# Model health certificate for blood products not intended for human consumption that could be used as feed material (BP)

# GBHC500 v1.0 May-23

Part I. Details of	of dispatch	ned cons	ignmen	t				
I.1 Consignor		I.2 Certificate reference no.		I.3 Central competent authority				
Name:								
Address:			<b>I.2.a</b> N	ot in use		I.4 Loca	I competent aut	hority
				J J.				
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	on of	Code	I.9 Country of	ISO	I.10 Region of	Code
origin	code	origi		Oode	destination	code	destination	Jouc
I.11 Place of or	igin				I.12 Place of des	stination		
Name:					Custom warehouse			
Approval number	er:				Name:			
Address:					Approval number:			
					Address:			
Name:								
Approval number	er:							
Address:								
Name:								
Approval number	er:							
Address:								
I.13 Place of lo	ading				I.14 Date of depart	arture		
I.15 Means of transport			I.16 Entry BCP					
Aeroplane								
Ship								
☐ Railway wagon								
☐ Road vehicle				I.17 Not in use				
Other								
Identification:								
Documentation references:								

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

I.18 Description of commodity				
I.19 Commodity code (HS co	de) I.21 Temperature	e of products	I.23 Seal / Container	No.
	☐ Ambient			
	☐ Chilled			
	Frozen			4
I.20 Quantity	I.22 Number of p	oackages	I.24 Type of packagi	ng
I.25 Commodity certified for	,			
Animal feedingstuff				
☐ Manufacture of petfood				
☐ Technical use				
I.26  For transit through Great Britain to third country		I.27  For import or admission into Great Britain		
Third country	ISO Code			
I.28 Identification of the com	modities	. ( )		
Species (Scientific name)	Nature of commodity		er of establishments ecturing plant	Batch number
	0			
	CV			

# Part II. Certification

# **Animal Health**

I, the undersigned official veterinarian, declare that I have read and understood the requirements of the relevant GB regulations and certify that the blood products described in Part I of this certificate consist of blood products that satisfy the health requirements below:

# AH/E102 Establishment requirements (plant)

have been prepared and stored in a plant approved and supervised by the competent authority in accordance with GB requirements;

# AH/P010 Product requirements (segregation)

the product has undergone all precautions to avoid contamination with pathogenic agents after treatment;

(\*)**AND** [has been stored in dry warehouse conditions under room temperature for a period of at least 6 weeks;]

# **AH/P104 Product requirements (composition)**

have been prepared exclusively with the following animal by-products as set out in the notes for completion  $(^{\circ})[A]$  and/or  $(^{\circ})[B]$ ;

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

# **AH/P151A Product requirements**

consist exclusively of blood products not intended for human consumption;

## AH/P506 Packaging and labelling

the end product was:

(\*) **EITHER** [packaged in new or sterilised bags;]

(\*) **OR** [transported in bulk in containers or other means of transport that were thoroughly cleaned and disinfected with a disinfectant approved by the competent authority before use;]

and which bear labels indicating 'NOT FOR HUMAN CONSUMPTION';

#### AH/P550B Storage

the end product was stored in enclosed storage;

#### AH/P700 Product treatment

in order to inactivate pathogenic agents, have been submitted:

(\*) EITHER (a) [to processing in accordance with processing method ...... in accordance with GB requirements;]

(\*) **OR** (b) [to a method and parameters which ensure that the product complies with the GB microbiological requirements;]

(\*) OR (c) [to treatment for blood products as set out in the notes for completion and meets the relevant requirements;]

#### AH/P800B Testing

the competent authority examined a random sample of the products immediately prior to dispatch and found it to comply with microbiological GB requirements for *Salmonella* and *Enterobacteriaceae*;

#### AH/P901 Product requirements (statement)

the product(s) described above contain or are derived from animal-by products of nonruminant origin, and are, according to the statement of the Consignor referred to in box reference I.1,

(\*) **EITHER** [not intended for the production of feed for farmed animals, other than fur animals;]

[intended for the production of feed for non-ruminant farmed animals, other than fur animals, and the Consignor has undertaken to ensure that the border control post of entry will be provided with the results of the analyses carried out in accordance with the methods set out in GB legislation;]

# AH/D200A TSE (scrapie)

the animal by-products described above:

(\*) **EITHER** [does not contain ovine or caprine milk or milk products or is not intended for feed for farmed animals, other than fur animals;]

(\*) **OR** [contains ovine or caprine milk or milk products intended as feed for farmed animals, other than fur animals, and the milk or milk products:

- (a) are derived from animals from countries which meet GB requirements in regards to scrapie controls;
- **(b)** originate from holdings where no official restrictions are imposed due to a suspicion of TSE;
- (c) originate from holdings where no case of classical scrapie has been diagnosed during the period of the preceding seven years or, following the confirmation of a case of classical scrapie:

