# CHAPTER 3(B)

# **Health certificate**

For processed petfood other than canned petfood, intended for dispatch to or for transit through  $\binom{2}{2}$  the European Union

# **COUNTRY: UNITED STATES**

# Veterinary certificate to EU

	I.1. Consignor	I.2. Certificate reference No I.2.a.			
	Name Address				
	Address	I.3. Central competent authority			
	Tal	APHIS-VS  I.4. Local competent authority			
	Tel.				
	I.5. Consignee Name	I.6. Person responsible for the load in EU Name			
	Address	Address			
	Postal code	Postal code			
	Tel.	Tel.			
ent					
muß					
Consi	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			
ıtched	I.11. Place of origin	I.12. Place of destination			
Part I : Details of dispatched consignment	Name Approval number Address	Custom warehouse Name Address			
etails		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  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 ☐				

### **COUNTRY: UNITED STATES**

# Processed petfood other than canned petfood

		1.2.	Certificate reference No		1.2.a.
I.23. Seal/Container No		1.24.	Type of packaging		
I.25. Commodities certified for:		<u> </u>			
Petfood □ Technical use	e 🗆				
I.26. For transit through EU to thir	d country	1.27.	For import or admission into EU		
Third country	ISO code				
I.28. Identification of the commod	ities				
Species (Scientific name)	Approval number of establishme Manufacturing plant	ents	Net weight	Batch num	ıber

II.

**COUNTRY: UNITED STATES** 

#### **Health information** II.a. Certificate reference No II.b. 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 Articles 8 and 10 thereof, and Commission Regulation (EU) No 142/2011(1b), and in particular Chapter II of Annex XIII and Chapter II of Annex XIV thereto, and certify that the petfood described above: has been prepared and stored in a plant approved and supervised by the competent authority in accordance with Article 24 of II.1. Regulation (EC) No 1069/2009; 11.2. has been prepared exclusively with the following animal by-products: carcases and parts of animals slaughtered or, in the case of game, bodies or parts of animals killed, and which are fit for human consumption in accordance with Union legislation, but are not intended for human consumption for commercial reasons;] (2)and/or carcases and the following parts originating either from animals that have been slaughtered in a ſslaughterhouse and were considered fit for slaughter for human consumption following an ante-mortem inspection or bodies and the following parts of animals from game killed for human consumption in accordance with Union legislation: carcases or bodies and parts of animals which are rejected as unfit for human consumption in accordance with Union legislation, but which did not show any signs of disease communicable to humans or animals; (ii) heads of poultry: (iii) hides and skins, including trimmings and splitting thereof, horns and feet, including the phalanges and the carpus and metacarpus bones, tarsus and metatarsus bones; (iv) pig bristles; (v) feathers;] (2)and/or animal by-products from poultry and lagomorphs slaughtered on the farm as referred to in Article 1(3)(d) of Regulation (EC) No 853/2004 of the European Parliament and of the Council(2a), which did not show any signs of disease communicable to humans or animals] blood of animals which did not show any signs of disease communicable through blood to humans or (2)and/or animals, obtained from animals that have been slaughtered in a slaughterhouse after having been considered fit for slaughter for human consumption following an ante-mortem inspection in accordance with Union legislation: (2)and/or animal by products arising from the production of products intended for human consumption, including degreased bone, greaves and centrifuge or separator sludge from milk processing;] products of animal origin, or foodstuffs containing products of animal origin, which are no longer intended (2)and/or for human consumption for commercial reasons or due to problems of manufacturing or packaging defects or other defects from which no risk to public or animal health arise;] (2)and/or petfood and feedingstuffs of animal origin, or feedingstuffs containing animal by products or derived products, which are no longer intended for feeding for commercial reasons or due to problems of manufacturing or packaging defects or other defects from which no risk to public or animal health arise;] blood, placenta, wool, feathers, hair, horns, hoof cuts and raw milk originating from live animals that did (2)and/or not show signs of any disease communicable through that product to humans or animals;] aquatic animals, and parts of such animals, except sea mammals, which did not show any signs of diseases (2)and/or communicable to humans or animals;] animal by-products from aquatic animals originating from plants or establishments manufacturing products (2)and/or for human consumption:1 (2)and/or the following material originating from animals which did not show any signs of disease communicable through that material to humans or animals: shells from shellfish with soft tissue or flesh: the following originating from terrestrial animals: hatchery by-products, egg by-products, including egg shells, day-old chicks killed for commercial reasons;] (2)and/or animal by-products from aquatic or terrestrial invertebrates other than species pathogenic to humans or animals and parts thereof of the zoological orders of Rodentia and Lagomorpha, except Category 1 material (2)and/or as referred to in Article 8(a)(iii), (iv) and (v) of Regulation (EC) No 1069/2009 and Category 2 material as referred to in Article 9(a) to (g) of that Regulation;] (2)and/or material from animals which have been treated with certain substances which are prohibited by Council Directive 96/22/EC(2b), the import of the material being permitted in accordance with Article 35(a)(ii) of Regulation (EC) No 1069/2009;] 11.3. (2)either [was subjected to a heat treatment of at least 90 °C throughout its substance;] [was produced as regards ingredients of animal origin using exclusively products which had been: in the case of animal by-products or derived products from meat or meat products subjected to a heat treatment of at least 90 °C throughout its substance; in the case of milk and milk based products,

# Processed petfood other than canned petfood

II.	Health informatio	n	II.a. Certificate reference No	II.b.				
			hird countries or parts of third countries listed in column B of Annex I to Commission No 605/2010( <sup>a</sup> ) submitted to a pasteurisation treatment sufficient to produce a atase test;					
		of Annex I to Re	d to less than 6 from third countries or parts of third countries listed in column C gulation (EU) No 605/2010, first submitted to a pasteurisation treatment sufficient ative phosphatase test; after countries or parts of third countries listed in column C of Annex I to Regulation IO, submitted to a sterilisation process or a double heat treatment where each ifficient to produce a negative phosphatase test on its own;					
		(iii) if they are from the (EU) No 605/20						
		(EU) No 605/201 12 months or w	nird countries or parts of third countries listed in column C of Ani 0, where there has been an outbreak of foot-and-mouth diseas here vaccination against foot-and-mouth disease has been hths, submitted to	e in the preceding				
		- a sterilisa or	ation process whereby an Fc value equal or greater than 3 is ac	hieved				
		<del>pasteuris</del>	heat treatment with a heating effect at least equal to the ation process of at least 72 °C for at least 15 seconds and suffireaction to a phosphatase test, followed by					
		- a second heat trea	heat treatment with a heating effect at least equal to that ach atment, and which would be sufficient to produce a negat tase test, followed, in the case of dried milk, or dried milk bacocess	ive reaction to a				
		<del>Or</del>	cation process such that the pH has been maintained at less	than 6 for at least				
	<del>(c)</del>	one hour;						
	<del>(d)</del>	in the case of hydrolysed minimise contamination of derived from ruminant his production, using only m	d protein produced using a production process involving approper fraw Category 3 material, and, in the case of hydrolysed protectes and skins produced in a processing plant dedicated only total training with a molecular weight below 10000 Dalton and a property 3 material by brining, liming and intensive washing follower	in entirely or partly hydrolysed protein cess involving the				
	(i) exposure of the n		material to a pH of more than 11 for more than three hours a and subsequently by heat treatment at more than 140 °C for 3	t a temperature of				
	treatment at 14 (e) in the case of egg prodi III of Annex IV to Regu		exposure of the material to a pH of 1 to 2, followed by a pH of more than 11, followed by heat treatment at 140 °C for 30 minutes at 3 bar;					
			sts submitted to any of the processing methods 1 to 5 or 7, as refe ation (EU) No 142/2011; or treated in accordance with Chapte EC) No 853/2004;					
	to a treatment involving v		ubmitted to a process ensuring that unprocessed Category 3 ma vashing, pH adjustment using acid or alkali followed by one or ma f preservatives other than those permitted by Union legislation I	ore rinses, filtration				
	<del>(g)</del>		lucts, produced using any of the processing methods 1 to 5 or pegulation (EU) No 142/2011;	7, as referred to in				
	<del>(h)</del>	7 and, in the case of por	n processed animal protein submitted to any of the processing cine blood, submitted to any of the processing methods 1 to 5 a heat treatment throughout its substance at a minimum temper	or 7 provided that				
	<del>(i)</del>		malian processed protein with the exclusion of fishmeal subm · 5 or 7 as referred to in Chapter III of Annex IV to Regulation (E					
	<del>(i)</del>	Annex IV to Regulation (	submitted to any of the processing methods 1 to 7 as referred (EU) No 142/2011 or to a method and parameters which ensure biological standards for derived products set out in Chapter 2011;	re that the product				
	<del>(k)</del>	method 6 in the case of or produced in accordar rendered fats from rumin	at, including fish oils, submitted to any of the processing metho fish oil) as referred to in Chapter III of Annex IV to Regulation are with Chapter II of Section XII of Annex III to Regulation (lant animals must be purified in such a way that the maximum lev does not excess 0,15 % in weight;	(EU) No 142/2011 EC) No 853/2004;				
	<del>(I)</del>	in the case of dicalcium	phosphate produced by a process that					

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II.	Health info	rmation		II.a.	Certificate reference No	II.b.	
		tre		e hydro	ry 3 bone-material is finely crushed and degreased w chloric acid (at a minimum concentration of 4 % and a pl two days:		
		<del>(ii) f</del> o	llowing the pro	cedure i	referred to in (i), applies a treatment of the obtained photo pitate of dicalcium phosphate at pH 4 to 7; and	sphoric liquor with	
					sipitate of dicalcium phosphate with inlet temperature o tween 30 °C and 65 °C ;	f 65 °C to 325 °C	
		(m) in the cas	e of tricalcium	<del>phosph</del>	ate produced by a process that ensures		
			at all Category one chips less		e-material is finely crushed and degreased in counter-flo-mm);	ow with hot water	
		<del>(ii) c</del> c	ontinuous cook	ing with	steam at 145 °C during 30 minutes at 4 bar;		
			eparation of the	<del>protei</del> i	n broth from the hydroxyapatite (tricalcium phosphate)	by centrifugation;	
				e tricalc	ium phosphate after drying in a fluid bed with air at 200 °	<del>,C :</del>	
		(n) in the cas	e of flavouring	nnards,	produced according to a treatment method and paramet to microbiological standards referred to in point II.4.]		
	<del>(²)or</del>	[was subject to a	treatment such	as dryir	ng or fermentation, which has been authorised by the con	npetent authority;]	
	<del>(²)or</del>				ertebrates other than species pathogenic to humans or a		
					authorised by the competent authority and which ensure nd animal health;]	s that the petfood	
II.4.		l by a random sar ant and complies v			samples from each processed batch taken during or af $ards(^4)$ :	ter storage at the	
	Salmonella:	а	bsence in 25g:	n = 5, c	= 0, m = 0, M = 0,		
II.5.	Enterobacteri				= 300 in 1 gramme; with pathogenic agents after treatment;		
II.6. (²)[II.7.	was packed in new packaging, which, if the petfood is not dispatched in ready-to-sell packages on which it is clearly indicate that the content is destined for feeding to pets only, bear labels indicating "NOT FOR HUMAN CONSUMPTION";  7. the petfood described above  (2) either [is derived from other ruminants than bovine, ovine or caprine animals.]  (4) or [is derived from bovine, ovine or caprine animals and does not contain and is not derived from:						
	· ·	( <del>²) either</del>	[bovine, continuous	vine a	nd caprine materials other than those derived from the digital of the same of the derived as placed in a country or region classified as place with Decision 2007/453/EC.]	<del>m animals born,</del>	
	<del>(²)or</del>		<u>[(a)</u>	pecified	I risk material as defined in point 1 of Annex V to Re 1 of the European Parliament and of the Council( <sup>5</sup> );	gulation (EC) No	
			( <del>d)</del> ************************************	nechani inimals, ilaughte iccordai	cally separated meat obtained from bones of bovine, except from those animals that were born, continuous and in a country or region classified as posing a neglinee with Commission Decision 2007/453/EC( <sup>6</sup> ), in whice prous BSE case,	ously reared and igible BSE risk in	
			† † †	nimals issue by avity, o hat wer	by product or derived product obtained from bovine, which have been killed, after stunning, by laceration of the means of an elongated rod-shaped instrument introduct by means of gas injected into the cranial cavity, exceptive born, continuously reared and slaughtered in a clause posing a negligible BSE risk in accordance 3/EC.]]]	ne central nervous ed into the cranial for those animals country or region	
Notes							
Part I:		noo I 6: Doroon ro	ananaibla far t	20 00no	ignment in the European Union; this hav is required to h	o filled in only if it	
-	is a certific		ty to be transite		ignment in the European Union: this box is required to b gh the European Union; it may be filled in if the certificate i		
=	<ul> <li>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 intransit may only be stored in free zones, free warehouses and custom warehouses.</li> </ul>						
-	<ul> <li>Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the border inspection post of entry into the</li> </ul>						
-	04.04; 04.	ence I.19: use the			ed System (HS) code under the following headings: 04. 5.02, 15.03, 15.04, 23.01, 23.09; 28.35.25; 28.35.26; 35		
-					er number and the seal number (if applicable) must be g an feeding of farmed animals, other than fur animals, a		
_	or manufa	cturing of pet food			ther it is a transit or an import certificate.	•	

### **COUNTRY: UNITED STATES**

# Processed petfood other than canned petfood

II.	Healt	h information	II.a.	Certificate reference No	II.b.			
	<ul> <li>Box reference I.28: Species: select from the following: Aves, Ruminantia, Suidae, Mammalia other than Ruminantia or Suidae, Pesca, Mollusca, Crustacea, Invertebrates other than Mollusca and crustacea. Part II:</li> </ul>							
Part		soa, Monasoa, Orastaoca, Invertebrates	Juici ui	an Wondsod and Grastacca. Tart II.				
(1a)								
(1b)	OJ L 54, 26.2.2011, p. 1.							
(2)	Delete a	as appropriate.						
( <sup>2a</sup> )	OJ L 13	39, 30.4.2004, p. 55.						
( <sup>2b</sup> )	OJ L 12	25, 23.5.1996, p. 3.						
(3)	OJ L 17	75, 10.7.2010, p. 1.						
(4)	Where:							
	n =	number of samples to be tested;						
	m = threshold value for the number of bacteria; the result is considered satisfactory if the number of bacteria in all sar does not exceed m;							
	M = maximum value for the number of bacteria; the result is considered unsatisfactory if the number of bacteria in one of more samples is M or more; and							
	c = number of samples the bacterial count of which may be between m and M, the sample still being considered accept if the bacterial count of the other samples is m or less.							
(5)	OJ L 14	7, 31.5.2001, p. 1.						
(6)	OJ L 17	72, 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 entry into the European Union.</li> </ul>							
Official veterinarian/Official inspector								
	Name	(in capital letters):	Qι	ualification and title:				
	Dot		C:	aneture				
	Date:		51	gnature:				
			St	tamp:				