## CHAPTER 3(F)

# Health certificate

For animal by-products  $^{(3)}$  for the manufacture of petfood, intended for dispatch to  $\frac{1}{2}$  or  $\frac{1}{2}$  the European Union

cou	INTRY: UNITED STATES	Veterinary certificate to EU				
	I.1. Consignor Name Address	I.2. Certificate reference No I.2.a.				
	Tel.	I.3. Central competent authority				
		APHIS-VS  I.4. Local competent authority				
		, ,				
Į.	I.5. Consignee Name Address Postcode Tel.	I.6. Person responsible for the load in EU Name Address Postcode Tel.				
men						
sign						
ed con	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 Name	I.12. Place of destination				
Part I: Details of dispatched consignment	Approval number Address	Custom warehouse  Name Approval number Address Postcode				
	Name Approval number Address  Name Approval pumber Address	I.14. Date of departure				
	I.13. Place of loading					
	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	Tallibor of packages				
	I.23. Seal/Container No	I.24. Type of packaging				

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**COUNTRY: UNITED STATES** 

# Animal by-products for the manufacture of petfood

			I.2. Certific	ate reference N	0	1.2.a.
I.25. Commodities	certified for:					
Manufacture o	f petfood $\square$	Further proc	ess 🗆		Tech	nical use □
I.26. For transit thro	ough EU to third country		I.27. For imp	ort or admission	into EU	
Third country	ISC	) code				
I.28. Identification of	of the commodities					
Species (Scientific name)	Nature of commodity	Approval number of es Manufacturing	stablishments plant	Number of packages	Net weight	Batch number

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	II. He	alth informati	on	II.a.	Certificate reference No	II.b.		
		of the Eu	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( <sup>1a</sup> ) and Commission Regulation (EU) No 142/2011( <sup>1b</sup> ), and in particular Chapter II of Annex XIV thereto, and certify that the animal by-products described above:					
u C		II.1.1. II.1.2.	I.1.1. consist of animal by-products that satisfy the animal health requirements below;					
catio		<del>(²)either</del>			se birth or for a period of at least three	months preceding		
Part II: Certification		<del>(<sup>2</sup>)or</del> ( <sup>2</sup> )or	[that were killed in the wild in this [that are animals of the zoologic or aquatic invertebrates;]		<del>'Y(<sup>1d</sup>);]</del> rs Rodentia or Lagomorpha, aquatic al	nimals or terrestrial		
<u> </u>	<del>(²)either</del>	[II.1.3.	have been obtained or produced any epizootic disease, and which		animals which were net slaughtered or	r killed to eradicate		
Pa			(a) come from holdings who	<del>ere</del>				
			Implementing Re rinderpest, News peried of the pres peried of the pres	es for which the animals are listed in (EU) 2018/1882, there has been no isease or highly pathogenic avian in 30 days, nor of classical or African swindays, nor in the holdings situated in the period of the preceding 30 days; and	o case/outbreak of fluenza during the ne fever during the			
	(ii) where ther period of th				ne case/outbreak of foot and mouth of 0 days, nor in the holdings situated in the period of the preceding 30 days; and			
		( <sup>2</sup> )either	departure and which ha	ave bec	of origin for a period of at least 40 days on transported directly to the slaughter h did not comply with the same health	rhouse without any		
	territory of the third				ler veterinary supervision in the third or origin from which imports of fresh me tions in accordance with Implementin nouse	at of ungulates are		
			preceding the tin	ne of s	nertem health inspection during the plaughter and have shown ne evidence the animals are susceptible; and			
			the relevant prov	/isions	re and at the time of slaughter or killing of Union legislation and have met red down in Chapters II and III of Council I	quirements at least		
	<del>(²)or</del>	[II.1.3.	( //11	d from	animals which were not killed to erad	icate any epizootic		
			' following disease Regulation (EU) disease or highly days, nor of clas	25 km s for wh 2018 pathog	n the wild in an area:  radius there has been no case/outb iich the animals are listed in accordance (1882: foot-and-mouth disease, rind enic avian influenza during the period African swine fover during the period	with Implementing erpest, Newcastle of the preceding 30		
			country not authouse the preceding 30	orised for days o	at least 20 km from any country or part or export to the European Union of pour of porcine material during the precedi	ıltry material during ng 40 days; and		
			chilling either to a colle	ection	orted within a period of 12 hours follo centre and immediately afterwards to ame-handling establishment;]			
		II.1.4.	no case/outbreak of the disease during the period of the precedin raw material for exportation to the	es referi g 30 da ne Euro	t around which, within a radius of 10 k red to in point II.1.3 for which the anim lys or, in the event of a case of disease pean Union has been authorised only a fection of the establishment under the	als are susceptible t, the preparation of after the removal of		

