) and can be viewed on the UK legislation website (legislation.gov.uk).

References to Great Britain in this certificate include Channel Islands and Isle of Man.

References to GB requirements refer to the requirement(s) of Great Britain as set out in the accompanying notes for completion.

Part I

Box reference I.11: The *place of origin* shall correspond to the semen collection centre of the semen

origin.

Box reference I.22: The *number of packages* shall correspond to the number of containers.

Box reference I.23: The identification of container and seal number shall be indicated.

Box reference I.28: The *donor identity* shall correspond to the official identification of the animal.

The date of collection shall be indicated in the following format dd/mm/yyyy.

Part II

Animal Health

Imports of equine semen are authorised from a third country listed in column 2 as set out in a document relating to 'equidae' published on gov.uk, in accordance with Regulation 2018/659 provided that the semen was collected in the part of the territory of the third country detailed in column 4 of that document from a donor stallion of the category of Equidae indicated in columns 11, 12 or 13 of that document (†)

AH/E351G Establishment requirement (Collection centre)

Only approved semen collection centres listed in accordance with Article 17(3)(b) of Directive 92/65.

The semen collection centre is approved and supervised by the competent authority in accordance with the conditions of the Chapters I(I)(1) and I(II)(1) of Annex D to Directive 92/65

AH/E605B Establishment requirements (freedom from disease)

(a) regionalisation according to Article 13 of Directive 2009/156

Not considered to be infected with African horse sickness in accordance with Article 5(2)(a) and (b) of Directive 2009/156 (†).

(b) Fulfilled the conditions for a holding laid down in Article 4(5) of Directive 2009/156 (†).

For the **EITHER** option:

- (i) from any type of equine encephalomyelitis for a period of at least 6 months, beginning on the day on which the 1quidae suffering from the disease are slaughtered.
- (ii) from equine infectious anaemia (EIA) for at least the period required to obtain a negative result in an agar gel immunodiffusion test (Coggins test) carried out on samples taken after the infected animals were slaughtered on two occasions 3 months apart from each of the remaining animals,
- (iii) from vesicular stomatitis (VS) for a period of at least 6 months from the last recorded case,
- (iv) from rabies for a period of at least one month from the last recorded case

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AH/A725B Animal requirements

- (d)(i)The agar gel immunodiffusion test (Coggins test) or the ELISA for equine infectious anaemia are not required for donor equidae which have continuously resided in Iceland since birth, provided that Iceland has remained officially free of equine infectious anaemia and no equidae and their semen, ova and embryos have been introduced into Iceland from outside prior to and during the period the semen was collected.
- (d) (iii) an agent identification test for contagious equine metritis (CEM) carried out on two occasions on samples collected with an interval of 7 days by isolation of *Taylorella equigenitalis* after a cultivation of 7 to 14 days from pre-ejaculatory fluid or a semen sample and from genital swabs taken at least from the penile sheath, urethra and urethral fossa with negative result in each case;
- (e)(i)The donor stallion was continuously resident on the semen collection centre for 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 during that time into direct contact with equidae of lower health status than the donor stallion.

The tests described in **point (d)** have been carried out on samples taken *Insert date* in the table in **point (f) of AH/A725B** to the first semen collection and at least 14 days following the date of the commencement of the residence period of at least 30 days.

(e)(ii)The donor stallion was resident on the semen collection centre for at least 30 days prior to the date of the first collection and during the period of collection of the semen described above, but has left the centre under the responsibility of the centre veterinarian for a continuous period of less than 14 days, or other equidae on the collection centre came into direct contact with equidae of lower health status.

The tests described in **point (d)** have been carried out on samples taken *Insert date* in the table in **point (f) of AH/A725B** prior to the date of the first semen collection of the breeding season or collection period in the year the semen described above was collected and at least 14 days following the date of the commencement of the residence period of at least 30 days,

The test described in **point** (d)(i) for equine infectious anaemia was last carried out on a sample of blood taken *Insert date* in the table in **point** (f) of AH/A725B) not more than 90 days before the semen described above was collected;

The test described in **point (d)(iii)** for contagious equine metritis was last carried out on samples taken *Insert date* in the table in **point (f) of AH/A725B** more than 60 days before the semen described above was collected.

(e)(ii) option 1

one of the tests described in **point (d)(ii)** for equine viral arteritis was last carried out on a sample taken *Insert date* in the table in **point (f) of AH/A725B** not more than 30 days before the semen described above was collected.

(e)(ii) option 2

a virus isolation test for equine viral arteritis was carried out with negative result on an aliquot of the entire semen of the donor stallion taken *Insert date* in the table in **point (f) of AH/A725B** not more than 6 months before the semen described above was collected and a blood sample taken on the same date *Insert date* in the table in **point (f) of AH/A725B** reacted positive in a serum neutralisation test for equine viral arteritis at a serum dilution of more than one in four,

(e)(iii)) The tests described in **point** (d) have been carried out on samples taken *Insert date* in the table in **point** (f) of AH/A725B prior to the date of the first semen collection of the breeding season or collection period in the year the semen described above was collected, and the tests described in **point** (d) have been carried out on samples taken *Insert date* in the table in **point** (f) of AH/A725B between 14 and 90 days after the collection of the semen described above.

Guidance for the completion of the table

Abbreviations:

VS Vesicular stomatitis (VS) testing if required in accordance with point (b) of AH/A781B

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

EIA-2 EIA testing second occasion

EVA-B1 Equine viral arteritis (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 Cem testing first occasion second sample taken 7 days after CEM-11

CEM-21 Cem testing second occasion first sample

Instructions:

CEM-22

For each semen identified in column A in correspondence with Box I.28, the test programme (points (e)(i) of AH/A725B, 5(e)(ii) of AH/A725B and/or (e)(iii) of AH/A725B) shall be specified in column B, and columns C and D shall be completed with the dates required.

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

The dates when samples were taken for laboratory testing prior to the first collection of the semen described above as required in points (e)(i) of AH/A725B, (e)(ii) of AH/A725B and (e)(iii) of AH/A725B, shall be entered in the upper line of columns 6 to 10 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.

The dates when samples were taken for repeat laboratory testing as required in accordance with point **(e)(ii) of AH/A725B** or **(e)(iii) of AH/A725B** are entered in the lower line of columns 6 to 10 in table, this being the boxes EIA-2, EVA-B2 or EVA-S2 and CEM-21 and CEM-22 in the example below.

Identification of semen	Test programme	Start date: Donor residence	Start date: Semen collection	Date of samplin g for health tests: VS point (b) of AH/A78	Date of samplin g for health tests: EIA point (d)(i) of AH/A72 5B	Date of samplin g for health tests: EVA point (d)(ii) of AH/A72 5B Blood sample	Date of samplin g for health tests: EVA point (d)(ii) of AH/A72 5B Semen sample	Date of samplin g for health tests: CEM point (d)(iii) of AH/A72 5B 1. sample	Date of samplin g for health tests: CEM d)(iii) of AH/A72 5B 2. sample
			_	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	EIA-1	EVA-B1	EVA-S1	CEM-11	CEM-12
A	В	С	D	VS	EIA-2	EVA-B2	EVA-S2	CEM-21	CEM-22

AH/A781B Animal requirements (freedom from disease)

(a) regionalisation in accordance with Article 13 of Directive 2009/156 (†).

For African Horse Sickness: It is not considered to be infected in accordance with Article 5(2)(a) and (b) of Directive 2009/156.

 (\dagger) .

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(b)(ii) The donor stallions were subjected were subjected to a virus neutralisation test for vesicular stomatitis (VS) carried out with negative result at a serum dilution of 1 in 12 on a blood sample taken (insert date in the table in **point** (f) of AH/A725B (Guidance on how to complete table in **point** f can be found at the bottom of the notes for completion) within 14 days prior to entering the centre.

AH/P463 Product requirements

Point (b) - Insert names and concentrations.

AH/P551D Product requirements (storage and transport)

GB requirements

- (a)In compliance with the requirements of Chapters II(I)(1) and III(I) of Annex D to Directive 92/65;
- (b) In accordance with point 1.4 of Chapter III(I) of Annex D to Directive 92/65 and bearing the number indicated in Box I.23.
- (†) The document(s) referred to above can be found at:

<u>EU and EFTA countries approved to export animals and animal products to Great Britain</u> (Available at: https://www.data.gov.uk/dataset/4698a65d-1a3b-42d1-981e-df869e04185b/eu-and-efta-countries-approved-to-export-animals-and-animal-products-to-great-britain)

Non-EU countries approved to export animals and animal products to Great Britain (Available at: https://www.data.gov.uk/dataset/b92627b0-dd7b-4e1d-ba36-e25424f55eeb/non-eu-countries-approved-to-export-animals-and-animal-products-to-great-britain)

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