CHAPTER 3(A)

Health certificate

For canned petfood intended for dispatch to or for transit through² the European Union

NTRY: UNITED STATES		ertificate to EU	
I.1. Consignor Name	I.2. Certificate reference No	I.2.a.	
Address			
	I.3. Central competent authority		
T.	APHIS-\	/S	
Tel.			
I.5. Consignee Name	I.6. Person responsible for the load in EU Name		
Address	Address		
Postal code	Postal code		
Tel.	Tel.		
I.7. Country of ISO code I.8. Region of Code		egion ofeod	
origin origin	destination code de	estination	
I.11. Place of origin	I.12. Place of destination		
Name	Custom wa	arabawaa 🗆	
Approval number	Name Approval i		
Address	Address		
	Postal code		
Name			
Approval number			
Address			
Nama			
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)		
	23.09.10		
	I.20. Quantity		
I.21. Temperature of product	I.22. Number of packages		

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Canned Petfood COUNTRY: UNITED STATES I.2. Certificate reference No I.2.a. I.23. Seal/Container No I.24. Type of packaging I.25. Commodities certified for: Animal feedingstuff Technical use I.26. For transit through EU to third country I.27. For import or admission into EU ISO code Third country I.28. Identification of the commodities Species Nature of commodity Batch number (Scientific name) Manufacturing plant

	l						
	II.	II. Health information			II.a. Certificate reference No	II.b.	
]	I, the unde	ersigne	d official veterinarian, decl	lare 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 Annex XIII, Chapter II and Annex XIV, Chapter II, thereof and certify that the petfood described above:					
ion	II.1.	. has been prepared and stored in an establishment or plant approved and supervised by the competent authority accordance with Article 24 of Regulation (EC) No 1069/2009;					
icat	II.2.	has been prepared exclusively with the following animal by-products:					
Part II: Certification		⁽²⁾ either	[-	carcases and parts of a killed, and which are fit f	inimals slaughtered or, in the case of game, bodie for human consumption in accordance with Union I sumption for commercial reasons;]		
Part		⁽²⁾ and/or	[-	carcases and the followi slaughterhouse and wer mortem inspection or b consumption in accordar	ing parts originating either from animals that have lead considered fit for slaughter for human consumpt sodies and the following parts of animals from gance with Union legislation:	tion following an ante- ame killed for human	
				consumption in a disease communi	dies and parts of animals which are rejected accordance with Union legislation, but which did n icable to humans or animals;		
	1			(ii) heads of poultry;			
					including trimmings and splitting thereof, horns a the carpus and metacarpus bones, tarsus and r an ruminants;		
				(iv) pig bristles;			
				(v) feathers;]			
		⁽²⁾ and/or	-[-	humans or animals, obta slaughterhouse after ha	a did not show any signs of disease communica ained from animals other than ruminants that have eving been considered fit for slaughter for human of tion in accordance with Union legislation;]	been slaughtered in a	
		(2)and/or	-[-	animal by-products arisi including degreased bor	ing from the production of products intended for ne, greaves and centrifuge or separator sludge from	human consumption, n milk processing;]	
		⁽²⁾ and/or	-[-	intended for human con	n, or foodstuffs containing products of animal origin nsumption for commercial reasons or due to proble other defects from which no risk to public or animal	ems of manufacturing	
		⁽²⁾ and/or	-[- -	petfood and feedingstuf derived products, which	ffs of animal origin, or feedingstuffs containing a n are no longer intended for feeding for commerce ing or packaging defects or other defects from whi	animal by-products or sial reasons or due to	
		⁽²⁾ and/or	-[blood, placenta, wool, fe	eathers, hair, horns, hoof cuts and raw milk original is of any disease communicable through that p		
		⁽²⁾ and/or	-[-	aquatic animals, and par	rts of such animals, except sea mammals, which d ble to humans or animals;]	id not show any signs	
		(2)and/or	-[-		rom aquatic animals originating from plants for human consumption;]	or establishments	
		(2)and/or	[-	the following material communicable through the	originating from animals which did not show a that material to humans or animals:	iny signs of disease	
				` '	sh with soft tissue or flesh;		
					inating from terrestrial animals:		
				 hatchery by- eggs, 	products,		
					ucts, including egg shells;		
				•••	led for commercial reasons;]		
		⁽²⁾ and/or	-[-	animal by-products fron	n aquatic or terrestrial invertebrates other than s	pecies pathogenic to	
				humans or animals;]			