## Animal by-products for the manufacture of petfood

II.	Health informatio	n	II.a. Certificate reference No	II.b.
	II.1.5.		red without contact with any other material that ove, and it has been handled so as to avoid co	
	II.1.6.	have been packed in new pack bearing the label indicating 'RA'	aging preventing any leakage and in officially s W MATERIAL ONLY FOR THE MANUFACTURE e establishment of destination in the European U	E OF PETFOOD'
	II.1.7.	consist only of the following anir	nal by-products:	
	<del>(²)either</del>	animals killed which we	animals slaughtered or, in the case of game, been deemed fit for human consumption in according to the case of game, but the dealers of the case of game, but the case of game,	<del>lance with Union</del>
	<del>(²)and/or</del>	[- carcases and the following consumption following	ibly declared as animal by-products for commerci lowing parts originating either from animals ghterhouse and were considered fit for slaug an ante-mortem inspection or bodies and the food and for human consumption in accordance with Un	that have been ghter for human ollowing parts of
		consumption in a of disease comm	lies and parts of animals which are rejected as accordance with Union legislation, but which did no nunicable to humans or animals;	
			; including trimmings and splitting thereof, horns a and the carpus and metacarpus bones, tarsus	
		(v) feathers;]		
	<del>(²)and/or</del>	. , .	rising from the production of products inten degreased bone, greaves and centrifuge or sepa	
	<del>(²)and/or</del>	longer intended for hur	n, or foodstuffs containing products of animal original consumption for commercial reasons or duging defects or other defects from which no risk to	e to problems of
	<del>(²)and/or</del>		rts of such animals, except sea mammals, which- nunicable to humans or animals;]	did not show any
	(²)and/or		om aquatic animals originating from plants or s for human consumption;]	r establishments
	<del>(<sup>2</sup>)and/or</del>	-	originating from animals which did not show any that material to humans or animals:	signs of disease
		(i) shells from shell	fish with soft tissue or flesh;	
		,,	ginating from terrestrial animals:	
		— hatchery by	<del>/ products,</del>	
		- egg by-pro	ducts, including egg shells;	
			illed for commercial reasons:1	
	<del>(²)and/or</del>	( )	a aquatic or terrestrial invertebrates, other than sp	ecies pathogenic
	(²)and/or		eof of the zoological orders of Rodentia and Lag	omornha evcent
	( )arraror	Category 1 material as	referred to in Article 8, point (a)(iii), (iv) and (v), of egory 2 material as referred to in Article 9, points	FRegulation (EC)
	<del>(<sup>2</sup>)and/or</del>	prohibited by Council D	which have been treated with certain substairective 96/22/EC( <sup>4a</sup> ), the import of the material b 35, point (a)(ii) of Regulation (EC) No 1069/2009	eing permitted in
	II.1.8.	Union legislation in such a way	ant of origin or have been preserved in accordanc that they will not spoil between dispatch and del Inion or during the transit through the European U	ivery to the plant
	II.1.9.	prohibited by Directive 96/22/E	ed from animals which have been treated with ce C for the manufacture of petfood, the import be t (a)(ii) of Regulation (EC) No 1069/2009:	

### Animal by-products for the manufacture of petfood

### Health information Certificate reference No it has been marked in the third country before entry into the territory of the European Union by a cross of liquefied charcoal or activated carbon on each outer side of each frozen block, or, when the raw material is transported in pallets which are not divided into separate consignments during transport to the petfood plant of destination in the European Union or during the transit through the European Union, on each outer side of each pallet, in a way that the marking covers at least 70 % of the diagonal length of the frozen block and be of at least 10 cm width: (b) in the case of material which is not frozen, the raw material has been marked in the third country before entry into the territory of the European Union by spraying it with liquefied charcoal or by applying charcoal powder in such a way that the charcoal is clearly visible on the material; and where the animal by-products are made up of raw material which has been treated as (c) referred to above and other non-treated raw material, all the raw materials have been marked as referred to in point (a) and (b) above. $(^{2})(^{5})[11.2.$ Specific requirements (2)(6)[II.2.1 The by products in this consignment come from animals that have been kept in the territory referred to in point (II.1.2), where vaccination programmes against foot and mouth disease are being regularly carried out and officially controlled in domestic bovine animals.] (2)(7)[II.2.2. The by products in this consignment consist only of animal by products derived from trimmed offal of domestic ruminants, which have maturated at an ambient temperature of more than + 2 °C for a period of at least three hours, or in the case of masseter muscles of bovine animals and deboned meat of domestic animals, for a period of at least 24 hours. Il (2)[II.3. the animal by products for the manufacture of petfood contains or is obtained from animal by products of ruminant origin and: [originate from a country or region, which is classified as posing a negligible bovine spongiform (2)either encephalopathy (BSE) risk in accordance with Commission Decision 2007/453/EC(8), and in which there has been no indigenous BSE case;]] [originate from a country or region classified as posing a negligible BSE risk in accordance with Decision 2007/453/EC in which there has been an indigenous BSE case, and the animal byproducts or derived products were derived from animals born after the date from which the ban on the feeding of ruminants with meat-and-bone meal and greaves derived from ruminants, as defined in the World Organisation for Animal Health (OIE) Terrestrial Animal Health Code, has been effectively enforced in that country or region, and [are derived from ruminants other than bovine, ovine or caprine animals.]]] fare derived from bevine, evine or caprine animals born, continuously reared and (2)or slaughtered in a country or region classified as posing a negligible BSE risk in accordance with Decision 2007/453/EC.]]] (do not contain: specified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001 of the European Parliament and of the Council(9): mechanically separated meat obtained from bones of bovine. 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 Decision 2007/453/EC, in which there has been no indigenous BSE case. animal by-product obtained from bovine, ovine or caprine animals which have been killed, after stunning, by laceration of the central nervous tissue by means of an elongated rod shaped instrument introduced into the cranial cavity, or by means of gas injected into the cranial cavity, except for those animals that were born, continuously reared and slaughtered in a country or region classified as posing a negligible BSE risk in accordance with Decision 2007/453/EC.111 Notes 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 a commodity to be imported into the European Union.

