CHAPTER 4(B)

Health certificate

For blood products not intended for human consumption that could be used as feed material, intended for dispatch to or for transit through² the European Union

COU	NTRY: UNITED STATES	Veterinary certificate to EU		
	I.1. Consignor Name Address	I.2. Certificate reference No I.2.a.		
	, addied	I.3. Central competent authority APHIS-VS		
	Tel.	I.4. Local competent authority		
	I.5. Consignee Name Address Postal code	I.6. Person responsible for the load in EU Name Address Postal code		
Part I : Details of dispatched consignment	Tel.	Tel.		
	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		
ls of disp	Name Approval number Address	Custom warehouse Name Approval number Address		
I : Detai		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		
	Ambient ☐ Chilled ☐ Frozen ☐			

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COUNTRY: UNITED STATES

Blood products not intended for human consumption that could be used as feed material

	I.2. Certificate reference No	1.2.a.
I.23. Seal/Container No	I.24. Type of packaging	
I.25. Commodities certified for:	-	
Animal feedingstuff \Box Manufacture of petfood \Box Technical use \Box		
I.26. For transit through EU to third country	I.27. For import or admission into EU	
Third country ISO code		
1.28. Identification of the commodities		
Species Nature of commodity Ap	oproval number of establishments Batch number	or
(Scientific name)	Manufacturing plant	51

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			could be used as feed material		
II.	Health inform	nation	II.a. Certificate reference No	II.b.	
	No 1069/20	09 of the European Parlia	declare that I have read and understood ment and of the Council(1a) and Comm he blood products described above:		
II.1.	consist of bl	ood products that satisfy th	ne health requirements below;		
II.2.	consist excl	usively of blood products n	ot intended for human consumption;		
II.3.	have been prepared and stored in a plant, approved and supervised by the competent authorit in accordance with Article 24 of Regulation (EC) No 1069/2009;				
II.4.	have been p		ne following animal by-products:		
II.4.	(²)either		nimals, which is fit for human consumpti but which is not intended for human		
	(2)and/or [blood of slaughtered animals, which has been rejected as unfit for consumption in accordance with Union legislation, but which did not signs of diseases communicable to humans or animals, which has been from carcases that have been slaughtered in a slaughterhouse and we considered fit for human consumption following an ante-mortem inspace accordance with Union legislation;]				
II.5.	in order to ir	nactivate pathogenic agent	-		
	(²)either	to processing in accor	dance with processing methodo Regulation (EU) No 142/2011;]	.(³) as set out i	
	(²)or		meters which ensure that the product of the set out in Chapter I of Annex X to Re		
	(²)or	porcine origin intended for temperature of at least 8	educts, including spray dried blood and for the feeding of porcine animals, to a h 0°C throughout the substance and the dr n more than 8% w/w moisture with a wat	eat treatment at a	
II.6.	the end prod	duct was:			
	(²)either	[packed in new or sterilis	sed bags;]		
	(²)0r		ontainers or other means of transport that with a disinfectant approved by the co		
and which bear labels indicating 'NOT FOR HUMAN CONSUMPTION'; II.7. the end product was stored in enclosed storage; II.8. the product has undergone all precautions to avoid contamination with pathogenic ager treatment;		FOR HUMAN CONSUMPTION';			
		genic agents afte			
	(²)and	porcine origin intended	oducts, including spray dried blood and for the feeding of porcine animals, has be	een stored in dr	
	b b		nder room temperature for a period of at	=	
II.9. have been examined prior to dispatch under the responsibility of the competent at a random sample during or on removal from storage which was found to comply standards(4):					
	Salmonella:	absence in 25g	y: n = 5, c = 0, m = 0, M = 0,		
	Enterobacteriaceae: n = 5, c = 2, m = 10, M = 300 in 1 gram;				
(²)[II.10.	•	oducts described above			
	(²)either [is derived from other ruminants than bovine, ovine or caprine animals.]]				
(2)or [is derived from bovine, ovine or caprine animals and does not contain and derived from:			ontain and is no		
	(2)	either [bovine, ovine	and caprine materials other than the	se derived from	

