COUNTRY: UNITED KINGDOM Part I: Details of dispatched consignment

Veterinary certificate to EU

I.1. Consignor Name		I.2. Certificate reference No		I.2.a.	
Address	I.3. Central Competent Authority				
	DEPARTMENT FOR ENVIRONMENT, FOOD AND RURAL AFFAIRS				
		I.4. Local Competent Authority			
Tel.					
I.5. Consignee		I.6. Person responsible for the load in the	ne EU		
Name Address		Name Address			
radioss		Address			
Postcode Tel.		Postcode Tel.			
	I.8. Region of origin, Code	I.9. Country of destination	ISO code	I.10. Region of destination	Code
UNITED KINGDOM GB					
I.11. Place of origin	A	I.12. Place of destination	Custom	warehouse	
Name Address	Approval number	Name			
		Address			
		Postal Code			
· ·		Approval number			
I.13. Place of loading		I.14. Date of departure			
I.15 Means of transport		I.16. Entry BIP in EU			
	hip Railway wagon	I.16. Entry BIP in EU			
Road vehicle	Other				
Identification:		I.17.			
Documentary references:					
I.18. Description of commodity		I.19. Commod	lity code (HS	S code)	
I.21. Temperature of product	1130	Quantity	122 N	Jumber of packages	
Ambient Chilled		Quantity	1.22. 1	vuiliber of packages	
I.23. Seal/Container No.			124 7	Type of packaging	
1.23. Sear/Container INO.			1.24.	Type of packaging	
I.25. Commodities certified for:					
Technical use					
I.26. For transit through EU to third country	JI.274	For import or admission into EU			
Third country	ISO code				
I.28. Identification of the commodities	<u> </u>				
	Approval number	ber of establishments			
Species (Scientific name)	Manuf	acturing plant		Batch number	
1				Y	

Untreated blood products, excluding those of equidae, for the manufacture of derived products for purposes outside the feed chain for farmed animals

II.a. Certificate reference No II. **Health information** I, the undersigned official veterinarian, declare that I have read and understood Regulation (EC) No 1069/2009 of the European Parliament and of the Council (1a), and in particular Article 8(c) and Article 8(d) and Article 10 thereof, and Commission Regulation (EU) No 142/2011 (1b), and in particular Chapter II of Annex XIV thereto, and certify the blood products described above consist of blood products that satisfy the health requirements below; II.1. Part II: Certification 11.2. they consist exclusively of blood products not intended for human or animal consumption; 11.3. they have been prepared and stored in a plant supervised by the competent authority or in the establishment of collection, exclusively with the following animal by-products: (2) either blood of slaughtered animals, which is fit for human consumption in accordance with Union legislation, but is not intended for human consumption for commercial reasons;] (2) and/or blood of slaughtered animals, which is rejected as unfit for human consumption in accordance with Union legislation, but which did not show any signs of diseases communicable to humans or animals, derived from carcases that have been slaughtered in a slaughterhouse and were considered fit for human consumption following an ante-mortem inspection in accordance with Union legislation;] blood of slaughtered animals, which did not show any signs of diseases communicable to (2) and/or humans or animals, obtained from animals that have been slaughtered in a slaughterhouse after having been considered fit for human consumption following an ante-mortem inspection in accordance with Union legislation;] (2) and/or blood and blood products derived from the production of products intended for human consumption;] (2) and/or blood and blood products originating from live animals that did not show signs of any disease communicable through that product to humans or animals;] animal by-products derived from animals which have been submitted to illegal treatment as (2) and/or defined in Article 1(2)(d) of Council Directive 96/22/EC (2a) or Article 2(b) of Council Directive 96/23/EC (2b);1 animal by-products containing residues of other substances and environmental contaminants (2) and/or listed in Group B(3) of Annex I to Directive 96/23/EC, if such residues exceed the permitted level laid down in Union legislation or, in the absence thereof, in national legislation;] 11.4. the blood, that such products were manufactured from, was collected in slaughterhouses approved in accordance with Union legislation, 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: (2) [II.5. 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 for a period of at least the preceding 12 months and in which vaccination has not been carried out against those diseases for a period of at least the preceding 12 months, and; [in third countries, territories or parts thereof (insert ISO country code in the case of a (2) either country, or codes (3) in the case of territories or parts thereof) where no case of foot-and-mouth disease has been recorded for a period of at least the preceding 12 months and in which vaccination has not been carried out against this disease for a period of at least the preceding 12 months, and] (2) or [in third countries, territories or parts thereof (insert ISO country code in the case of a country or codes (3) for territories or parts thereof) where no case of foot-and-mouth disease has been recorded for a period of at least the preceding 12 months and in which vaccination programmes against foot-and-mouth disease are being officially carried out and controlled in domestic ruminant animals for a period of at least the preceding 12 months (4), and]]

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Health information

II.

