CHAPTER 8

Health certificate

For animal by-products to be used for purposes outside the feed chain or for trade samples (2), intended for dispatch to or for transit through (2) the European Union

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	
nent	I.5. Consignee Name Address Postcode Tel.	I.6. Person responsible for the load in EU Name Address Postcode Tel.	
consignn	I.7. Country of ISO code I.8. Region of Code origin	I.9. Country of ISO I.10. Region of destination code destination	Code
ched	I.11. Place of origin	I.12. Place of destination	-
Part I : Details of dispatched consignment	Name Approval number Address	Custom warehouse Name Approval number Address Postcode	1
	Name Approval number Address Name Approval number		
	Address I.13. Place of loading	I.14. Date of departure	
	I.15. Means of transport Aeroplane Ship Railway wagon Road vehicle Other Identification	I.16. Entry BIP in EU	
	Documentation references		
	I.18. Description of commodity	I.19. Commodity code (HS code)	
		I.20. Quantity	
	I.21. Temperature of product Ambient □ Chilled □ Frozen □	I.22. Number of packages	
	I.23. Seal/Container No	I.24. Type of packaging	

Page	of

COUNTRY: UNITED STATES

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

	I.2. Certifi	cate reference N	lo	1.2.a.
I.25. Commodities certified for:				
Technical use □				
I.26. For transit through EU to third country	1.27. For im	port or admission	into EU	
Third country ISO code				
1.28. Identification of the commodities				
Species Nature of commodity Approval r (Scientific name) Ma	number of establishments nufacturing plant	Number of packages	Net weight	Batch number

Page ___ of ___

	II.	Health information			II.a. Certificate reference No	II.b.		
		No 1069/20	009 of th	ne European Parliame	declare that I have read and understont and of the Council(1a), and Commission of Annex XIV thereto, and certify that the	Regulation (EU) No		
	described above							
ıtion		(²)either	analy: Regul	ses as referred to i	onsist of animal by products intended for the definition of trade samples in poin 11, that bear the label 'TRADE SAMPLE	t 39 of Annex I to		
ifice		(²)or	[satisfy the animal health requirements set out in point II.1.];					
Cert	II.1. II.1.1.	The animal have been	by prod	ducts described above				
Part II: Certification		(²)either	[(a)		t, territory or part thorised to export			
Ъ		(²)and/or	[(b)	obtained in t	e exporting third country, te			
					(³) fron	n animals that		
				fresh meat to	d in that third country, territory or part ther the European Union since birth or for a p se months before the date of slaughter; ar	period of at least the		
					the wild in that third country, territory or pa			
		(²)and/or	[(c)		milk, rodents, lagomorphs, or aquatic anir	nals or terrestrial or		
	(²)[II.1.2.				না als derived from eggs, milk, rodents, lagon vertebrates and unprocessed furs, have			
		(²)either	[(a)	coming from holding	s:			
				there has n disease, New period of the the period o	e following diseases for which the animal bet been any case/outbreak of rinderpe castle disease or highly pathogenic avian preceding 30 days, nor of classical or Africa the preceding 40 days; nor in the holding a 10 km radius, during the period of the	st, swine vesicular influenza during the in swine fever during ings situated in their		
				during the pe	has not been any case/outbreak of foot- riod of the preceding 60 days, nor in the vithin a 25 km radius, during the period	holdings situated in		
			(b)	which:				
				(i) were not kille	d to eradicate any epizootic disease;			
				[(²)either				
				the date of slaughterhou	their holdings of origin for a period of at longer transport se without contact with other animals which the conditions;	ed directly to the		
				(²)or				
				or part of the meat of ungu	d on holdings under veterinary supervision erritory of the third country of origin from w ates are authorised without any restriction mplementing Regulation (EU) 2021/404]	hich imports of fresh		
				period of 24	erhouse, passed the ante-mortem health i lours before the time of slaughter and sho referred to above for which the animals ar	wed no evidence of		

