CHAPTER 4(C)

Health certificate

For untreated blood products, excluding of equidae, for the manufacture of derived products for purposes outside the feed chain for farmed animals, intended for dispatch to or for transit through(2) the European Union

COU	NTRY: UNITED STATES	Veterinary certificate to EU		
	I.1. Consignor Name Address	I.2. Certificate reference No I.2.a.		
		I.3. Central competent authority APHIS-VS		
	Tel.	I.4. Local competent authority		
	I.5. Consignee Name Address	I.6. Person responsible for the load in EU Name Address		
	Postal code Tel.	Postal code Tel.		
gnmen				
d consig	I.7. Country of ISO code I.8. Region of Code origin	I.9. Country of ISO I.10. Region of Code destination code destination		
atche	I.11. Place of origin	I.12. Place of destination		
Part I : Details of dispatched consignment	Name Approval number Address	Custom warehouse Approval number Address		
I : Detail		Postal code		
Part				
	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 BIP in EU		
	Aeroplane □ Ship □ Railway wagon □ Road vehicle □ Other □	1.17.		
	Identification			
	Documentation references			
	I.18. Description of commodity	I.19. Commodity code (HS code)		
		I.20. Quantity		
	I.21. Temperature of product	I.22. Number of packages		
		1.22. Inditibet of packages		
	Ambient Chilled Frozen			

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COUNTRY: UNITED STATES

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		I.2. Certificate reference No	I.2.a.
I.23. Seal/Container No		I.24. Type of packaging	
I.25. Commodities certified for:			
Technical use			
I.26. For transit through EU to third	country	I.27. For import or admission into EU	
Third country	ISO code		
I.28. Identification of the commodition	25		
1.20. Identification of the commodition			
Species (Scientific name)	Approval number of establishr Manufacturing plant	ments Batch number	
	31		

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	II.	Health information	II.a. Certificate reference No	II.b.
	II.1.	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 that: the blood products described above consist of blood products that satisfy the health requirements below;		
	II.2.	they consist exclusively of blood produ	cts not intended for human or animal consumptio	n;
Part II: Certification	II.3.	establishment of collection, exclusively (²) either [- blood of slaughte	I in a plant supervised by the competent auth with the following animal by-products: red animals, which is fit for human consumption tion, but is not intended for human consumption	in accordance
Part II: C		(²)and/or [- blood of slaughter accordance with communicable to slaughtered in a s	red animals, which is rejected as unfit for human Union legislation, but which did not show any signification or animals, derived from carcases to slaughterhouse and were considered fit for human mortem inspection in accordance with Union legit	gns of diseases hat have been an consumption
		communicable to slaughtered in a	ered animals, which did not show any sigr humans or animals, obtained from animals t slaughterhouse after having been considered owing an ante-mortem inspection in accordar	hat have been I fit for human
		(²)and/or [- blood and blood human consumpt	products derived from the production of production of production on;]	ets intended for
			products originating from live animals that did no nunicable through that product to humans or anir	
		treatment as define	ts derived from animals which have been sub- ned in Article 1(2)(d) of Council Directive 96/22/ rective 96/23/EC ^(2b) ;]	
		contaminants liste	ts containing residues of other substances and in Group B(3) of Annex I to Directive 96/23/EC, ted level laid down in Union legislation or, in the a tion;]	if such residues
	II.4.	accordance with Union legislation, in	nufactured from, was collected in slaughterhous slaughterhouses approved and supervised by r from live animals in facilities approved and su ollection;	the competent
	(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;		
	(²)either	ISO country code in the thereof) where no case least the preceding 12 m	s thereof	rritories or parts or a period of at
		ISO country code in the one case of foot-and-mopreceding 12 months a disease are being official	pries or parts thereof	of at least the foot-and-mouth
	(²)[II.5.1.	(²)either [no case of vesicular sto animals) has been recor	e and Tayassuidae, in third countries or regions matitis and bluetongue(²) (including the presence ded for a period of at least the preceding 12 mont n carried out against those diseases for a period	of seropositive hs and in which

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II.	Health information		II.a. Certificate reference No	II.b.	
	(2) or [voois	aular atomatitis and b	uotongua(2) garangaitiva animala ara procent ⁽⁴⁾	0.11	
(²)[II.5.2.	(²)or [vesicular stomatitis and bluetongue(²) seropositive animals are present(⁴);]] 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: (²)either [no case of vesicular stomatitis (including the presence of seropositive animals) has been recorded for a period of at least the preceding 12 months and in which vaccination has				
	.0.		t this disease for a period of at least the preced positive animals are present(4);]]]	ing 12 months;]]	
(²)[II.6.					
	which has been free from Newcastle disease and highly pathogenic avian influenza as defined in the Terrestrial Animal Health Code of the OIE, which for a period of at least the preceding 12 months has not carried out vaccination against avian influenza, where the animals from which the products are derived, have not been vaccinated against Newcastle disease with vaccines prepared from a Newcastle disease master strain showing a higher pathogenicity than lentogenic virus strains;]				
II.7.	the products were:				
	(²)either [pack	ed in new or sterilise	d bags or bottles,]		
			niners or other means of transport that were the fectant approved by the competent authority by the co		
the outer packaging or containers bear labels indicating 'NOT FOR HUMAN CONSUMPTION';				I OR ANIMAL	
II.8.	the products were s	tored in enclosed sto	rage;		
II.9.	all precautions wer transport;	all precautions were taken to avoid contamination of the products with pathogenic agents during			
(²)[II.10.	the untreated blood	products described a	bove		
	• •	rived from other rumi	nants than bovine, ovine or caprine animals.]]		
	(²)or [is de from:		ne or caprine animals and does not contain ar	nd is not derived	
	(²) either	continuously reared	caprine materials other than those derived from and slaughtered in a country or region classion in accordance with Decision 2007/453/EC.]]		
	(²)or		sk material as defined in point 1 of Annex V to I1 of the European Parliament and of the Coul		
		caprine and reared and negligible	ly separated meat obtained from bones of the mals, except from those animals that were bounded in a country or region classification accordance with Committed C(²), in which there has been no indigenous E	rn, continuously ed as posing a ssion Decision	
		caprine an the centra instrument into the c continuous	product or derived product obtained from be mals which have been killed, after stunning, nervous tissue by means of an elonga introduced into the cranial cavity, or by means ranial cavity, except for those animals the y reared and slaughtered in a country or regionalished by the country or segligible BSE risk in accordance with Decision	by laceration of ted_rod-shaped s of gas injected hat_were_born, ion_classified_as	
Notes					
Part I:					
_	filled in only if it is a certific filled in if the certificate is f Box reference I.11 and I.12 been issued by the compe	cate for a commodity or a commodity that is 2: Approval number: the tent authority.	onsignment in the European Union: this box is that is to be transited through the European Union. It to be imported into the European Union. The registration number of the establishment or	Jnion; it may be plant, which has	
_	 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.

II.	Health information	II.a.	Certificate reference No	II.b.	
	 Box reference I.15: Registration number (railwaname (ship) is to be provided. In the case of unust inform the border inspection post of the post of t	nloadir int of e em (H ntainer r than	g and reloading in the European Union, to the European Union. S) code under the following headings: 05 number and the seal number (if applicated feeding of farmed animals, other than furth it is a transit or an import certificate.	he consignor .11; 30.02 or ble) must be animals, and	
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		• , ,	,	
(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.				
(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)	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 difference. Note for the person responsible for the consignment purposes and must accompany the consignment into the European Union. 	ent in	the European Union: this certificate is only		
Offic	ial veterinarian/Official inspector				
	Name (in capital letters): Qualifi	cation	and title:		
	Date: Signa	ture:			

Stamp:

