## 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 through<sup>2</sup> 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 I.4. Local competent authority
	Tel.	
	I.5. Consignee Name Address	I.6. Person responsible for the load in EU Name Address
	Postal code Tel.	Postal code Tel.
gnment		
consi	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
tched	I.11. Place of origin	I.12. Place of destination
ls of dispa	Name Approval number Address	Custom warehouse Name Approval number Address
Part I : Details of dispatched consignment		Postal code
	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 Chilled	
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OUNTRY: UNITED STAT	TES .	Blood products not intended for hum could be used as feed material	an consumption that					
		I.2. Certificate reference No	I.2.a.					
.23. Seal/Container No	)	I.24. Type of packaging						
.25. Commodities certi	fied for:							
Animal feedingstuff 🗆	Manufacture of petfood   Technica	I use 🗆						
.26. For transit through	EU to third country	I.27. For import or admission into EU	I.27. For import or admission into EU					
Third country	ISO code							
.28. Identification of the	e commodities							
Species (Scientific name)	Nature of commodity	Approval number of establishments Manufacturing plant	Batch number					
Page o								

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

	П.	Health inform	nation	II.a.	Certificate reference No	II.b.					
	II.1.	No 1069/20 (EU) No 142	09 of the European Parlia	ment a	re that I have read and understoo and of the Council( <sup>1a</sup> ) and Comm od products described above: Ith requirements below;						
	II.2.	consist exclusively of blood products not intended for human consumption;									
ition	II.3.		prepared and stored in a pl ce with Article 24 of Regula		pproved and supervised by the co EC) No 1069/2009;	mpetent authority					
fice	11.4.	have been prepared exclusively with the following animal by-products:									
Part II: Certification		<ul> <li>(<sup>2</sup>)either [blood of slaughtered animals, which is fit for human consumption in a with Union legislation, but which is not intended for human consu commercial reasons;]</li> <li>(<sup>2</sup>)ent/(ent)</li> </ul>									
Par		(²)and/or	r [blood of slaughtered animals, which has been 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, which has been derived from carcases that have been slaughtered in a slaughterhouse and which were considered fit for human consumption following an ante-mortem inspection in accordance with Union legislation;]								
	II.5.	in order to ir	nactivate pathogenic agent	s, have	e been submitted						
		( <sup>2</sup> )either	( <sup>2</sup> ) <i>either</i> [to processing in accordance with processing method( <sup>3</sup> ) as set out in Chapter III of Annex IV to Regulation (EU) No 142/2011;]								
		(²)or	( <sup>2</sup> )or [to a method and parameters which ensure that the product complies with the microbiological standards set out in Chapter I of Annex X to Regulation (EU) No 142/2011;]								
		( <sup>2</sup> )or [in the case of blood products, including spray dried blood and blood plasma, of porcine origin intended for the feeding of porcine animals, to a heat treatment at a temperature of at least 80°C throughout the substance and the dry blood and blood plasma does not contain more than 8% w/w moisture with a water activity (Aw) of less than 0,60.]									
	II.6.	the end product was:									
		( <sup>2</sup> )either	[packed in new or sterilis	sed ba	gs;]						
		( <sup>2</sup> ) <i>or</i> [transported in bulk in containers or other means of transport that were thoroughly cleaned and disinfected with a disinfectant approved by the competent authority before use,]									
		and which bear labels indicating 'NOT FOR HUMAN CONSUMPTION';									
	II.7.	the end proc	duct was stored in enclose	d stora	ge;						
	II.8.	the product treatment;	has undergone all precau	tions to	o avoid contamination with patho	genic agents after					
		(²)and	porcine origin intended	for the	, including spray dried blood and feeding of porcine animals, has l oom temperature for a period of a	been stored in dry					
	II.9.	have been examined prior to dispatch under the responsibility of the competent authority by taking a random sample during or on removal from storage which was found to comply with the following standards( <sup>4</sup> ):									
		Salmonella:	absence in 25g	g: n = 5	, c = 0, m = 0, M = 0,						
		Enterobacteriaceae: n = 5, c = 2, m = 10, M = 300 in 1 gram;									
	( <sup>2</sup> )[II.10.	the blood pr	oducts described above								
	()L	( <sup>2</sup> ) <i>either</i> [is derived from other ruminants than bovine, ovine or caprine animals.]]									
		<sup>(2)</sup> or [is			caprine animals and does not c						
		(2)			caprine materials other than the ously reared and slaughtered in a						