## Animal by-products for the manufacture of petfood

II.		Health information	II.a.	Certificate reference No	II.b.				
	_	Box reference I.12: Place of destination: this box is to be filled i stored in free zones, free warehouses and custom warehouses.	n only if	it is a certificate for a transit commodity. Products in tra	insit may only be				
	<ul> <li>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.</li> </ul>								
	_	Box reference I.19: use the appropriate Harmonized System (HS	S) code:	05.04; 05.06; 05.07; 05.11.91 or 05.11.99; 23.01; 41.01					
	_	Box reference I.23: for bulk containers, the container number ar		, , , ,					
	<ul> <li>Box reference I.25: technical use: any use other than feeding of farmed animals, other than fur animals, and the production or manufacturing of petfood.</li> </ul>								
	_	Box reference I.26 and I.27: fill in according to whether it is a tra	nsit or a	n import certificate.					
	_	Box reference I.28:  ———————————————————————————————————	NTIA S	UIDAE, MAMMALIA OTHER THAN RUMINANTIA OR S	SUIDAE PESCA				
		MOLLUSCA, CRUSTACEA, INVERTEBRATES O  manufacturing plant: provide the veterinary control	THER T	HAN MOLLUSCA AND CRUSTACEA;	.0.27.2, 1.2007.,				
Part II	:								
( <sup>1a</sup> )	OJ I	L 300, 14.11.2009, p. 1.							
( <sup>1b</sup> )		L 54, 26.2.2011, p. 1.							
(1c)	The	name and ISO code number of the exporting country as laid dow	n in:						
	_	Part 1 of Annex XIII to Commission Implementing Regulation (		1/404 (OJ L 114, 31.3.2021, p. 1);					
		Part 1 of Annex XIV to Implementing Regulation (EU) 2021/40 Part 1 of Annex I to Commission Regulation (EC) No 119/2009		9, 10.2.2009, p. 12).					
	In a	ddition the ISO code of regionalisation in the abovementioned Ani			nust be included.				
( <sup>1d</sup> )	Only Unio	y for countries from which game meat intended for human consur on.	nption of	the same animal species is authorised for importation in	nto the European				
( <sup>2</sup> )		ete as appropriate.							
( <sup>3</sup> )	of th	luding raw blood, raw milk, hides and skins, hooves and horn, pig lese products).	bristles a	and feathers (see relevant specific certificates in that An	nex for the import				
( <sup>4</sup> )		L 303, 18.11.2009, p. 1.							
( <sup>4a</sup> ) ( <sup>5</sup> )		L 125, 23.5.1996, p. 3. Interpretary guarantees to be provided when the material of don	nestic ru	minants originated in the territory of a South American	or South African				
( )	Supplementary guarantees to be provided when the material of domestic ruminants originated in the territory of a South American or South African country or part thereof from where only maturated and deboned fresh meat of domestic ruminants for human consumption is permitted for exportation to the European Union. The whole masseter muscles of bovine animals, incised in accordance with Section IV, Chapter I, Part B.1, of Annex I to Regulation (EC) No 854/2004 of the European Parliament and of the Council (OJ L 139, 30.4.2004, p. 206), are also permitted.								
( <sup>6</sup> )	-	y for certain South American countries.		, , , , , , , , , , , , , , , , , , , ,					
( <sup>7</sup> )	Only for certain South American and South African countries.								
(8)	OJ L 172, 30.6.2007, p. 84.								
( <sup>9</sup> )	OJ L 147, 31.5.2001, p. 1.								
	<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	ial ve	eterinarian/Official inspector							
	Na	me (in capital letters): Qualif	cation	and title:					
	_		4						
	Da	ate: Signa	iure:						
		Stam	p:						
		J	•						