CHAPTER 8

Health certificate

For animal by-products to be used for purposes outside the feed chain or for trade samples², intended for dispatch to or for transit through² the European Union

cou	NTRY: UNITED STATES	Veterinary certifi	cate to EU			
	I.1. Consignor Name Address	I.2. Certificate reference No	I.2.a.			
		I.3. Central competent authority				
	Tel.	I.4. Local competent authority				
	Consignee Border inspection post through which consignment is intended to leave the EU	I.6. Person responsible for the load in EU Name Address				
	Name Address	Postal code Tel.				
nment	Postal code Tel.					
l consig	I.7. Country of origin ISO code I.8. Region of origin Origin	I.9. Country of ISO I.10. Region of destination code destination	eode			
tchec	I.11. Place of origin	I.12. Place of destination				
Part I : Details of dispatched consignment	Name Approval number Address	Custom warehouse Ap number Address	pproval			
: Deta		Postal code				
Part I						
	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 □	I.17.				
	Identification	1.17.				
	Documentation references					
	I.18. Description of commodity	I.19. Commodity code (HS code	e)			
		I.20. Quantity				
	I.21. Temperature of product Ambient □ Chilled □	I.22. Number of packages				
	I.23. Seal/Container No	I.24. Type of packaging				

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Animal by-products to be used for purposes outside the feed chain or for trade samples $^{(2)}$

			I.2. Certific	ate reference N	0	1.2.a.	
I.25. Commodities of	ertified for:						
Technical use □							
			T				
I.26. For transit thro	ugh EU to third country		I.27. For imp	ort or admission	into EU		
Third country	ISO	code					
I.28. Identification of	f the commodities						
Species (Scientific name)	Nature of commodity	Approval number of es Manufacturing	stablishments plant	Number of packages	Net weight	Batch number	

Animal by-products to be used for purposes outside the feed chain or for trade samples (2)

	II.	Health information		II.a. Certificate reference No	II.b.				
	-	European Parliamen	and of the	inarian, declare that I have read and understood Regulation (EC) e Councii ^(1a) and Commission Regulation (EU) No 142/2011 ^(1b) , and ify that the animal by-products described above:	No 1069/2009 of the d in particular Annex				
	⁽²⁾ Ⅱ.1.	definition No 39 of A	are trade samples which consist of animal by products intended for particular studies or analyses as referred to i definition No 39 of Annex I to Commission Regulation (EU) No 142/2011, that are bearing the label 'TRADE SAMPLI NOT FOR HUMAN CONSUMPTION'; or						
Z	⁽²⁾ II.2.	satisfy the animal hea		•					
ij	II.2.1.	have been							
ifica		⁽²⁾ either [(a)	obtained thereof:.	from materials imported from third country, t	erritory or part e species to the EU;]				
Part II: Certification		⁽²⁾ and/or [(b)	obtained from anii either	in the exporting country, territory or part thereof:nals	(3)				
art II:			(i) 7	That have remained in this territory or in a region eligible to exposible to the EU since birth or for at least the last three months beformed in the wild in this territory. (A)					
ď		⁽²⁾ and/or [(c)	are deriv	ved from eggs, milk, rodents, lagomorphs, or aquatic animals or rates;]	•				
	II.2.2.			than derived from eggs, milk, rodents, lagomorphs, or aquatic anin obtained from animals:	mals or terrestrial or				
		(2) either [(a)		rom holdings:					
			f F	where, for the following diseases for which the animals are suscep- neither case/outbreak of rinderpest, swine vesicular disease, Newca- nathogenic avian influenza during the prior 30 days, nor of classi ever during the prior 40 days; nor in the holdings situated in their during the prior 30 days; and	stle disease or highly ical or African swine				
			(ii) v	where there has been neither case/outbreak of foot-and-mouth disc 50 days, nor in the holdings situated in their vicinity within 25 km lays; and	0 1				
		——————————————————————————————————————	which:						
			(ii) t	vere not killed to eradicate any epizootic disease; have remained in their holdings of origin for at least 40 days before have been transported directly to the slaughterhouse without contac which did not comply with the same health conditions;					
			`´´ ‡	at the slaughterhouse, have passed the ante-mortem health inspection during the 24 hours before the slaughter and have shown no evidence of the diseases referred to above for which the animals are susceptible; and					
		(2)	8 4	have been handled in the slaughterhouse before and at the time of accordance with the relevant provisions of Union legislation and ha at least equivalent to those laid down in Chapters II and III of Counci 1099/2009;	ve met requirements				
		(2) or [(a)	•	l and killed in the wild in an area:					
			f e	n which within 25 km there has been no case/outbreak of any of the or which the animals are susceptible: foot and mouth disease, ri- disease or highly pathogenic avian influenza during the prior 30 day Mrican swine fever during the prior 40 days; and	nderpest, Newcastle				
			(ii) t	hat is situated at a distance that exceeds 20 km from the borders erritory of a country or part thereof, which is not authorised at thes his material to the European Union; and					
		(b)		ter killing were transported within 12 hours for chilling either to a detely afterwards to a game establishment, or directly to a game establishment, or directly to a game establishment.					
	II.2.3.	establishment around II.2.2 for which the ar of raw material for ex	(2) in the case of materials other than materials derived from wild caught fish or invertebrates, have been obtained in an establishment around which, within a radius of 10 km, there has been no case/outbreak of diseases referred to in point II.2.2 for which the animals are susceptible during the prior 30 days or, in the event of a case of disease, the preparation of raw material for exportation to the European Union has been authorised only after removal of all meat, and the total cleaning and disinfection of the establishment under the control of an official veterinarian;						
	II.2.4.		have been obtained and prepared without contact with other material not complying with the conditions required above, and it has been handled so as to avoid contamination with pathogenic agents;						
	II.2.5.								