II. H	ealth information			II.a. Certificate reference No	II.b.
			killing in accord	n the slaughterhouse before and at the time of dance with the relevant provisions of Union leg requirements at least equivalent to those la I III of Council Regulation (EC) No 1099/2009(5)]	gislation and aid down in
	(²)or	[(a)	captured and killed in	the wild in an area:	
				25 km radius there has been no case/outbreak	
			disease, rinder during the perio	ses for which the animals are susceptible: foo pest, Newcastle disease or highly pathogenic avi and of the preceding 30 days nor of classical or A period of the preceding 40 days; and	an influenza
			another territor	at a distance that exceeds 20 km from the border y of a third country or part thereof, which is not a the exportation of such material to the Europear	uthorised at
		(b)	a collection centre and	e transported within a period of 12 hours for chil Himmediately afterwards to a game establishmer	
(²)[II.1.3.	been obtain case/outbre of the prece of raw mater	ed in a ak of dis ding 30 rial for e	an establishment arouseases referred to in pole days or, in the event of exportation to the Europ	Is derived from fish or invertebrates caught in the nd which, within a radius of 10 km, there haint II.1.2 for which the animals are susceptible during a case/outbreak of one of those diseases, the sean Union was authorised only after the removate establishment under the control of an official variations.	as been no ring a period preparation I of all meat,
II.1.4.				contact with other material which does not comen handled so as to avoid contamination with	
II.1.5.	cleaned and in container 'ANIMAL BY OUTSIDE T	disinfers seale -PROD HE FE	cted before use and, in d under the responsibi DUCTS ONLY FOR TH	ich prevents any leakage or in packaging whice the case of consignments shipped other than via lity of the competent authority, bearing the lab E MANUFACTURE OF DERIVED PRODUCTS ame and address of the establishment of destir	parcel post, el indicating FOR USES
II.1.6.	European U		ollowing animal by-pro	duate:	
11.1.0.	(²)either		• • • •	animals slaughtered or, in the case of game, bo	dies or parts
	()ouro r	ı	of animals killed which	were deemed fit for human consumption in according to the cons	ordance with
	(²)and/or	[-	slaughtered in a slaug consumption following	ollowing parts originating either from animals ghterhouse and were considered fit for slaughte an ante-mortem inspection or bodies and the fole killed for human consumption in accordance	er for human llowing parts
			human consum	dies and parts of animals which were rejected ption in accordance with Union legislation, but w of disease communicable to humans or animals	hich did not
			` '	s, including trimmings and splitting thereof, hor nalanges and the carpus and metacarpus bones	
			(iv) pig bristles;		
			(v) feathers;]		
	(²)and/or	[-	referred to in Article 1(rom poultry and lagomorphs slaughtered on 3), point (d), of Regulation (EC) No 853/2004 of the Council (20), which did not show any signs cans or animals;]	ne European

II. Heal	n information II.a. Certificate reference No II.b.
	(2) and/or [blood of animals which did not show any signs of disease communicable through blood to humans or animals, obtained from animals that have been slaughtered
	a slaughterhouse after having been considered fit for slaughter for hum
	consumption following an ante-mortem inspection in accordance with Un
	legislation;]
	(2)and/or [animal by products arising from the production of products intended for hum
	consumption, including degreased bone, greaves and centrifuge or separa
	sludge from milk processing;]
	(2)and/or [- products of animal origin, or foodstuffs containing products of animal origin, wh
	are no longer intended for human consumption for commercial reasons or due
	problems of manufacturing or packaging defects or other defects from which risk to public or animal health arises;]
	(²)and/or [- petfood and feedingstuffs of animal origin, or feedingstuffs containing animal
	products or derived products, which are no longer intended for feeding
	commercial reasons or due to problems of manufacturing or packaging defects
	other defects from which no risk to public or animal health arises;]
	(2)and/or [blood, placenta, wool, feathers, hair, horns, hoof cuts and raw milk originating fr
	live animals that did not show signs of any disease communicable through t product to humans or animals;]
	(²)and/or [aquatic animals, and parts of such animals, except sea mammals, which didshow any signs of diseases communicable to humans or animals;]
	(2)and/or [- animal by products from aquatic animals originating from establishments or pla
	manufacturing products for human consumption;]
	(2)and/or [- the following material originating from animals which did not show any signs
	disease communicable through that material to humans or animals:
	(i) shells from shellfish with soft tissue or flesh;
	(ii) the following originating from terrestrial animals:
	- hatchery by products;
	
