# Model health certificate for animal by-products to be used for purposes outside the feed chain or for trade samples (ABPO) GBHC580 v1.1 Aug-23

Part I. Details of the	dispatche	d consignment				
I.1 Consignor			I.2 Certificate reference no.			
Name: Address:						
Tel:			10.5	l 2 Camtual agreementary	4 1 4 1	l samustant
101.			I.2.a	I.3 Central competen authority	auth	
			Not in	authority	auti	Officy
			use	APHIS-VS		
I.5 Consignee				1.6 Person responsi	hle for the	e load in Great
Name:				Britain	DIC IOI LII	, load III Olcat
Address:				Name:		
Tel:				Address:		
				Tel:		
I.7 Country of origin	ISO code	I.8 Region of	Code	I.9 Country of	ISO	I.10 Region of Code destination
		origin	110.4	destination	code	
USA	US	US	US-1			
I.11 Place of origin Name:				1.12 Place of destina	ation	
Approval number:				Custom warehouse		
Address:				Name:		
				Approval number: Address:		
Name:						
Approval number:						
Address:						
Name:						
Approval number: Address:						
I.13 Place of loading				I.14 Date of departu	ro	
1.13 Flace of loading				1.14 Date of departu	16	
I.15 Means of transpo	ort			I.16 Entry BCP		
☐ Aeroplane	☐ Shi	р				
☐ Railway wagon	Roa	ad vehicle		I.17 Not in use		
Other				•		
Identification:						
Doormont-ti						
Documentation referer	ices:					

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Animal by-products to be used for purposes outside the feed chain or for trade samples (ABPO) GBHC580

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

I.18 Description	of commodity					
I.19 Commodity code (HS code)			I.21 Temperature of products		I.23 Seal / Container No.	
		Ambient (	Chilled			
L20 Quantity		I.22 Number of pack	/2000	I 24 Type	of nackaging	
I.20 Quantity		1.22 Number of pack	kages	1.24 Type 0	of packaging	
1.25 Commodity		X Technical use	127 V For it	mnost or admic	paian into Creat Britain	
Third country	through Great Britis	ISO Code			sion into Great Britain	
,						
I.28 Identification	n of the commodi	ties				
Species (Scientific name)	Nature of commodity	Approval number of establishments / Manufacturing plant	Number of packages	Net weight	Batch number	

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II.a. Certificate reference no.	N.b.	
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# Part II. Certification

#### **Animal Health**

- I, the undersigned official veterinarian, declare that I am aware of the relevant requirements of the GB regulations, and certify that the animal by-products described in Part I of this certificate:
- (\*) **EITHER** [are trade samples which consist of animal by-products intended for particular studies or analyses as referred to in the definition of trade samples in GB regulations, that bear the label 'TRADE SAMPLE NOT FOR HUMAN CONSUMPTION';]
- (\*) **OR** [satisfy the animal health requirements below;]

## AH/T109 Territory requirements

- (the animal by-products in this consignment come from animals that have been obtained in the country, territory or part thereof referred to in AH/T, where vaccination programmes against footand-mouth disease are regularly carried out and officially controlled in domestic bovine animals;
- (\*) AND/OR [the animal by products in this consignment consist of animal by products derived from offal or deboned meat;]

## AH/T501 Territory requirements

have b	eer
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- - (i) have remained in that third country, territory or part thereof eligible to export fresh meat to Great Britain since birth or for a period of at least the preceding three months before the date of slaughter; and/or
  - (ii) were killed in the wild in that third country, territory or part thereof;]
- (c) [derived from eggs, milk, rodents, lagomorphs, or aquatic animals or terrestrial or aquatic invertebrates:]

# AH/E411 Establishment requirements

(\*)[in the case of materials other than materials derived from eggs, milk, rodents, lagomorphs, wool grease, aquatic animals, terrestrial or aquatic invertebrates and unprocessed furs, have been obtained from animals: (\*) **EITHER** [(a) coming from holdings:

- (i) where, for the following diseases for which the animals are susceptible, there has not been any case/outbreak of rinderpest, swine vesicular disease, Newcastle disease or highly pathogenic avian influenza during the period of the preceding 30 days, nor of classical or African swine fever during the period of the preceding 40 days; nor in the holdings situated in their vicinity within a 10 km radius, during the period of the preceding 30 days; and
- (ii) where there has not been any case/outbreak of foot-and-mouth disease during the period of the preceding 60 days, nor in the holdings situated in their vicinity within a 25 km radius, during the period of the preceding 30 days; and
- (b) which:
  - (i) were not killed to eradicate any epizootic disease;
  - (ii) remained on their holdings of origin for a period of at least 40 days before the date of departure and which were transported directly to the slaughterhouse without contact with other animals which did not comply with the same health conditions;
  - (iii) at the slaughterhouse, passed the ante-mortem health inspection during the period of 24 hours before the time of slaughter and showed no evidence of the diseases referred to above for which the animals are susceptible; and