Animal by-products to be used for purposes outside the feed chain or for trade samples (2)

II. Hea	alth information	ı	II.a. Certificate reference No	II.b.			
II.2.6.		of the fo	ollowing animal by-products:				
	⁽²⁾ either	- -	carcases and parts of animals slaughtered or, in the case of game, bodie killed, and which are fit for human consumption in accordance with Union kintended for human consumption for commercial reasons;]				
	⁽²⁾ and/or	-	carcases and the following parts originating either from animals that have to slaughterhouse and were considered fit for slaughter for human consumption mortem inspection or bodies and the following parts of animals from ga	on following an anto			
			consumption in accordance with Union legislation: (i) carcases or bodies and parts of animals which are rejected consumption in accordance with Union legislation, but which did not be consumption in accordance with Union legislation, but which did not be consumption in accordance with Union legislation.				
			disease communicable to humans or animals; (ii) heads of poultry;				
			(iii) hides and skins, including trimmings and splitting thereof, horns at phalanges and the carpus and metacarpus bones, tarsus and metatar				
			(iv) pig bristles; (v) feathers;]				
	⁽²⁾ and/or	-[-	animal by-products from poultry and lagomorphs slaughtered on the farm at 1(3)(d) of Regulation (EC) No 853/2004, which did not show any signs of di to humans or animals;				
	⁽²⁾ and/or	[-	blood of animals which did not show any signs of disease communical humans or animals, obtained from animals other than ruminants that have tslaughterhouse after having been considered fit for slaughter for human c	peen slaughtered in			
	⁽²⁾ and/or	[-	an ante-mortem inspection in accordance with Union legislation;] animal by-products arising from the production of products intended for				
	⁽²⁾ and/or	[-	 including degreased bone, greaves and centrifuge or separator sludge from products of animal origin, or foodstuffs containing products of animal origin intended for human consumption for commercial reasons or due to problem 	, which are no longe			
	⁽²⁾ and/or	-1-	packaging defects or other defects from which no risk to public or animal her petfood and feedingstuffs of animal origin, or feedingstuffs containing a	alth arises;]			
			derived products, which are no longer intended for feeding for commerci problems of manufacturing or packaging defects or other defects from whice animal health arises;]	al reasons or due			
	⁽²⁾ and/or	[-	 blood, placenta, wool, feathers, hair, horns, hoof cuts and raw milk original that did not show signs of any disease communicable through that pr animals;] 				
	⁽²⁾ and/or	[-	 aquatic animals, and parts of such animals, except sea mammals, which di of diseases communicable to humans or animals; 	d not show any sigr			
	⁽²⁾ and/or	[-	animal by-products from aquatic animals originating from establishments or products for human consumption;]	plants manufacturir			
	⁽²⁾ and/or	-[-	the following material originating from animals which did not show as communicable through that material to humans or animals:	ny signs of diseas			
			(i) shells from shellfish with soft tissue or flesh;				
			(ii) the following originating from terrestrial animals:				
			(iii) day-old chicks killed for commercial reasons;				
	⁽²⁾ and/or	-[-	animal by-products from aquatic or terrestrial invertebrates, other than sphumans or animals;	pecies pathogenic			
	⁽²⁾ and/or	[-	animals and parts thereof of the zoological orders of Rodentia and Category 1 material as referred to in Article 8(a)(iii), (iv) and (v) of Regulatio and Category 2 material as referred to in Article 9(a) to (a) of that Regulation	n (EC) No 1069/200			
	⁽²⁾ and/or	- [-	fur originating from dead animals that did not show clinical signs of any di through that product to humans or animals;]				
II.2.7.			zen at the plant of origin or have been preserved in accordance with EU legis ill between dispatch and delivery to the plant of destination.	slation in such a wa			
⁽²⁾⁽⁵⁾ [II.2.8.	Specific requirements						
⁽²⁾⁽⁶⁾ II.2.8.1.	(II.2.1), wh∈	re vacci	this consignment come from animals that have been obtained in the territorination programmes against foot-and-mouth disease are being regularly carries bevine animals.				
⁽²⁾⁽⁷⁾ II.2.8.2.			his consignment consist of animal by-products derived from offal or deboned m	eat.]			
II.2.9.	⁽²⁾ either	[the	product does not contain and is not derived from specified risk material as dulation (EC) No 999/2001 of the European Parliament and of the Counc	efined in Annex V			

Animal by-products to be used for purposes outside the feed chain or for trade samples⁽²⁾

II.	Health information		II.a. Certificate reference No	II.b.			
		this product is d cavity or killed b	botained from bones of bovine, ovine or caprine animals; and the erived have not been slaughtered after stunning by means of gas injudy the same method or slaughtered by laceration of central nervous d-shaped instrument introduced into the cranial cavity;	ected into the crania			
	⁽²⁾ or	derived from ar	es not contain and is not derived from bovine, ovine or caprine mate nimals born, continuously reared and slaughtered in a country or gible BSE risk by a decision in accordance with Article 5(2) of	region classified a			
II.2.10.	in addition as	regards TSE:					
	⁽²⁾ either	ovine or caprine kept continuous restriction is imp the last three ye		e derived have bee			
		(ii) no class	een subject to regular official veterinary checks; sical scrapie case, as defined in point 2(g) of Annex I to Regulation in diagnosed or, following the confirmation of a classical scrapie case				
			animals in which classical scrapie was confirmed have been killed a				
		all rai	goats and sheep on the holding have been killed and destroyed, ms of the ARR/ARR genotype and breeding ewes carrying at least VRQ allele:	except for breeding			
		introduc	nd caprine animals, with the exception of sheep of the ARR/ARR ed into the holding only if they come from a holding which nents set out in points (i) and (ii).]				
	⁽²⁾ or	ovine or caprine (EC) No 546/20	nal by-products intended for feeding ruminants and containing mill be origin, and destined to a Member State listed in the Annex to Co 106 ⁽⁹⁾ , the ovine and caprine animals from which these products are	mmission Regulation e derived have been			
		kept continuously since birth or for the last seven years on a holding where no of restriction is imposed due to a suspicion of TSE and which has satisfied the following r the last seven years:					
		(ii) no class	pen subject to regular official veterinary checks; sical scrapie case, as defined in point 2(g) of Annex I to Regulation				
	has been diagnosed or, following the confirmation of a classical scrapic case —— all animals in which classical scrapic was confirmed have been killed a						
		- all rai	goats and sheep on the holding have been killed and destroyed, ms of the ARR/ARR genotype and breeding ewes carrying at least VRQ allele:	except for breedin			
	(iii) ovine and caprine animals, with the exception of sheep of the ARR/ARR prion introduced into the holding only if they come from a holding which comprequirements set out in points (i) and (ii).]						
Notes							
Part I:			e for the consignment in the European Union: this box is to be f	illed in only if it is			
_	Box reference I.11: In		be filled in if the certificate is for import commodity. ments for the particular technological studies or analyses: indicate i	name and address of			
_	establishment only. Box reference I.11 and I.12: Approval number: the registration number of the establishment or plant, which has been issued by the competent authority.						
_	Box reference I.12: Place of destination: this box is to be filled in: — products for the manufacture of derived products for uses outside the feed chain: 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. — products for the particular technological studies or analyses: the EU plant indicated in authorisation of competent authority when appropriate.						
_	Box reference I.15: F be provided. In case	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 BIP of entry into the EU.					
_	30.01.						
_	Box reference I.25: to	reference I.23: for bulk containers, the container number and the seal number (if applicable) should be included. reference I.25: technical use: any use other than for animal consumption. reference I.25: for the purposes of the certificate, 'technical use' includes use as a trade sample.					
_	Box reference I.26: for the purposes of the certificate, technical dee includes due as a trade sample. Box reference I.26 and I.27: except for trade samples, which are not sent in transit, fill in according to whether it is a transit or an import certificate. Box reference I.28:						
	— products		are of derived products for uses outside the feed chain: Manufacturi of the approved establishment.	ng plant: provide th			

Animal by-products to be used for purposes outside the feed chain or for trade samples⁽²⁾

II.	Health information	II.a.	Certificate reference No	II.b.			
	 products for the particular technological studies or analyses: the EU plant indicated in authorisation of competent authority when appropriate. Species: select from the following: Aves, Ruminantia, Mammalia other than Ruminantia, Pesca, Mollusca, Crustacea, invertebrates other than Mullusca and Crustacea. 						
(1a) (1b) (2) (3) (4) (5)	(tb) OJ L 54, 26.2.2011, p. 1. Delete as appropriate. The name and ISO code number of the exporting country as laid down in: — Part 1 of Annex II to Regulation (EU) No 206/2010; — the Annex to Regulation (EC) No 119/2009. In addition the ISO code of territories and parts thereof referred to in Regulations mentioned in this footnote (where applicable for the susceptible species concerned) should be included. (4) Only for countries from where game meat intended for human consumption of the same animal species is authorised for importation into the European Union. (5) 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 Annex I, Section IV, Chapter I, Part B(1) of Regulation (EC) No 854/2004 of the European Parliament and of the Council, are also permitted. (6) Only for certain South American countries. Only for certain South American and South African countries. OJ L 147, 31.5.2001, p. 1.						
	Date:		Signature: Stamp:				