Canned Petfood

II.	Health info	rmation	II.a. Certificate reference No	II.b.	
	⁽²⁾ and/or	pursuant to Directive 96	which have been treated with certain substances which have been treated with certain substances which with the substances which is also with the certain substances which have been treated with high permitted in ation (EC) No 1069/2009;]		
II.3.	has been s	subjected to heat treatment to a m	inimum Fc value of 3 in hermetically sealed containers	;	
II.4.	was analysed by a random sampling of at least five containers from each processed batch by laboratory diagnostic methods to ensure adequate heat treatment of the whole consignment as foreseen under point II.3;				
II.5.	has undergone all precautions to avoid contamination with pathogenic agents after treatment.				
II.6.	[the product does not contain and is not derived from specified risk material as defined in Annex Regulation (EC) No 999/2001 of the European Parliament and of the Council ⁽³⁾ or mechar separated meat obtained from bones of bovine, ovine or caprine animals; and the animals from this product is derived have not been slaughtered after stunning by means of gas injected into cranial cavity or killed by the same method or slaughtered by laceration of central nervous tissumeans of an elongated rod-shaped instrument introduced into the cranial cavity;]				
	⁽²⁾ or	those derived from animals bor	and is not derived from bovine, ovine or caprine main, continuously reared and slaughtered in a country or k by a decision in accordance with Article 5(2) of Re	region classified	
II.7.	in addition (2) either	ovine or caprine origin, the ovideen kept continuously since by restriction is imposed due to a the last three years: (i) it has been subject to require (ii) no classical scrapic case has been diagnosed or, the last sheet all animals in which all goats and sheet rams of the ARR/A no VRQ allele, (iii) ovine and caprine animal introduced into the ho	intended for feeding ruminants and containing milk or ine and caprine animals from which these products in irth or for the last three years on a holding where no consuspicion of TSE and which has satisfied the following gular official veterinary checks; e, as defined in point 2(g) of Annex I to Regulation (Elellowing the confirmation of a classical scrapic case: classical scrapic was confirmed have been killed and destroyed, expending the holding have been killed and destroyed, expending entry and breeding ewes carrying at least one als, with the exception of sheep of the ARR/ARR pridling only if they come from a holding which contains the contains the contains and contains the contain	are derived have official movement requirements for Grant Region 1. The second requirements for destroyed, and cept for breeding ARR allele and on genetype, are	
	⁽²⁾ or	requirements set out in p [in case of animal by products ovine or caprine origin, and des (EC) No 546/2006 ⁽⁴⁾ , the ovine kept continuously since birth or restriction is imposed due to a sthe last seven years: (i) it has been subject to req (ii) no classical scrapie case has been diagnosed or, if all animals in which all goats and sheel rams of the ARR/A no VRQ allele; (iii) ovine and caprine animals	points (i) and (ii).] intended for feeding ruminants and containing milk or stined to a Member State listed in the Annex to Command caprine animals from which these products are dor for the last seven years on a holding where no consuspicion of TSE and which has satisfied the following gular official veterinary checks; e, as defined in point 2(g) of Annex I to Regulation (Effollowing the confirmation of a classical scrapic case: classical scrapic was confirmed have been killed and depend the holding have been killed and destroyed, extended and provided the provided and breeding ewes carrying at least one cals, with the exception of sheep of the ARR/ARR pridling only if they come from a holding which contains the contents of the provided and the confirmation of the provided and the provided and the contents of the provided and t	r milk products of ission Regulation erived have been official movement requirements for EC) No 999/2001, destroyed, and cept for breeding a ARR allele and on genetype, are	
Note	es				

II.	Health information	II.a.	Certificate reference No	II.b.

Part I:

- Box reference I.6: Person responsible for the consignment in the European Union: this box is to be filled in only if it is
 a certificate for transit commodity; it may be filled in if the certificate is for import commodity.
- Box reference I.12: Place of destination: this box is to be filled in only if it is a certificate for transit commodity. The products in transit can 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 event of unloading and reloading.
- Box reference I.23: for bulk containers, the container number and the seal number (if applicable) should be given.
- Box reference I.25: technical use: any use other than for animal consumption.
- Box reference I.26 and I.27: fill in according to whether it is a transit or an import certificate.

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⁾ OJ L 147, 31.5.2001, p. 1.
- ⁽⁴⁾ OJ L 94, 1.4.2006, p. 28.
- 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 has to accompany the consignment until it reaches the border inspection post.

	1 1	1 1
Offici	al veterinarian/Official inspector	
	Name (in capital letters):	Qualification and title:
	Date:	Signature:
		Stamp: