Commission Regulation (EU) No 294/2013 of 14 March 2013 amending and correcting Regulation (EU) No 142/2011 implementing Regulation (EC) No 1069/2009 of the European Parliament and of the Council laying down health rules as regards animal by-products and derived products not intended for human consumption and implementing Council Directive 97/78/EC as regards certain samples and items exempt from veterinary checks at the border under that Directive (Text with EEA relevance)

ANNEX

The Annexes to Regulation (EU) No 142/2011 are amended as follows:

- (1) Annex I is amended as follows:
 - (a) point 19 is replaced by the following:
 - 19. "**petfood**" means feed, other than material referred to in Article 24(2), for use as feed for pet animals, and dogchews consisting of animal by-products or derived products which:
 - (a) contain Category 3 material, other than material referred to in Article 10(n), (o) and (p) of Regulation (EC) No 1069/2009; and
 - (b) may contain imported Category 1 material comprising of animal by-products derived from animals which have been submitted to illegal treatment as defined in Article 1(2)(d) of Directive 96/22/EC or Article 2(b) of Directive 96/23/EC;;
 - (b) point 23 is replaced by the following:
 - 23. "digestion residues" means residues, including the liquid fraction, resulting from the transformation of animal by-products in a biogas plant;
- (2) in Annex IV, Chapter IV, Section 3 is amended as follows:
 - (a) point 1 is amended as follows:
 - (i) point (a)(iii) is replaced by the following:
 - (iii) transformed into biogas. In such case the digestion residues must be disposed of in accordance with point (i) or (ii), except where the material results from processing in accordance with point 2(a) or (b) where the residues can be used in accordance with the conditions set out in point 2(a) or point 2(b)(iii) as appropriate; or;
 - (ii) point (b)(i) is replaced by the following:
 - (i) disposed of as provided for in point 1(a)(i) or (ii), with or without prior processing as provided for in Article 13(a) and (b) and Article 14(a) and (b) of Regulation (EC) No 1069/2009;;
 - (b) points 2(b)(ii) and (iii) are replaced by the following:
 - (ii) in the case of potassium sulphate, used for direct application to land or for the production of derived products for application to land;
 - (iii) in the case of glycerine derived from Categories 1 and 2 material which has been processed in accordance with processing method 1 as set out in Chapter III:
 - used for technical purposes,

- transformed into biogas, in which case the digestion residues may be applied to land within the national territory of the producing Member State, subject to the decision of the competent authority, or
- used for denitrification in a waste water treatment plant, in which case the residues of the denitrification may be applied to land in accordance with Council Directive 91/271/EEC⁽¹⁾;
- (iv) in the case of glycerine derived from Category 3 material:
 - used for technical purposes,
 - transformed into biogas, in which case the digestion residues may be applied to land, or
 - used for feeding, provided that the glycerine is not derived from Category 3 material referred to in Article 10(n), (o) and (p) of Regulation (EC) No 1069/2009;;
- (c) point 3 is replaced by the following:
 - 3. Any waste other than animal by-products and derived products provided for in point 2, resulting from the processing of animal by-products in accordance with this Section, such as sludge, filter contents, ash and digestion residues, shall be disposed of in accordance with Regulation (EC) No 1069/2009 and with this Regulation.;
- in Annex V, Chapter I, Section 1, point 2(d) is replaced by the following:
 - (d) animal by-products which may be applied to land without processing in accordance with Article 13(f) of Regulation (EC) No 1069/2009 and with this Regulation, if the competent authority does not consider them to present a risk of spreading any serious transmissible disease to humans or animals;;
- (4) in Annex VI, Chapter II, Section 1, the introductory phrase is replaced by following:

Categories 2 and 3 materials as referred to in Article 18(1) of Regulation (EC) No 1069/2009 may be fed to the animals referred to in paragraph (1)(a), (b), (d), (f), (g) and (h) of that Article subject to compliance with at least the following conditions, in addition to any conditions laid down by the competent authority in accordance with Article 18(1) of that Regulation:;

- (5) Annex VIII is amended as follows:
 - (a) in Chapter II, point 2(b), point (xix) is replaced by the following:
 - (xix) in the case of manure which has been subject to the lime treatment set out in point I of Section 2 of Chapter IV of Annex IV, the words "manure-lime-mixture";
 - in the case of processed manure which has been subject to the treatment set out in point (b) and (c) of Section 2 of Chapter I of Annex XI, the words "processed manure".;
 - (b) the following Chapter VI is added:

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

CHAPTER VI

TRANSPORT OF DEAD PET ANIMALS

The conditions in points 1 to 3 of Article 48 of Regulation (EC) No 1069/2009 regarding the advance authorisation by the competent authority in the Member States of destination and the use of TRACES shall not be required in the case of the transport of a dead pet animal for incineration in an establishment or plant located in the border region of another Member State sharing a common border when the Member States conclude a bilateral agreement on the condition of the transport.;

- (6) in Annex X, Chapter II is amended as follows:
 - (a) in Section 4, Part II, point 1 is replaced by the following:
 - 1. The requirements laid down in points 2 and 3 of this Part shall apply to the processing, use and storage of milk, milk-based products and milk-derived products which are Category 3 material, as referred to in Article 10(e) of Regulation (EC) No 1069/2009, other than centrifuge or separator sludge, and milk, milk-based products and milk-derived products referred to in Article 10(f) and (h) of that Regulation, that have not been processed in accordance with Part I of this Section.;
 - (b) Section 10 is replaced by the following:

Section 10

Specific requirements for feeding to farmed animals, other than fur animals, of certain Category 3 material referred to Article 10(f) of Regulation (EC) No 1069/2009

Category 3 material comprising of foodstuffs containing products of animal origin originating from Member States which are no longer intended 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, referred to in Article 10(f) of Regulation (EC) No 1069/2009, may be placed on the market for feeding to farmed animals, other than fur animals, without further treatment, provided that the material:

- (i) has undergone processing as defined in Article 2(1)(m) of Regulation (EC) No 852/2004 or in accordance with this Regulation;
- (ii) is composed of or contain one or more of the following Category 3 materials referred to in Article 10(f) of Regulation (EC) No 1069/2009:
 - milk,
 - milk-based products,
 - milk-derived products,
 - eggs,

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

- egg products,
- honey,
- rendered fats,
- collagen,
- gelatine;
- (iii) has not been in contact with any other Category 3 materials; and
- (iv) all necessary precautions have been taken to prevent the contamination of the material.;
- in Annex XI, Chapter II, Section 1, point 1(b) is replaced by the following:
 - (b) using processed animal protein, including processed animal protein produced in accordance with point B.1(b)(ii) of Section 1 of Chapter II of Annex X, which has been produced from Category 3 material in accordance with Section 1 of Chapter II of Annex X, or materials which have been subject to another treatment, where such materials may be used for organic fertilisers and soil improvers in accordance with this Regulation; or;
- (8) Annex XIII is amended as follows:
 - (a) in Chapter VI, points C(1)(c) and (d) are replaced by the following:
 - (c) have been subject to an anatomical preparation such as by plastination;
 - (d) are animals of the biological class Insecta or Arachnida which have been subject to a treatment, such as drying, to prevent any transmission of diseases communicable to humans or animals; or
 - (e) are objects in natural history collections or for the promotion of science and they have been:
 - (i) preserved in media, such as alcohol or formaldehyde, which allow display of the items; or
 - (ii) embedded completely on micro-slides;
 - (f) are processed DNA samples intended for repositories for the promotion of biodiversity research, ecology, medical and veterinary science or biology.;
 - (b) in Chapter XI, the following point is added:
 - 3. End point for products derived from rendered fats:

Fat derivatives which have been processed as referred to in point 1 may be placed on the market for uses indicated in point 2 without restrictions in accordance with this Regulation.;

- (9) Annex XIV is amended as follows:
 - (a) in Chapter I, Section 1 is amended as follows:
 - (i) points (c), (d) and (e) are replaced by the following:

- (c) they must come from a third country or part of a third country listed in the column "third countries' list" of Table 1;
- (d) they must come from an establishment or plant which is registered or approved by the competent authority of the third country, as applicable, and which is on the list of such establishments and plants referred to in Article 30; and
- (e) they must be:
 - (i) accompanied during transportation to the point of entry into the Union where the veterinary checks take place by the health certificate referred to in the column "certificates/model documents" of Table 1; or
 - (ii) presented at the point of entry into the Union where the veterinary checks take place accompanied by a document corresponding to the model referred to in the column "certificates/model documents" of Table 1.;
- (ii) point (f) is deleted;
- (b) in Chapter II, Section 1 is amended as follows:
 - (i) points (c), (d) and (e) are replaced by the following:
 - (c) they must come from a third country or part of a third country listed in the column "third countries' list" of Table 2;
 - (d) they must come from an establishment or plant which is registered or approved by the competent authority of the third country, as applicable, and which is on the list of such establishments and plants referred to in Article 30; and
 - (e) they must be:
 - (i) accompanied during transportation to the point of entry into the Union where the veterinary checks take place by the health certificate referred to in the column "certificates/model documents" of Table 2; or
 - (ii) presented at the point of entry into the Union where the veterinary checks take place accompanied by a document corresponding to the model referred to in the column "certificates/model documents" of Table 2.;
 - (ii) point (f) is deleted;
 - (iii) Table 2 is amended as follows:

row No 13 is replaced by the following:

	nMaterial		Third	Annex
innards	referred		gountries	
for the	to in	innards	listed	Chapter
manufac		must	in Part	
of	35(a)	have	1 of	
petfood		been	Annex	
		produced		
		in	Regulati	on
		accordar		
		with	No	
		Chapter	206/2010),
		III of	from	
		Annex	which	
		XIII.	Member	
			States	
			authorise	.
			imports	
			of fresh	
			meat	
			from	
			the	
			same	
			species	
			and	
			where	
			only	
			bone in	
			meat is	
			authorise	ed.
			In the	
			case of	
			flavourir	ng
			innards	
			from	
			fish	
			materials	5,
			third	,
			countries	5
			listed	
			in	
			Annex	
			II to	
			Decision	
			2006/760	
			EC.	
			In the	
			case of	
			flavourir]]g
			innards	5
			LILLIAI GO	

		poultry meat third countries listed in Part 1 of Annex I to Regulation (EC) No 798/2008, from which Member States authorise imports of fresh poultry	

- in row No 14, point (a) in the third column is replaced by the following:
 - (a) Category 3 materials referred to in Article 10(a) to (m).;
- rows Nos 15 and 16 are replaced by the following:

15	Animal	Category	The	Third	Annex
	by-	3	products	countries	sXV,
	products	materials	shall	listed	Chapter 3(D).
	for use	referred	comply	in Part	•
	as raw	to in	with	1 of	
	petfood	Article	the	Annex	
	_	10(a)	requirem	elhtso	
		and	set	Regulati	on
		Article	out in	(EU)	
		10(b)	Section	No	
		(i) and	8.	206/2010)
		(ii).		or in	
				Annex	
				I to	
				Regulati	on
				(EC)	
				No	
				798/2008	3,
				from	
				which	
				Member	
				States	
				authorise	

				imports	
				of fresh	
				meat	
				from	
				the	
				same	
				species	
				and	
				where	
				only	
				bone in	
				meat is	
				authorise	ed.
				In the	
				case	
				of fish	
				materials	5,
				third	
				countries	S
				listed	
				in	
				Annex	
				II to	
				Decision	
				2006/766	5/
				EC.	
16	Animal	Category	The	Third	Annex
	by-	3		countries	
		materials		listed	Chapter 3(D).
	for use	referred		in part	1
	in feed	to in	with	1 of	
	for fur	Article	the	Annex	
	animals	10(a) to	requirem	eth tso	
		(m)	set	Commis	sion
			out in	Regulati	on
			out in Section	Regulati	on
				Regulati	on
			Section	Regulati (EU)	
			Section	Regulati (EU) No	
			Section	Regulation (EU) No 206/2010	
			Section	Regulation (EU) No 206/2010 or in Annex I to),
		,	Section	Regulati (EU) No 206/2010 or in Annex I to Regulati),
			Section	Regulati (EU) No 206/2010 or in Annex I to Regulati (EC)),
			Section	Regulati (EU) No 206/2010 or in Annex I to Regulati (EC) No	on
			Section	Regulati (EU) No 206/2010 or in Annex I to Regulati (EC) No 798/2008	on
			Section	Regulati (EU) No 206/2010 or in Annex I to Regulati (EC) No 798/2008 from	on
			Section	Regulati (EU) No 206/2010 or in Annex I to Regulati (EC) No 798/2008 from which	on
			Section	Regulati (EU) No 206/2010 or in Annex I to Regulati (EC) No 798/2003 from which Member	on
			Section	Regulati (EU) No 206/2010 or in Annex I to Regulati (EC) No 798/2000 from which Member States	on 3,
			Section	Regulati (EU) No 206/2010 or in Annex I to Regulati (EC) No 798/2000 from which Member States authorise	on 3,
			Section	Regulati (EU) No 206/2010 or in Annex I to Regulati (EC) No 798/2008 from which Member States authorise imports	on 3,
			Section	Regulati (EU) No 206/2010 or in Annex I to Regulati (EC) No 798/2000 from which Member States authorise	on 3,

		meat
		from
		the
		same
		species
		and
		where
		only
		bone in
		meat is
		authorised.
		In the
		case
		of fish
		materials,
		third
		countries
		listed
		in
		Annex
		II to
		Decision
		2006/766/
		EC.

- in row No 17, third column, point (a) is replaced by the following:
 - (a) In the case of materials destined for the production of biodiesel or oleochemical products: Categories 1, 2 and 3 materials referred to in Articles 8, 9 and 10.;
- row No 18 is replaced by the following:

18	Fat	(a)	Time fat	Any	(a)	In
	derivati	ves	decivativ	ethird		the
			slast	country.		case
			co mply			of
			waitth			fat
			therivativ	es		derivatives
			feq uiren	ents		for
			sustes			uses
			outside			outside
			Section			the
			f@ed			feed
			chain			chain
			for			for
			farmed			farmed
			animals:			animals:
			Category	y		Annex
			1			XV,
			material	s		Chapter
			referred			14(A).

	to in Article 8(b),	(b)	In the case of
	(c) and (d), Category		fat derivatives for use as
	materials referred to in Article		feed: Annex XV, Chapter 14(B).
	9(c) and (d) and		14(<i>D)</i> .
	Article 9(f) (i) and Category		
	materials referred to in		
(b)	Article 10. In the		
	case of fat derivatives		
	for use as feed: Category		
	materials other than		
	materials referred to in		
	Article 10(n), (o)		

	and	
	(p);	