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II.a. Certificate reference No

(²) [II.5.1.	in the case of animals other than Suidae and Tayassuidae, in third countries or regions in which :									
	(²) either	has been	recorde	ular stomatitis and d for a period of a gainst those dise	t least the pr	eceding 12	months and	d in which v	vaccination h	
	(²) or	[vesicular	stomatit	is and bluetongue	e (²) seroposi	tive animals	are preser	nt (4);]]		
(²) [II.5.2.	classical swin	e fever and on has not b	African een car	suidae, in third c swine fever has ried out against th	been record	ed for a per	riod of at le	ast the pre	eceding 12 n	nonths
	(²) either	for a perio	d of at	ular stomatitis (inc least the preceding se for a period of	ng 12 months	s and in wh	ich vaccina			
	(2) or	[vesicular	stomatit	is seropositive ar	imals are pre	esent (4);]]]				
(²) [II.6.				rived from poultry on with code		an species t	the animals	and the p	roducts com	e from
	which has bee Animal Health			astle disease and	I highly path	ogenic avia	n influenza	as defined	d in the Ten	restrial
	which for a pe	riod of at lea	ast the p	preceding 12 mon	ths has not c	arried out v	accination a	against avia	an influenza,	
	where the animals from which the products are derived, have not been vaccinated against Newcastle disvaccines prepared from a Newcastle disease master strain showing a higher pathogenicity than lentog strains;]									
II.7.	the products w	vere:								
	(²) either	[packed in	new or	sterilised bags or	bottles,]					
	(²) or			alk in containers disinfectant appro					ughly cleane	ed and
	the outer pack	kaging or containers bear labels indicating 'NOT FOR HUMAN OR ANIMAL CONSUMPTION';			JMPTION';					
II.8.	the products w	were stored in enclosed storage;								
II.9.	all precautions	s were taker	n to avoi	d contamination of	of the produc	ts with path	ogenic ager	nts during t	ransport;	
(²) [II.10.	the untreated	ntreated blood products described above								
	(²) either	[is derived	from ot	her ruminants tha	n bovine, ov	ine or caprir	ne animals.]	l]		
	(²) or	[is derived	from bo	ovine, ovine or ca	orine animals	s and does r	not contain	and is not o	derived from:	:
		(²) either	continu	e, ovine and ca lously reared an ble BSE risk in ac	d slaughtere	ed in a co	untry or re	gion class		
		(²) or	[(a)	specified risk m No 999/2001 of						n (EC)
			(b)	mechanically se animals, except slaughtered in a accordance with no indigenous B	from those country or Commission	animals th region class	at were bo sified as po	orn, continu sing a neg	uously reare gligible BSE	d and risk in

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II.	Health information	II.a. Certificate reference No	II.b.

(c) 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 in accordance with Decision 2007/453/EC.]]]

Notes

Part I:

- Box reference I.6: Person responsible for the consignment in the European Union: this box is required to be filled in only if
 it is a certificate for a commodity that is to be transited through the European Union; it may be filled in if the certificate is
 for a commodity that is to be imported into the European Union.
- 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 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.
- 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 in the European Union, the consignor must inform the border inspection post of the point of entry into the European Union.
- Box I.19: use the appropriate Harmonized System (HS) code under the following headings: 05.11; 30.02 or 35.02.
- 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, Reptilian.

Part II:

- (1a) OJ L 300, 14.11.2009, p. 1.
- (1b) OJ L 54, 26.2.2011, p. 1.
- (2) Delete as appropriate.
- (^{2a}) OJ L 125, 23.5.1996, p. 3.
- (^{2b}) OJ L 125, 23.5.1996, p. 10.
- (3) Code of the territory as it appears in Part 1 of Annex II to Regulation (EU) No 206/2010 (OJ L 73, 20.3.2010, p. 1).
- (4) In this case following the veterinary checks provided for in Directive 97/78/EC (OJ L 24, 30.1.1998, p. 9), and in accordance with the conditions laid down in Article 8(4) of that Directive, the products must be transported directly to the establishment at the place of destination.

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II.	Health information	II.a. Certificate reference No	11.0.				
(5)	(5) Code of the territory as it appears in Part 1 of Annex I to Commission Regulation (EC) No 798/2008 (OJ L 226, 23.8.2008, p. 1).						
(⁶)	(⁶) OJ L 147, 31.5.2001, p. 1.						
(7)	OJ L 172, 30.6.2007, p. 84.						
_	 The signature and the stamp must be in a different colour to that of the printing. Note for the person responsible for the consignment in the European Union: this certificate is only for veterinary purposes and must accompany the consignment until it reaches the border inspection post of the point of entry into the European Union. 						
Officia	al veterinarian/Official inspector						
	Name (in capital letters): Date:	Qualification and title:					
	Date: Stamp:	Signature:					

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