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(iv) were handled in the slaughterhouse before and at the time of slaughter or killing in accordance with GB requirements;]

(\*) Captured and killed in the wild in an area:

- (i) where within a 25 km radius there has been no case/outbreak of any of the following diseases for which the animals are susceptible: foot and mouth disease, rinderpest, Newcastle disease or highly pathogenic avian influenza during the period of the preceding 30 days nor of classical or African swine fever during the period of the preceding 40 days; and
- (ii) that is situated at a distance that exceeds 20 km from the borders separating another territory of a third country or part thereof, which is not authorised at these dates for the exportation of such material to Great Britain; and
- (b) which, after killing, were transported within a period of 12 hours for chilling either to a collection centre and immediately afterwards to a game establishment, or directly to a game establishment;]]

# AH/E412A Establishment requirements

(\*)[in the case of materials other than materials derived from fish or invertebrates caught in the wild, have been obtained in an establishment around which, within a radius of 10 km, there has been no case/outbreak of diseases referred to in AH/E411 for which the animals are susceptible during a period of the preceding 30 days or, in the event of a case/outbreak of one of those diseases, the preparation of raw material for exportation to Great Britain was authorised only after the removal of all meat, and the total cleaning and disinfection of the establishment under the control of an official veterinarian;]

#### AH/P003 Production requirements

have been obtained and prepared without contact with other materials that do not comply with the conditions required above, and have been handled so as to avoid contamination with pathogenic agents;

# AH/P101A Product requirements (composition)

consist only of the following Category 3 materials as set out in the notes for completion: (\*)[A] (\*)[B] (\*)[C] (\*)[D] (\*)[E] (\*

# AH/P504 Product requirements

have been packed in new packaging which prevents any leakage or in packaging which has been cleaned and disinfected before use and, in the case of consignments shipped other than via parcel post, in containers sealed under the responsibility of the competent authority, bearing the label indicating 'ANIMAL BY-PRODUCTS ONLY FOR THE MANUFACTURE OF DERIVED PRODUCTS FOR USES OUTSIDE THE FEED CHAIN' and the name and address of the establishment of destination in Great Britain:

## **AH/P611A Product requirements**

have been deep-frozen at the plant of origin or have been preserved in accordance with GB regulations in such a way that they will not spoil between the time of dispatch and the time of delivery to the plant of destination;

#### 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 scrapic 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:

<del>(*)</del> EITHER	[all ovine and caprine animals on the holding have been killed and destroyed or
	slaughtered, except for animals which meet GB requirements;]
<del>(*)</del> 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:

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II.a. Certificate reference no.	М.b.	7

# **Public Health**

# PH/D011A Bovine spongiform encephalopathy (BSE)

the products described in Part I

(\*)OR

(\*) **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;]

[(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;]]

