### CHAPTER 4(D)

### Health certificate

For treated 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  $\binom{2}{2}$  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		
t	Postal code Tel.	Postal code Tel.		
ignmen				
ed cons	I.7. Country of ISO code I.8. Region of Code origin origin	I.9. Country of ISO I.10. Region of Code destination code destination		
atch	I.11. Place of origin	I.12. Place of destination		
Part I : Details of dispatched consignment	Name Approval number Address	Custom warehouse Name Approval number Address		
: Detail		Postal code		
Part I				
	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		
	Ambient C Chilled Frozen			
l	Page of			

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123. Seal/Container No       124. Type of packaging         125. Commodities certified for:		uenveu pro	I.2. Certificate reference No	I.2.a.
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 commodities         Species       Approval number of establishments         Batch number	I.23. Seal/Container No		I.24. Type of packaging	
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 commodities         Species       Approval number of establishments         Batch number	I.25. Commodities certified for:			
Third country     ISO code       I.28. Identification of the commodities       Species     Approval number of establishments     Batch number	Technical use			
I.28. Identification of the commodities         Species       Approval number of establishments         Batch number	I.26. For transit through EU to third country		I.27. For import or admission into EU	
Species Approval number of establishments Batch number	Third country ISO	code		
Species (Scientific name)     Batch number	I.28. Identification of the commodities			
Species Approval number of establishments Batch number (scientific name)				
	Species // (Scientific name)	Approval number of esta Manufacturing pla	ablishments Batch number nt	

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	П.	Health inform	ation	II.a. Certificate reference No	II.b.	
ا. I, the undersigned official veterinarian, declare that I have read and understood Regulation (EC) No 10 European Parliament and of the Council( <sup>1a</sup> ), and in particular Article 8(c) and Article 8(d) and Article Commission Regulation (EU) No 142/2011( <sup>1b</sup> ), and in particular Chapter II of Annex XIV thereto, and certify t					e 10 thereof, and	
	II.1.	the blood produc	ts described above consis	t of blood products that satisfy the requirements below;		
<ul> <li>II.2. they consist exclusively of blood products not intended for human or animal consumption;</li> <li>II.3. they have been prepared and stored in a plant supervised by the competent authority, exclusively with by-products;</li> </ul>			e following animal			
ation		<del>(<sup>2</sup>)either [-</del>		animals, which is fit for human consumption in accordance with or human consumption for commercial reasons;]	Union legislation,	
Part II: Certification		( <sup>2</sup> )and/or [-	Union legislation, bu derived from carcas	I animals, which is rejected as unfit for human consumption in t which did not show any signs of diseases communicable to h es that have been slaughtered in a slaughterhouse and were following an ante-mortem inspection in accordance with Union	umans or animals, considered fit for	
Part II		<del>(<sup>2</sup>)and/or [-</del>	animals, obtained fr	⊢animals, which did not show any signs of diseases communic om animals that have been slaughtered in a slaughterhouse nan consumption following an ante-mortem inspection in acco	after having been	
		( <sup>2</sup> )and/or [-		ducts originating from live animals that did not show clinical sig gh these products to humans or animals;]	ins of any disease	
		<del>(<sup>2</sup>)and/or [-</del>	blood and blood proc	ducts derived from the production of products intended for hum	an consumption;]	
		(²)and/or [-		which have been derived from animals which have been so d in Article 1(2)(d) of Council Directive 96/22/EC <sup>(2a)</sup> or Articl <sup>(b)</sup> ;]		
		( <sup>2</sup> )and/or [-	Group B(3) of Annex	containing residues of other substances and environmental con k I to Directive 96/23/EC, if such residues exceed the permitte or, in the absence thereof, in national legislation;]		
	II.4.	the blood that these products were manufactured from was been 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.				
	( <sup>2</sup> )[II.5.	[II.5. In the case of blood products derived from Artiodactyla, Perissodactyla and Proboscidea including their crossbreed than Suidae and Tayassuidae, the products have undergone one of the following treatments, guaranteeing the abs pathogens of foot-and-mouth disease, vesicular stomatitis, rinderpest, peste des petits ruminants, Rift Valley fe bluetongue:				
		( <del>²)either</del>	•	perature of 65 °C for at least three hours, followed by an effecti	veness check;]	
		<del>(<sup>2</sup>)and/or</del>		gamma rays, followed by an effectiveness check;]		
		<del>(<sup>2</sup>)and/or</del>		r two hours, followed by an effectiveness check;]		
	(2)[1] C	( <sup>2</sup> )and/or	•	st 80 °C throughout their substance, followed by an effectivene		
	<del>(<sup>-</sup>)[11.0.</del>	( <sup>2</sup> )[II.6. In the case of blood products derived from Suidae, Tayassuidae, poultry and other avian species, the products undergone one of the following treatments guaranteeing the absence of pathogens of the following diseases: foot-and-r disease, vesicular stomatitis, swine vesicular disease, classical swine fever, African swine fever, Newcastle disease highly pathogenic avian influenza, as appropriate to the species:		es: foot-and-mouth astle disease and		
		( <sup>2</sup> )either	•	perature of 65 °C for at least three hours, followed by an effecti	veness check;]	
		<del>(<sup>2</sup>)and/or (<sup>2</sup>)and/or</del>		gamma rays, followed by an effectiveness check;] st 80 °C for Suidae/Tayassuidae( <sup>2</sup> ) and at least 70°C for poult	ny and other avian	
		t <del>janu/or</del>		e substance of the product, followed by an effectiveness check		
	( <sup>2</sup> )[II.7. In the case of blood products derived from species other than those listed in point II.5 or II.6, the products have up of the following treatment (please specify):					
	II.8.	The products we	re:			
			acked in new or sterilised l			
( <sup>2</sup> )or [transported in bulk in containers or other means of transport that were thoroughly cleaned a with a disinfectant approved by the competent authority before use;] and						
				DN';		
	II.9.					
	II.10.       all precautions were taken to avoid the contamination of the products with pathogenic agents after treatment;         ( <sup>2</sup> )[II.11.       The treated blood products described above         ( <sup>2</sup> )either       [is derived from other ruminants than bovine, ovine or caprine animals.]]		ent;			
					<b></b>	
			ither [bovine, ovin reared and s	or caprine animals and does not contain and is not derived from the and caprine materials other than those derived from animals slaughtered in a country or region classified as posing a neg with Decision 2007/453/EC.]]	born, continuously	
		(²)0/	r [(a) spe	ecified risk material as defined in point 1 of Annex V to Re 9/2001 of the European Parliament and of the Council( <sup>3</sup> );	egulation (EC) No	

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П.	Health information	II.a. Certificate reference No	II.b.			
	(b) 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 in accordance with Commission Decision 2007/453/EC( <sup>4</sup> ), in which there has been no indigenous BSE case.					
	wh me by bo	nimal by-product or derived product obtained from bovine, ovine or caprine animals hich have been killed, after stunning, by laceration of the central nervous tissue by leans of an elongated rod-shaped instrument introduced into the cranial cavity, or y means of gas injected into the cranial cavity, except for those animals that were orn, continuously reared and slaughtered in a country or region classified as posing negligible BSE risk in accordance with Decision 2007/453/EC.]]]				
Notes						
Part I: -	Box reference I.6: Person responsible for the cor certificate for a commodity to be transited through	nsignment in the European Union: this box is required to be filled n the European Union; it may be filled in if the certificate is for a c				
-	imported into the European Union. Box reference I.11 and I.12: Approval number: the competent authority.	e registration number of the establishment or plant, which has bee	en issued by the			
-	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 BIP of entry into the European Union.					
-	Box I.19: use the appropriate Harmonized System (HS) code under the following headings: 05.11, 30.02, 35.02 or 35.04.					
-		ner number and the seal number (if applicable) must be included. r than feeding of farmed animals, other than fur animals, and th				
	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 in case of Species: select from the following: Aves, Ruminantia, Suidae, Mammalia other than Ruminantia or Suidae, Pesca, Reptilian.					
Part II	:					
(1a) (1b)	OJ L 300, 14.11.2009, p. 1.					
(10)	OJ L 54, 26.2.2011, p. 1.					
(2a)	Delete as appropriate. OJ L 125, 23.5.1996, p. 3.					
(2b)	OJ L 125, 23.5.1996, p. 10.					
(3)	OJ L 147, 31.5.2001, p. 1.					
(4)	OJ L 172, 30.6.2007, p. 84.					
-	<ul> <li>The signature and the stamp must be in a different colour to that of the printing.</li> <li>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 European Union.</li> </ul>					
Offic	Official veterinarian/Official inspector					
	Name (in capital letters):	Qualification and title:				
	Date:	Signature:				
		Stamp:				
		examp.				