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

(\*) EITHER [all ovine and caprine animals on the holding have been killed and

destroyed or slaughtered, except for animals which meet GB

requirements;]

(\*)OR [all animals in which classical scrapie was confirmed have been killed and

destroyed, and the holding has been subjected to TSE monitoring which

meets GB requirements;]]

#### **Public Health**

# PH/D011A Bovine spongiform encephalopathy (BSE)

the products described in Part I

(\*) **EITHER** [come from other ruminants than bovine, ovine or caprine animals;]

(\*) **OR** [come from bovine, ovine or caprine material:

(\*) **EITHER** [(a) derived from animals that were born, continuously reared and slaughtered in a country or region with a negligible BSE risk as set out in a document published on GOV.UK in accordance with GB regulations;]

(\*) **OR** [(b) that does not contain and is not derived from:

- (i) specified risk material and mechanically separated meat, in compliance with GB regulations;
- (ii) animal by-product or derived product obtained from animals which have not been killed in compliance with GB regulations in regards laceration of certain tissues after stunning;]]

Official Veterinarian / Official Inspector		
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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<sup>(\*)</sup> Keep as appropriate.

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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.6:	Person responsible for the consignment in Great Britain: this box is required to
	be filled in only if it is a certificate for a commodity to be transited through Great
	Britain; it may be filled in if the certificate is for a commodity that is to be imported

into Great Britain.

Box reference I.12: Place of destination: this box is to be filled in only if it is a certificate for transit

commodity. Products in transit may only be stored in free zones, free

warehouses and custom warehouses.

Box reference I.15: Registration number (railway wagons or container and lorries), flight number

(aircraft) or name (ship) is to be provided. In the case of unloading and reloading, the consignor must inform the border control post of the point of entry into Great

Britain.

Box reference I.19: Use the appropriate Harmonized System (HS) code under the following heading:

05.11.91, 05.11.99, 35.02 or 35.04.

Box reference I.23: For bulk containers, the container number and the seal number (if applicable)

must be included.

Box reference I.25: Technical use: any use other than feeding of farmed animals, other than fur

animals, and the production or manufacturing of pet food.

Box reference I.26 and I.27: Fill in according to whether it is a transit or an import certificate.

Box reference I.28: Species: select from the following: Aves, Ruminantia, Suidae, Mammalia other

than Ruminantia or Suidae, Pesca, Reptilia.

#### Part II

#### **Animal Health**

By signing this certificate, you, the official veterinarian, are certifying that you have read and understood Regulation (EC) No 1069/2009 and Commission Regulation (EU) No 142/2011.

# AH/E102 Establishment requirements (plant)

The establishment is approved and supervised by the competent authority in accordance with Article 24 of Regulation (EC) No 1069/2009.

#### AH/P010 Product requirements (segregation)

The 'and' attestation is applicable in the case of blood products, including spray dried blood and blood plasma of porcine origin intended for the feeding of porcine animals.

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# Blood products (BP) GBHC500

# **AH/P104 Product requirements (composition)**

- **A:** Blood of slaughtered animals, which is fit for human consumption in accordance with retained EU law, but which is not intended for human consumption for commercial reasons.
- **B:** Blood of slaughtered animals, which has been rejected as unfit for human consumption in accordance with retained EU law, but which did not show any signs of diseases communicable to humans or animals, which has been derived from carcases that have been slaughtered in a slaughterhouse and which were considered fit for human consumption following an ante-mortem inspection in accordance with retained EU law.

# **AH/P151A Product requirements**

No further notes for completion.

## AH/P506 Packaging and labelling

No further notes for completion.

# AH/P550B Storage

No further notes for completion.

#### AH/P700 Product treatment

- (a) Insert method 1 to 5 or method 7 as applicable, as set out in Chapter III of Annex IV to Regulation (EU) to 142/2011.
- (b) Microbiological requirements as set out in Chapter I of Annex X to Regulation (EU) No 142/2011.
- (c) In the case of blood products, including spray dried blood and blood plasma, of porcine origin intended for the feeding of porcine animals, to a heat treatment at a temperature of at least 80°C throughout the substance and the dry blood and blood plasma does not contain more than 8% w/w moisture with a water activity (Aw) of less than 0.60.

#### AH/P800B Testing

The products must be examined immediately prior to dispatch under the responsibility of the competent authority by taking a random sample during or on removal from storage which complies with the following standards:

- Salmonella: absence in 25 g: n=5, c=0, m=0, M=0
- Enterobacteriaceae: n=5, c=2, m=10, M=300 in 1 gram

#### Where:

- n = number of samples to be tested;
- m = threshold value for the number of bacteria; the result is considered satisfactory if the number of bacteria in all samples does not exceed m;
- M = maximum value for the number of bacteria; the result is considered unsatisfactory if the number of bacteria in one or more samples is M or more; and
- c = number of samples the bacterial count of which may be between m and M, the sample still being considered acceptable if the bacterial count of the other samples is m or less.

#### AH/P901 Product requirements (statement)

## 'or' attestation:

The person responsible for the load referred to in box reference I.6 must ensure that, if the blood products described in this health certificate are intended to be used for the production of feed for non-ruminant farmed animals, other than fur animals the consignment must be analysed, in accordance with the methods set out in Annex VI to Regulation (EC) No 152/2009, in order to verify the absence of unauthorised constituents of animal origin. The information on the result of such analysis must be attached to this health certificate when presenting the consignment at a border control post of Great Britain.

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# Blood products (BP) GBHC500

# AH/D200A TSE (scrapie)

Where the animal by-products described above contains milk or milk products of ovine or caprine animal origin and is intended for feed for farmed animals, other than fur animals, they must:

- (a) be derived from ovine and caprine animals which were kept continuously since birth in a country where the following conditions are fulfilled:
  - (i) classical scrapie is compulsorily notifiable;
  - (ii) an awareness, surveillance and monitoring system is in place for classical scrapie;
  - (iii) official restrictions apply to holdings of ovine or caprine animals in the case of a suspicion of TSE or the confirmation of classical scrapie;
  - (iv) ovine and caprine animals affected with classical scrapie are killed and destroyed;
  - (v) the feeding to ovine and caprine animals of meat-and-bone meal or greaves, as defined in the Terrestrial Animal Health Code of the World Organisation for Animal Health (WOAH (formerly OIE)), of ruminant origin has been banned and effectively enforced in the whole country for a period of at least the preceding seven years.
- **(b)** originate from holdings where no official restrictions are imposed due to a suspicion of TSE.
- (c) originate from holdings where no case of classical scrapie has been diagnosed during the period of the preceding seven years or, following the confirmation of a case of classical scrapie:
  - **EITHER**

all ovine and caprine animals on the holding have been killed and destroyed or slaughtered, except for breeding rams of the ARR/ARR genotype, breeding ewes carrying at least one ARR allele and no VRQ allele and other ovine animals carrying at least one ARR allele;

OR

all animals in which classical scrapie was confirmed have been killed and destroyed, and the holding has been subjected for a period of at least two years since the date of confirmation of the last classical scrapie case to intensified TSE monitoring, including testing with negative results for the presence of TSE in accordance with the laboratory methods set out in point 3.2 of Chapter C of Annex X to Regulation (EC) No 999/2001, of all of the following animals which are over the age of 18 months, except ovine animals of the ARR/ARR genotype:

- animals which have been slaughtered for human consumption; and
- animals which have died or been killed on the holding but which were not killed in the framework of a disease eradication campaign.

#### **Public Health**

# PH/D011A Bovine spongiform encephalopathy (BSE)

The products described in Part I of the certificate:

**EITHER** 

are derived from other ruminants than bovine, ovine or caprine animals.

OR

are derived from bovine, ovine or caprine animals, and do not contain and are not derived from:

**EITHER** 

(a) bovine, ovine and caprine materials other than those derived from animals born, continuously reared and slaughtered in a country or region classified as posing a negligible BSE risk as set out in a document relating to 'BSE risk status' published on GOV.UK, in accordance with Regulation (EC) No 999/2001(‡):

OR

- **(b)** the following:
  - (i) specified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001 of the European Parliament and of the Council and mechanically separated meat obtained from bones of bovine, ovine or caprine animals, except from those animals that were born,

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- continuously reared and slaughtered in a country or region classified as posing a negligible BSE risk as set out in a document relating to 'BSE risk status' published on GOV.UK, in accordance with Regulation (EC) No 999/2001<sup>(‡)</sup>, in which there have been no indigenous BSE cases.
- (ii) animal by-product or derived product obtained from bovine, ovine or caprine animals which have been killed, after stunning, by laceration of the central nervous tissue by means of an elongated rod-shaped instrument introduced into the cranial cavity, or by means of gas injected into the cranial cavity, except for those animals that were born, continuously reared and slaughtered in a country or region classified as posing a negligible BSE risk as set out in a document relating to 'BSE risk status' published on GOV.UK, in accordance with Regulation (EC) No 999/2001.<sup>(‡)</sup>
- (‡) A document relating to the 'Bovine Spongiform Encephalopathy (BSE) risk status' of approved trading partners published by the Secretary of State, with the consent of the Scottish and Welsh Ministers, can be found at:

<u>Animal health status of countries approved to export animals and animal products to Great Britain -</u> data.gov.uk

(Available at: https://www.data.gov.uk/dataset/b7712d2e-debb-4996-8e79-d27ca7492a00/animal-health-status-of-countries-approved-to-export-animals-and-animal-products-to-great-britain)

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