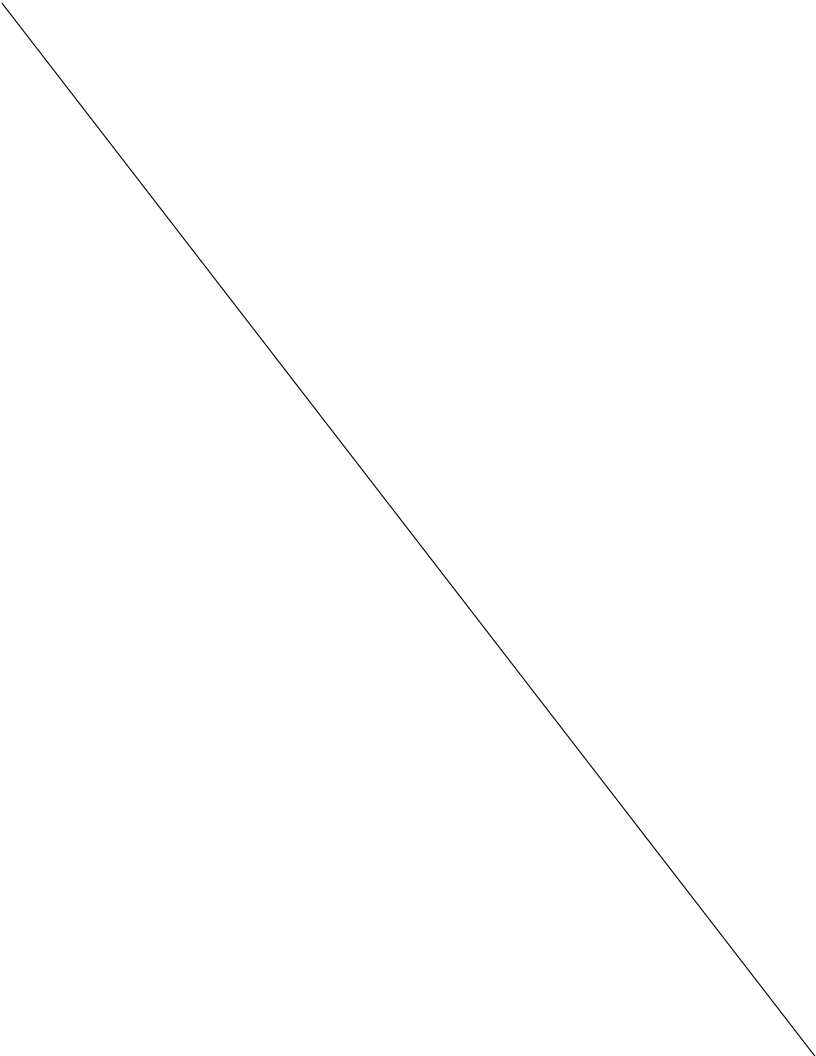
Official Veterinarian / Official Inspector	
By signing this certificate, I certify that the renotes for completion have been met.	quirements laid out above and in the accompanying
Name (in capital letters):	Qualification and title:
Date:	Signature:
Stamp:	

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

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



# 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.11: In the case of consignments for trade samples or analyses: indicate the name and address of the establishment only.

Box reference I.11 and I.12: *Approval number*: the registration number of the establishment or plant, which has been issued by the competent authority.

Box reference I.12: Place of destination: this box is to be filled in:

- products for the manufacture of derived products for uses outside the feed chain: only if it is a certificate for a transit commodity. Products in transit may only be stored in free zones, free warehouses and custom warehouses.

- products for trade samples or analyses: the plant in Great Britain indicated in the authorisation of the competent authority where appropriate.

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 headings: 04.01; 04.02; 04.03; 04.04; 04.08; 05.05; 05.06, 05.07; 05.11.91; 05.11.99, 23.01 or 30.01.

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.25: For the purposes of the certificate, 'technical use' includes use as a trade sample.

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

Box reference I.28: - Products for the manufacture of derived products for uses outside the feed chain: *Manufacturing plant*: provide the veterinary control number of the approved establishment.

- Products for the particular technological studies or analyses: the plant in Great Britain indicated in the authorisation of the competent authority where appropriate.

- Species: select from the following: Aves, Ruminantia, Suidae, Mammalia other than Ruminantia or Suidae, Pesca, Mollusca, Crustacea, invertebrates other than Mollusca and Crustacea.

#### 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, and in particular Chapter II of Annex XIV thereto.

Definition of trade samples is in point 39 of Annex I to Regulation (EU) No 142/2011.

# **AH/T109 Territory requirements**

This attestation is only required when supplementary guarantees are to be provided where the material of domestic ruminants originated in the territory of a South American or South African country or part thereof from where only maturated and deboned fresh meat of domestic ruminants for human consumption is authorised for exportation to Great Britain. The whole masseter muscles of bovine animals, incised in accordance with the requirements of Article 19, paragraph 1(a) to (EU) Regulation 2019/627 are also permitted.

The 'either' attestation is only for certain South American countries.

The 'and/or' attestation is only for certain South American and South African countries.

# **AH/T501 Territory requirements**

Insert the name and ISO code number of the exporting country as laid down in:

- a document relating to 'fresh meat of ungulates' published on GOV.UK, in accordance with Commission Regulation (EU) No 206/2010<sup>(†)</sup>
- a document relating to 'poultry and poultry products' published on GOV.UK, in accordance with Commission Regulation (EC) No 798/2008<sup>(†)</sup>
- a document relating to 'meat of wild leporidae, certain wild land mammals and of farmed rabbits' published on GOV.UK, in accordance with Commission Regulation (EC) No 119/2009<sup>(†)</sup>

In addition, the ISO code of territories and parts thereof referred to in the documents published in accordance with Regulations (EU) No 206/2010, (EC) No 798/2008 and (EC) No 119/2009 referred to in this note (where applicable for the susceptible species concerned) must be included where applicable.<sup>(†)</sup>

Subparagraph (b)(ii) refers only for countries from where the game meat intended for human consumption of the same animal species is authorised for importation into Great Britain.

# AH/E411 Establishment requirements

Subparagraph (b)(iv) refers in compliance with provisions of retained EU law and complied with requirements at least equivalent to those laid down in Chapters II and III of Council Regulation (EC) No 1099/2009.

## AH/E412A Establishment requirements

No further notes for completion.

# AH/P003 Production requirements

No further notes for completion.

# AH/P101A Product requirements (composition)

- A: Carcases and parts of animals slaughtered or, in the case of game, bodies or parts of animals killed which were deemed fit for human consumption in accordance with retained EU law until irreversibly declared as animal by-products for commercial reasons.
- **B:** Carcases and the following parts originating either from animals that were slaughtered in a slaughterhouse and were considered fit for slaughter for human consumption following an ante-mortem inspection or bodies and the following parts of animals from game killed for human consumption in accordance with retained EU law:
  - (i) carcases or bodies and parts of animals which were rejected as unfit for human consumption in accordance with retained EU law, but which did not show any signs of disease communicable to humans or animals:

- (ii) heads of poultry;
- (iii) hides and skins, including trimmings and splitting thereof, horns and feet, including the phalanges and the carpus and metacarpus bones, tarsus and metatarsus bones;
- (iv) pig bristles;
- (v) feathers.
- C: Animal by-products from poultry and lagomorphs slaughtered on the farm as referred to in Article 1(3)(d) of Regulation (EC) No 853/2004 of the European Parliament and of the Council, which did not show any signs of disease communicable to humans or animals.
- **D:** Blood of animals which did not show any signs of disease communicable through blood to humans or animals, obtained from animals that have been slaughtered in a slaughterhouse after having been considered fit for slaughter for human consumption following an ante-mortem inspection in accordance with GB regulations.
- **E:** Animal by products arising from the production of products intended for human consumption, including degreased bone, greaves and centrifuge or separator sludge from milk processing.
- F: Products of animal origin, or foodstuffs containing products of animal origin, which are no longer intended for human consumption for commercial reasons or due to problems of manufacturing or packaging defects or other defects from which no risk to public or animal health arises.
- G: Petfood and feedingstuffs of animal origin, or feedingstuffs containing animal by-products or derived products, which are no longer intended for feeding for commercial reasons or due to problems of manufacturing or packaging defects or other defects from which no risk to public or animal health arises.
- **H:** Blood, placenta, wool, feathers, hair, horns, hoof cuts and raw milk originating from live animals that did not show signs of any disease communicable through that product to humans or animals.
- **l:** Aquatic animals, and parts of such animals, except sea mammals, which did not show any signs of diseases communicable to humans or animals.
- **J:** Animal by-products from aquatic animals originating from establishments or plants manufacturing products for human consumption.
- **K:** The following material originating from animals which did not show any signs of disease communicable through that material to humans or animals:
  - (i) shells from shellfish with soft tissue or flesh;
  - (ii) the following originating from terrestrial animals:
    - hatchery by-products;
    - eggs
    - egg by-products, including egg shells;
  - (iii) day old chicks killed for commercial reasons.
- L: Animal by-products from aquatic or terrestrial invertebrates, other than species pathogenic to humans or animals.
- M: Animals and parts thereof of the zoological orders of Rodentia and Lagomorpha, except Category 1 material as referred to in Article 8(a)(iii), (iv) and (v) of Regulation (EC)No 1069/2009 and Category 2 material as referred to in Article 9(a) to (g) of that Regulation.
- N: Furs originating from dead animals that did not show clinical signs of any disease communicable through that product to humans or animals.

# AH/P504 Product requirements

No further notes for completion.

#### AH/P611A Product requirements

No further notes for completion.

# 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 scrapic has been diagnosed during the period of the preceding seven years or, following the confirmation of a case of classical scrapic:

EITHER

OR

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; 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 scrapic 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 909/2001, 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 cradication campaign.

#### **Public Health**

OR

## 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.

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, 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(i), 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.(1)

(†) The document(s) referred to above can be found at:

EU and EFTA countries approved to export animals and animal products to Great Britain

(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)

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# 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)

(‡) 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:

Animal health status of countries approved to export animals and animal products to Great Britain - 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)