	 egg by products, including egg shells;
	(iii) day old chicks killed for commercial reasons;]
	(2)and/or [animal by products from aquatic or terrestrial invertebrates, other than spec pathogenic to humans or animals;]
	(²)and/or [- animals and parts thereof of the zoological orders of Rodentia and Lagomorp
	except Category 1 material as referred to in Article 8, point (a)(iii), (iv) and (v) Regulation (EC) No 1069/2009 and Category 2 material as referred to in Article
	points (a) to (g), of that Regulation;]
	(2)and/or [furs originating from dead animals that did not show clinical signs of any disection communicable through that product to humans or animals;]
II.1.7.	have been deep-frozen at the plant of origin or have been preserved in accordance with Europe Union legislation in such a way that they will not spoil between the time of dispatch and the time delivery to the plant of destination.
(²)(⁶)[II.1.8.	
(2)(²)	
	he animal by products in this consignment come from animals that have been obtained in the coun
•	territory or part thereof referred to in point II.1.1, where vaccination programmes against foot and mo disease are regularly carried out and officially controlled in domestic bovine animals.]]
(²)(⁸)	
	The animal by products in this consignment consist of animal by products derived from offal or debor meat.]]
(²)[II.1.9.	the animal by products described above
()[{²)either _ [are derived from ether ruminants than bovine, evine or caprine animals.]]
	(2)or [are derived from bovine, ovine or caprine animals and does not contain and is not derived.

Page	of	

II. H	ealth informatior	ı		II.a. Certificate reference No	II.b.
		(²) either	born, continuous posing a negli	 nd caprine materials other than those do ly reared and slaughtered in a country or gible bovine spongiform encephalopa	region classified a
		(²)or	-[(a) specifie	Commission Decision 2007/453/EC(⁹).]] drisk material as defined in point 1 of An	nex V to Regulatio
			, ,	999/2001 of the European Parliament ar ically separated meat obtained from bor	•
			continuc as posi	ne animals, except from those anima ously reared and slaughtered in a country ng a negligible BSE risk in accorda 3/EC, in which there has been no indige	/ or region classifie ance with Decisio
			(c) animal t	by product or derived product obtained from the same of the students which have been killed, after students of the students of the same of	om bovine, ovine c
			instrume injected born, ce classifi e	entral nervous tissue by means of an election introduced into the cranial cavity, or into the cranial cavity, except for those entinuously reared and slaughtered in a dasposing a negligible BSE risk in accor	or by means of ga animals that wer a country or regio
II.1.10	the animal	by-products	2007/45 described above:	3/EC.]]]	
	(²)either	-	•	roducts of ovine or caprine animal origin other than fur animals.]	or are not intende
	(²)or	•	•	of ovine or caprine animal origin and a In fur animals, and the milk or milk produ	
		` '	nce birth in a cour classical (ne and caprine animals which have bee try where the following conditions are ful cerapie is compulsorily notifiable; ness, surveillance and monitoring systematic;	filled:
		(ii	case of a	trictions apply to holdings of ovine or ca suspicion of transmissible spongiform on firmation of classical scrapie;	prine animals in the common transfer of the c
		(i)	v) ovine and destroyed	caprine animals affected with classical s	crapie are killed ar
		(√	greaves, ; Organisal banned a	ig to ovine and caprine animals of mea as defined in the Terrestrial Animal Healt ion for Animal Health (OIE), of rumina nd effectively enforced in the whole coun preceding seven years;	h Code of the Wor ant origin has bee
		` '		ings where no official restrictions are	imposed due to
		dı	riginate from holdi uring the period of ase of classical sci	ngs where no case of classical scrapie he the preceding seven years or, following t apie:	nas been diagnose the confirmation of
		(2	destroyed genotype	and caprine animals on the holding had or slaughtered, except for breeding rare breeding ewes carrying at least one ARI other ovine animals carrying at least one	ns of the ARR/AR Rallele and no VR
		(2	or [all anima and destr least two scrapie c negative laboratory Regulatio	Is in which classical scrapie was confirm byed, and the holding has been subject years since the date of confirmation case to intensified TSE monitoring, incresults for the presence of TSE in a methods set out in Chapter C, point a (EC) No 999/2001, of all of the following of 18 months, except ovine anima	ed have been killed for a period of of the last classic cluding testing with the coordance with the 3.2, of Annex X g animals which a

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

II.	Health information	II.a.	Certificate reference No	II.b.
	and	uio Wiii	ch have been slaughtered for human c	onoumption,
	were	not k	illed in the framework of a disease	

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 to be transited through the European Union; it may be filled in if the certificate is for a commodity to be imported into the European Union.
- Box reference I.11: In the case of consignments for trade samples or analyses: indicate the name and address
 of the 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 for the following products:
 - products for the manufacture of derived products for uses outside the feed chain: only if it is a certificate for a transit commodity. Products in transit may only be stored in free zones, free warehouses and custom warehouses.
 - products for trade samples or analyses: the plant in the European Union indicated in the authorisation of the competent authority where appropriate.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In the case of unloading and reloading in the European Union, the consignor must inform the border inspection point of the point of entry into the European Union.
- Box reference I.19: use the appropriate Harmonized System (HS) code under the following headings: 04.01;
 04.02; 04.03; 04.04; 04.08; 05.05; 05.06, 05.07; 05.11.91; 05.11.99, 23.01 or 30.01.
- Box reference I.23: for bulk containers, the container number and the seal number (if applicable) must be included.
- 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;
 - for the purposes of the certificate, 'technical use' includes use 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 for the manufacture of derived products for uses outside the feed chain: Manufacturing plant: provide the veterinary control number of the approved establishment;
 - products for the particular technological studies or analyses: the plant in the European Union indicated in the authorisation of the competent authority where appropriate;
 - 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:

- ^(1a) OJ L 300, 14.11.2009, p. 1.
- (1b) OJ L 54, 26.2.2011, p. 1.
- (2) Delete as appropriate.
- ^(2a) OJ L 139, 30.4.2004, p. 55.
- The name and ISO code number of the exporting country as laid down in:
 - Part 1 of Annex XIII to Commission Implementing Regulation (EU) 2021/404 (OJ L 114, 31.3.2021, p. 1);
 - Part 1 of Annex XIV to Implementing Regulation (EU) 2021/404, and
 - Annex I to Commission Regulation (EC) No 119/2009 (OJ L 39, 10.2.2009, p. 12).

In addition the ISO code of territories and parts thereof referred to in the Annexes to Implementing Regulation (EU) 2021/404 and to Regulation (EC) No 119/2009 referred to in this note (where applicable for the susceptible species concerned) must be included where applicable.

II.	Health information	II.a.	Certificate reference No	II.b.
(4)	Only for countries from where the game meat inter	ded fo	human consumption of the same	animal species is
(5)	authorised for importation into the European Union. OJ L 303, 18.11.2009, p. 1.			
(6)	Supplementary guarantees to be provided where the a South American or South African country or part the domestic ruminants for human consumption is aumasseter muscles of bovine animals, incised in acc B.1, of Annex I to Regulation (EC) No 854/2004 (30.4.2004, p. 206), are also permitted.	ereof fi horised ordance	om where only maturated and debo for exportation to the European with the requirements of Section	oned fresh meat of Union. The whole IV, Chapter I, Part
(7)	Only for certain South American countries.			
(8)	Only for certain South American and South African	ountrie	5.	
(9) (10)	OJ L 172, 30.6.2007, p. 84.			
(10)	OJ L 147, 31.5.2001, p. 1.			
_	The signature and the stamp must be in a different or Note for the person responsible for the consignment purposes and must accompany the consignment until the European Union.	in the	European Union: this certificate is	
Offic	cial veterinarian/Official inspector			
	Name (in capital letters): Qualif	cation	and title:	
	Data.	turo.		
	Date: Signa	ture.		