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

II.	Health inform	mation			II.a.	Certi	ficate r	eferen	ce No			II.b.	
				ied as po 153/EC.]]		a neg	ligible	BSE	risk ir	accol	rdance	e with	Decisio
	(2	)or	[(a)	specified Regulation the Cour	d risł ion (E incil( <sup>5</sup> )	EC) No );	999/2	:001 c	f the E	Europe	an Pa	rliamer	nt and o
			(b)	mechani ovine or born, con classified Commiss no indige	r capi ontinu ed as ssion	rine ar ously i posin Decisi	nimals eared g a no on 200	, exce and s egligit	pt fro slaugh ble BS	m thos tered i SE risk	se ani in a co c in ac	mals th ountry o ccordai	nat were or region nce with
			(c)	animal b or caprir laceratio rod-shap means c animals country accordar	ine an on of t ped in of ga that v or re	nimals he cer nstrum s injec were b gion c	which ntral ne ent in sted in orn, co lassifi	n have ervous troduc to the ontinu ed as	e bee s tissu ced in e cran ously posir	n kille e by m to the ial cav reared ng a n	d, afte leans o crania /ity, ex and s	er stun of an el al cavit cept f laughte	ning, by longated ty, or by or those ered in a
II.11.	the blood pr	roducts d	escribed a	above:									
	( <sup>2</sup> )either			milk or m d for farm								origin	or is no
	( <sup>2</sup> ) <i>or</i> [contain milk feed for farme										in and	is inte	nded fo
		(a)		ived from ously sinc									
			(i)	classical	l scra	pie is	compu	Isorily	/ notifi	able;			
			(ii)	an aware classical			eillanc	e and	l moni	toring	syster	n is in l	place fo
			(iii)	official re the case scrapie;	e of a								
			(iv)	ovine ar killed and				als aff	ected	with o	classio	al scr	apie are
			(v)	the feedi or greav the Work has beer a period	ves, a ld Org n ban	as defi ganisa ned ar	ned in tion foi nd effe	the 1 Anim ctivel	errest al Hea / enfoi	rial Ar alth (O ced in	nimal I IE), of the w	Health rumina	Code c ant origi
		(b)	•	e from hol on of TSE;	•	s wher	e no o	fficial	restrio	tions a	are im	posed	due to a
		(c)	diagnos	e from ho ed during g the conf	g the	perio	d of a	t leas	t the	preced	ding s		
				all ovine destroye ARR/AR allele an one ARF	e and ed or RR ge nd no	caprin slaug notype VRQ a	e anin ghtere e, bree	nals o d, ex eding	n the h cept ewes	olding for bro carryii	i have eeding ng at	rams least o	of the
			(²)or	[all anim killed an period of last class	nals ir nd de of at le	n whic stroye east tw	d, and /o yea	l the I rs sin	nolding ce the	g has l date o	been : of con	subject firmatio	ted for a

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Blood products not intended for human consumption that 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 o Chapter C of Annex X to Regulation (EC) No 999/2001, of all o the following animals which are over the age of 18 months, excep ovine animals of the ARR/ARR genotype:
	<ul> <li>animals which have been slaughtered for human consumption; and</li> </ul>
	<ul> <li>animals which have died or been killed on the holding bu which were not killed in the framework of a disease eradication campaign.]]</li> </ul>
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,
	( <sup>2</sup> )either [not intended for the production of feed for farmed animals, other than fur animals.
	( <sup>2</sup> )( <sup>7</sup> )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( <sup>8</sup> ).]
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 transic commodity. Products in transit may only be stored in free zones, free warehouses and custon warehouses.
	Box reference I.15: Registration number (railway wagons or container and lorries), flight numbe (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 fu 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 othe than Ruminantia or Suidae , Pesca, Reptilia.
Part II:	
( <sup>1a</sup> ) O	J L 300, 14.11.2009, p. 1.
( <sup>1b</sup> ) O	J L 54, 26.2.2011, p. 1.
	elete as appropriate.
( <sup>2</sup> ) D	
	sert method 1 to 5 or method 7 as applicable.
( <sup>3</sup> ) In	sert method 1 to 5 or method 7 as applicable. /here:

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

		could be used as leed material	
II.	Health information	II.a. Certificate reference No	II.b.
		nber of bacteria; the result is considered satis	sfactory if the numb
	of bacteria in all samples d M = maximum value for the num	pes not exceed m; ber of bacteria; the result is considered unsati	sfactory if the numb
	of bacteria in one or more		Side for y in the numb
		cterial count of which may be between m ar le if the bacterial count of the other samples is	
(5)	OJ L 147, 31.5.2001, p. 1.		
( <sup>6</sup> )	OJ L 172, 30.6.2007, p. 84.		
(7)	described in this health certificate farmed animals, other than fur an methods set out in Annex VI to unauthorised constituents of anim	ad referred to in Box I.6 must ensure that, are intended to be used for the production of mals, the consignment must be analysed, in Regulation (EC) No 152/2009, in order to v al origin. The information on the result of su then presenting the consignment at a border	feed for non-rumin accordance with t verify the absence uch analysis must
(8)	OJ L 54, 26.2.2009, p. 1.		
	- Note for the person responsible	ust be in a different colour to that of the printi for the consignment in the European Union: at accompany the consignment until it reaches e European Union.	this certificate is o
Offic	cial veterinarian/Official inspector		
	Name (in capital letters):	Qualification and title:	
	Date:	Signature: Stamp:	
		Stamp.	

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