- (c) in Chapter II, Section 2, point 2 is replaced by the following:
 - 2. The blood from which blood products for the manufacture of derived products for uses outside the feed chain for farmed animals are produced must have been collected under veterinary supervision:
 - (a) in slaughterhouses:
 - (i) approved in accordance with Regulation (EC) No 853/2004; or
 - (ii) approved and supervised by the competent authority of the country of collection; or
 - (b) from live animals in facilities approved and supervised by the competent authority of the country of collection.;
- (d) in Chapter II, Section 3, point 1 is replaced by the following:
 - 1. The blood must comply with the conditions set out in point 1(a) of Chapter IV of Annex XIII and must be collected under veterinary supervision:
 - (a) in slaughterhouses:
 - (i) approved in accordance with Regulation (EC) No 853/2004; or
 - (ii) approved and supervised by the competent authority of the country of collection; or
 - (b) from live equidae in facilities approved and furnished with a veterinary approval number and supervised by the competent authority of the country of collection for the purpose of collecting blood from equidae for the production of blood products for purposes other than feeding.;
- (e) in Chapter II, Section 3, point 2(d) is replaced by the following:
 - in the case of blood products other than serum and plasma, vesicular stomatitis for a period of at least six months.;
- (f) in Chapter II, Section 9, point (a)(i) is replaced by the following:
 - (i) in the case of materials destined for the production of biodiesel or oleochemical products, animal by-products referred to in Articles 8, 9 and 10 of Regulation (EC) No 1069/2009;;
- (10) Annex XV is amended as follows:
 - (a) Chapter 3(B) is replaced by the following:

CHAPTER 3(B)

Health certificate

For processed petfood other than canned petfood, intended for dispatch to or for transit through $\binom{2}{1}$ the European Union

_	INTR	те Еигореан Оп 1	ion								,
	l.1.	Consignor				1.2.	Certifica	te refer	ence No		1.2
		Name									_
		Address				1.3.	Central	compet	ent authority		
		Tel.				1.4.	Local co	mpeter	nt authority		_
Ħ	1.5.	Consignee				1.6.	Person	respons	ible for the lo	ad in	El
Ĕ		Name					Name				
sign		Address					Address				
cou		Postcode					Postcod	۵			
eq		Tel.					Tel.	•			
tch										T	_
sba	1.7.	Country of origin ISO of	ode I.8	Region of origin	Code	1.9.	Country destinati		ISO code	1.10). I
đ		1			1		dootiilati	···			`
Part I: Details of dispatched consignment	111	Place of origin				112	Place of	decting	ation		_
etai		-				"		acount			
ă		Name Address		Approval nur	nber		Name Address			stom prova	
Έ		Name		Annual number			Address		Αþ	prova	u n
ď		Address		Approval nur	nber		Postcod	e			
		Name		Approval nur	mher						
		Address		Approvarria	11001						
	I.13.	Place of loading				1.14.	Date of	departu	re		_
	I.15.	Means of transport				1.16.	Entry BI	P in EU	J		_
			Ship 🔲	Bailway			-				
			. —		wagon 🗌						
		_	Other	l		1.17.					_
		Identification									-
		Documentation references									
	I.18.	Description of commodity						I.19. C	commodity co	de (H	S
							l				
										1.20.	Qi
											_
	I.21.	Temperature of product		_					_	1.22.	Νι
		Ambient		Chilled				Frozen			
	1.23.	Seal/Container No								1.24.	Ту
	1.25.	Commodities certified for:									_
		Animal feedingstuff			Technical	use 🗌	l				
	1.26.	For transit through EU to	third cour	ntry		1.27.	For impo	rt or ad	mission into E	ΞU	Т
		Third country		ISO code							
	1.28.	Identification of the commo	odities			•					_
		Species (Scientific name)		Approv	al number of Manufacturi				Net we	ight	

СО	UNTRY			Processed petfood of
	II.	Health informa	ation	II.a. Certificate reference No II.b.
E		and of the Cou		and understood Regulation (EC) No 1069/2009 reof, and Commission Regulation (EU) No 142, and certify that the petfood described above:
Part II: Certification	II.1.	has been prepa (EC) No 1069/3		ised by the competent authority in accordance v
e E	II.2.	has been prepared	ared exclusively with the following animal by-pr	oducts:
Part		(²) either [, in the case of game, bodies or parts of animals on legislation, but are not intended for human c
		(²) and/or [considered fit for slaughter for human consum	ther from animals that have been slaughtered in aption following an ante-mortem inspection or bo umption in accordance with Union legislation:
				which are rejected as unfit for human consumptic signs of disease communicable to humans or a
			(ii) heads of poultry;	
			(iii) hides and skins, including trimmings and metacarpus bones, tarsus and metatarsu	splitting thereof, horns and feet, including the phase bones;
			(iv) pig bristles;	
			(v) feathers;]	
		(²) and/or [phs slaughtered on the farm as referred to in Assigns of disease communicable to humans or a
		(²) and/or [from animals other than ruminants that have	ns of disease communicable through blood to hi been slaughtered in a slaughterhouse after hav n ante-mortem inspection in accordance with U
		(²) and/or [- animal by-products arising from the production greaves and centrifuge or separator sludge f	on of products intended for human consumption, rom milk processing;]
		(²) and/or [aining products of animal origin, which are no to problems of manufacturing or packaging de ;]
		(²) and/or [feedingstuffs containing animal by-products or or reasons or due to problems of manufacturing or al health arises;]
		(²) and/or [blood, placenta, wool, feathers, hair, horns, he any disease communicable through that produced the state of	oof cuts and raw milk originating from live animal duct to humans or animals;]
		(²) and/or [aquatic animals, and parts of such animal communicable to humans or animals; 	ls, except sea mammals, which did not sho
		(²) and/or [- animal by-products from aquatic animals ori consumption;]	ginating from plants or establishments manufac
		(²) and/or [the following material originating from anima material to humans or animals:	als which did not show any signs of disease of
			(i) shells from shellfish with soft tissue or fl	esh;
			(ii) the following originating from terrestrial a	nimals:
			 hatchery by-products, 	
			— eggs,	
			- egg by-products, including egg shells	;
			(iii) day-old chicks killed for commercial reas	sons;]

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

II.		mation		Processed pe	
11.	Health infor	mation	II.a. C	Certificate reference No	II.b.
	(²) and/or	[- animal by-produ	ucts from aquatic or terrestrial inverte	brates other than species patho	ogenic (
	(²) and/or		rts thereof of the zoological orders of (iii), (iv) and (v) of Regulation (EC) No ulation;]		
	(²) and/or		animals which have been treated wit import of the material being permit		
II.3.					
	(²) either	[was subjected to	a heat treatment of at least 90 °C thr	roughout its substance;]	
	(²) or	[was produced as	regards ingredients of animal origin to	using exclusively products which	h had b
			animal by-products or derived product roughout its substance;	cts from meat or meat products	subject
		(b) in the case of	milk and milk based products,		
			e from third countries or parts of third 605/2010 (³) submitted to a pasteuris		
		Commissi	reduced to less than 6 from third co ion Regulation (EU) No 605/2010, fire phosphatase test;		
		605/2010	e from third countries or parts of thir , submitted to a sterilisation process a negative phosphatase test on its ow	or a double heat treatment whe	
		605/2010	e from third countries or parts of thir , where there has been an outbreak of oot-and-mouth disease has been carri	f foot-and-mouth disease in the la	ast 12 i
		either			
		— a steri	ilisation process whereby an Fc value	e equal or greater than 3 is ach	ieved
		or			
			ial heat treatment with a heating effect 72 °C for at least 15 seconds and sed by		
		either			
		which	ond heat treatment with a heating effe would be sufficient to produce a nego or dried milk-based products by a dry	ative reaction to a phosphatase	
		or			
		— an aci	dification process such that the pH h	nas been maintained at less tha	n 6 for
		treatment with	gelatine, produced using a process the acid or alkali, followed by one or more ated, extraction by heat, followed by	re rinses with subsequent adjust	tment o

(d) in the case of hydrolysed protein produced using a production process involving approprontal contamination of raw Category 3 material, and, in the case of hydrolysed protein entirely or hides and skins produced in a processing plant dedicated only to hydrolysed protein prowith a molecular weight below 10000 Dalton and a process involving the preparation of brining, liming and intensive washing followed by:

(i) exposure of the material to a pH of more than 11 for more than three hours at a temp and subsequently by heat treatment at more than 140 °C for 30 minutes at more than

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

Health information

II.

OUNTRY	Processed petfood of

(ii) exposure of the material to a pH of 1 to 2, followed by a pH of more than 11, followed for 30 minutes at 3 bar;

II.a. Certificate reference No

II.b.

- (e) in the case of egg products submitted to any of the processing methods 1 to 5 or 7, as Annex IV to Regulation (EU) No 142/2011; or treated in accordance with Chapter II of Regulation (EC) No 853/2004 of the European Parliament and of the Council (4);
- (f) in the case of collagen submitted to a process ensuring that unprocessed Category 3 treatment involving washing, pH adjustment using acid or alkali followed by one or more rin the use of preservatives other than those permitted by Union legislation being prohibited;
- (g) in the case of blood products, produced using any of the processing methods 1 to 5 or 7, as Annex IV to Regulation (EU) No 142/2011;
- (h) in the case of mammalian processed animal protein submitted to any of the processing met case of porcine blood, submitted to any of the processing methods 1 to 5 or 7 provided that heat treatment throughout its substance at a minimum temperature of 80 °C has been ap
-) in the case of non-mammalian processed protein with the exclusion of fishmeal submitte methods 1 to 5 or 7 as referred to in Chapter III of Annex IV to Regulation (EU) No 142/
- in the case of fishmeal submitted to any of the processing methods or to a method and pa the products complies with the microbiological standards for derived products set out in Regulation (EU) No 142/2011;
- (k) in the case of rendered fat, including fish oils, submitted to any of the processing methods the case of fish oil) as referred to in Chapter III of Annex IV to Regulation (EU) No 142/2011 with Chapter II of Section XII of Annex III to Regulation (EC) No 853/2004; rendered fats to be purified in such a way that the maximum level of remaining total insoluble impurities weight;
- (I) in the case of dicalcium phosphate produced by a process that
 - (i) ensures that all Category 3 bone-material is finely crushed and degreased with hot w hydrochloric acid (at a minimum concentration of 4 % and a pH of less than 1,5) over a
 - (ii) following the procedure under (i), applies a treatment of the obtained phosphoric liquid precipitate of dicalcium phosphate at pH 4 to 7; and
 - (iii) finally, air dries the precipitate of dicalcium phosphate with inlet temperature of 6 temperature between 30 °C and 65 °C;
- (m) in the case of tricalcium phosphate produced by a process that ensures
 - (i) that all Category 3 bone-material is finely crushed and degreased in counter-flow with than 14 mm);
 - (ii) continuous cooking with steam at 145 °C during 30 minutes at 4 bar;
 - (iii) separation of the protein broth from the hydroxyapatite (tricalcium phosphate) by cent
 - (iv) granulation of the tricalcium phosphate after drying in a fluid bed with air at 200 $^{\circ}\text{C}$;
- (n) in the case of flavouring innards, produced according to a treatment method and parame product complies with the microbiological standards referred to under point II.4.]
- (2) or [was subject to a treatment such as drying or fermentation, which has been authorised by
- (²) or [in the case of aquatic and terrestrial invertebrates other than species pathogenic to humans treatment which has been authorised by the competent authority and which ensures the unacceptable risks to public and animal health;]
- II.4. was analysed by a random sampling of at least five samples from each processed batch taken during or after plant and complies with the following standards (5):

Salmonella: absence in 25g: n = 5, c = 0, m = 0, M = 0,

Enterobacteriaceae: n = 5, c = 2, m = 10, M = 300 in 1 gram;

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Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

cou

II.

II.5.

II.6.

11.7.

II.8.