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II.	Health information		II.a. Certificate reference No	II.b.	
		classified as po 2007/453/EC.]]	osing a negligible BSE risk in acco	ordance with Decision	
	(²)or		d risk material as defined in poi ion (EC) No 999/2001 of the Europo ncil(⁵);		
		ovine o born, co classifie Commis	rically separated meat obtained for caprine animals, except from the continuously reared and slaughtered as posing a negligible BSE rist csion Decision 2007/453/EC(6), in venous BSE case,	se animals that were in a country or region k in accordance with	
		or capri laceration rod-sha means animals country	by-product or derived product obtain ine animals which have been killed on of the central nervous tissue by n ped instrument introduced into the of gas injected into the cranial ca that were born, continuously reared or region classified as posing a runce with Decision 2007/453/EC.]]]	ed, after stunning, by neans of an elongated recranial cavity, or by vity, except for those d and slaughtered in a	
II.11.	the blood products	s described above:			
			milk products of ovine or caprine a ned animals, other than fur animals.		
		-			
	(a)		n ovine and caprine animals wh ce birth in a country where the fol		
		(i) classica	ıl scrapie is compulsorily notifiable;		
			eness, surveillance and monitoring ll scrapie;	system is in place for	
			estrictions apply to holdings of ovince e of a suspicion of TSE or the cor :		
		(iv) ovine a	nd caprine animals affected with nd destroyed;	classical scrapie are	
		o r grea the Wor has bee	ling to ovine and caprine animals of ves, as defined in the Terrestrial A ld Organisation for Animal Health (C in banned and effectively enforced in I of at least the preceding seven yea	nimal Health Code of NE), of ruminant origin the whole country for	
	(b)		oldings where no official restrictions		
	(c)	diagnosed during	oldings where no case of classic g the period of at least the prece firmation of a case of classical scra	ding seven years or,	
		destroy ARR/AF allele ar	e and caprine animals on the holding ed or slaughtered, except for br RR genotype, breeding ewes carry nd no VRQ allele and other ovine an R allele;]	eeding rams of the ing at least one ARR	
		killed ar period o	nals in which classical scrapie was nd destroyed, and the holding has of at least two years since the date scical scrapie case to intensified TSE	been subjected for a of confirmation of the	

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		could be used as feed material
II.	Health information	II.a. Certificate reference No II.b.
		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 999/2001, of all of the following animals which are over the age of 18 months, except evine animals of the ARR/ARR genetype: — 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 eradication campaign.]]
II.12.	the blood products described above contain or are derived from animal-by products of non-ruminant origin, and are, according to the statement of the Consignor referred to in Box I.1, (2)either [not intended for the production of feed for farmed animals, other than fur animals.] (2)(7)or [intended for the production of feed for non-ruminant farmed animals, other than fur animals, and the Consignor has undertaken to ensure that the border inspection post of entry will be provided with the results of the analyses carried out in accordance with the methods set out in Annex VI to Commission Regulation (EC) No 152/2009(8).]	
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.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); information is to be provided in the case of unloading and reloading in the European Union.
- Box reference I.19: use the appropriate HS code: 05.11.91, 05.11.99, 35.02 or 35.04.
- Box reference I.23: for bulk containers, the container number and the seal number (if applicable) should 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, Reptilia.

Part II:

- (^{1a}) OJ L 300, 14.11.2009, p. 1.
- (1b) OJ L 54, 26.2.2011, p. 1.
- (2) Delete as appropriate.
- (3) Insert method 1 to 5 or method 7 as applicable.
- (4) Where:
 - n = number of samples to be tested;

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II. **Health information** II.a. Certificate reference No II.b. threshold value for the number of bacteria; the result is considered satisfactory if the number m = of bacteria in all samples does not exceed m: maximum value for the number of bacteria; the result is considered unsatisfactory if the number M = of bacteria in one or more samples is M or more; and number of samples the bacterial count of which may be between m and M, the sample still being considered acceptable if the bacterial count of the other samples is m or less. OJ L 147, 31.5.2001, p. 1. OJ L 172, 30.6.2007, p. 84. The person responsible for the load referred to in Box I.6 must ensure that, if the blood products described in this health certificate are intended to be used for the production of feed for non-ruminant farmed animals, other than fur animals, the consignment must be analysed, in accordance with the methods set out in Annex VI to Regulation (EC) No 152/2009, in order to verify the absence of unauthorised constituents of animal origin. The information on the result of such analysis must be attached to this health certificate when presenting the consignment at a border inspection post of the European Union. OJ L 54, 26.2.2009, p. 1. 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. Official veterinarian/Official inspector Name (in capital letters): Qualification and title:

Signature:

Stamp:

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Date: