Model health certificate for untreated blood products, excluding those of equidae, for the manufacture of derived products for purposes outside the feed chain for farmed animals (BP-U) GBHC502 v1.1 Aug-23

Part I. Details of the dispatched consignment I.1 Consignor I.2 Certificate reference no. Name: Address: Tel: I.2.a I.3 Central competent I.4 Local competent authority authority Not in **APHIS-VS** use I.5 Consignee 1.6 Person responsible for the load in Great Name: Britain Address: Name: Tel: Address: Tel: ISO code I.7 Country of origin I.8 Region of Code I.9 Country of ISO I.10 Region of Code origin destination code destination 1.12 Place of destination I.11 Place of origin Name: Custom warehouse Approval number: Address: Name: Approval number: Address: Name: Approval number: Address: Name: Approval number: Address: I.13 Place of loading I.14 Date of departure I.15 Means of transport I.16 Entry BCP ☐ Aeroplane Ship I.17 Not in use Railway wagon Road vehicle Other Identification: Documentation references:

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

I.18 Description of commodity							
I.19 Commodity code (HS code)	I.21 Temperature of products	S I.23 Seal / Container No.					
, ,	Ambient Chilled						
	_						
	Frozen						
I.20 Quantity	I.22 Number of packages		I.24 Type of packaging				
n.20 Quantity	n== mannon or puonageo		in 1 1 y po or puckaging				
I.25 Commodity certified for	Fechnical use						
I.26 For transit through Great Brita	in to third country	1.27	For import or admission into				
	_		Britain				
Third country	ISO Code						
I.28 Identification of the commodities							
Species (Scientific name)	Approval number of establishments / Manufacturing plant		Batch number				

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II.a. Certificate reference no.



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/T111 Territory requirements

- (*)[(a) in the case of blood products obtained from animals belonging to the taxa *Artiodactyla*, *Perissodactyla* and *Proboscidea*, including crossbreds between species of those taxa, the blood was collected in a country or region where no case of rinderpest, peste des petits ruminants and Rift Valley fever has been recorded and in which vaccination has not been carried out in accordance with GB requirements set out in the notes for completion; and
- (*)[(b) in the case of animals other than *Suidae* and *Tayassuidae*, in third countries or regions in which:

 (*)EITHER [no case of (*)[vesicular stomatitis] and (*)[bluetongue] (including the presence of serope
 - (*) **EITHER** [no case of (*)[vesicular stomatitis] and (*)[bluetongue] (including the presence of seropositive animals) has been recorded in accordance with GB requirements as set out in the notes for completion;]
 - (*) OR [(*) [vesicular stomatitis] and (*) [bluetongue] seropositive animals are present;]]
- (*)[(c) in the case of *Suidae* and *Tayassuidae*, in third countries or regions in which no case of swine vesicular disease, classical swine fever and African swine fever has been recorded for a period of at least the preceding 12 months and vaccination has not been carried out against those diseases for a period of at least the preceding 12 months in the susceptible species and:
 - (*) **EITHER** [no case of vesicular stomatitis (including the presence of seropositive animals) has been recorded in accordance with GB requirements as set out in the notes for completion;]
 - (*) **OR** [vesicular stomatitis seropositive animals are present;]]

AH/T112 Territory requirements

AH/E103 Establishment requirements

have been prepared and stored in a plant and supervised by the competent authority or in the establishment of collection, exclusively with the following animal by-products as set out in the notes for completion: (*)[A] (*)[B] (*)[C] (*)[D] (*)[E] (*)[G];

AH/E302 Establishment requirements (slaughterhouse)

the blood, that these products were manufactured from, was collected in slaughterhouses approved in accordance with GB regulations, in slaughterhouses approved and supervised by the competent authority of the country of collection or from live animals in facilities approved and supervised by the competent authority of the country of collection;

AH/P011 Product requirements (segregation)

all precautions were taken to avoid contamination of the products with pathogenic agents during transport;

AH/P151B Product requirements

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II.a.	Certifica	te referenc	e no.

N.b.

consist exclusively of blood products not intended for human or animal consumption;

AH/P509 Packaging and labelling

the products were:

(*) **EITHER** [packed in new or sterilised bags or bottles;]

(*) 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 the outer packaging or containers bear labels indicating 'NOT FOR HUMAN OR ANIMAL CONSUMPTION';

AH/P550A Storage

the product was stored in enclosed storage;

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:	_				

The signature and the stamp must be in a different colour to that of the printing.

^(*) Keep as appropriate.