JNT	DV		Processed pe	offeed o			
INI	Health infor	mation	II.a. Certificate reference No	II.b.			
	has undergor	ne all precautions to avoid contamination with patho	ogenic agents after treatment;				
	was packed in new packaging, which, if the petfood is not dispatched in ready-to-sell packages on which it is content is destined for feeding to pets only, bear labels indicating "NOT FOR HUMAN CONSUMPTION";						
	(²) either	[the product does not contain and is not derived the 999/2001 of the European Parliament and of the Covine or caprine animals; and the animals from with means of gas injected into the cranial cavity or kit tissue by means of an elongated rod-shaped institute.	council (6) or mechanically separated match this product is derived have not alled by the same method or slaughter	neat obta been sla ed by la			
	(²) or	[the product does not contain and is not derived animals born, continuously reared and slaughtere decision in accordance with Article 5(2) of Regula	d in a country or region classified as				
	in addition as	regards TSE:					
	(²) either	[in case of animal by-products intended for feeding the ovine and caprine animals from which these pi three years on a holding where no official move satisfied the following requirements for the last th	roducts are derived have been kept comment restriction is imposed due to a	ntinuous			
		(i) it has been subject to regular official veterina	ry checks;				
		(ii) no classical scrapie case, as defined in point		999/200			

(2) or

[in case of animal by-products intended for feeding ruminants and containing milk or milk product and destined to a Member State listed in the Annex to Commission Regulation (EC) No 546/200 animals from which these products are derived have been kept continuously since birth or for holding where no official movement restriction is imposed due to a suspicion of TSE and which requirements for the last seven years:

- all animals in which classical scrapie was confirmed have been killed and destroyed, a

all goats and sheep on the holding have been killed and destroyed, except for breed genotype and breeding ewes carrying at least one ARR allele and no VRQ allele;

(iii) ovine and caprine animals, with the exception of sheep of the ARR/ARR prion genotype, are only if they come from a holding which complies with the requirements set out in points (i)

(i) it has been subject to regular official veterinary checks;

following the confirmation of a classical scrapie case:

- (ii) no classical scrapie case, as defined in point 2(g) of Annex I to Regulation (EC) No 999/200 following the confirmation of a classical scrapie case:
 - all animals in which classical scrapie was confirmed have been killed and destroyed, a
 - all goats and sheep on the holding have been killed and destroyed, except for breed genotype and breeding ewes carrying at least one ARR allele and no VRQ allele;
- (iii) ovine and caprine animals, with the exception of sheep of the ARR/ARR prion genotype, are only if they come from a holding which complies with the requirements set out in points (i)

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 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 be stored in free zones, free warehouses and custom warehouses.

Processed petfood of

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

II.	Health information	II.a. Certificate reference No	II.b.
-	Box reference I.15: Registration number (railway wagons or containe case of unloading and reloading, the consignor must inform the BIP		or name
-	Box reference I.19: use the appropriate Harmonized System (HS) cool 15.03, 15.04, 23.01, 23.09 or 35.02.	de under the following headings: 04.08	, 05.04,
-	Box reference I.23: for bulk containers, the container number and th	e seal number (if applicable) should b	e given.
-	Box reference I.25: technical use: any use other than for animal con	sumption.	
-	Box reference I.26 and I.27: fill in according to whether it is a transit	or an import certificate.	
-	Box reference I.28: Species: select from the following: Aves, Mamma	alia - Ruminantia, Pesca, Mollusca, C	rustacea
Pa	t 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 175, 10.7.2010, p. 1.		
(4)	OJ L 139, 30.4.2004, p. 55.		
(⁵)	Where:		
	n = number of samples to be tested;		
	m = threshold value for the number of bacteria; the result is consid m;	ered satisfactory if the number of bact	eria in al
	$M = \mbox{maximum value}$ for the number of bacteria; the result is consider or more; and	ered unsatisfactory if the number of ba	acteria in
	c = number of samples the bacterial count of which may be betwee count of the other samples is m or less.	en m and M, the sample still being co	nsidered
(⁶)	OJ L 147, 31.5.2001, p. 1.		
(7)	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 Europaccompany the consignment until it reaches the border inspection p		or veteri
Off	cial veterinarian/Official inspector		
	Name (in capital letters):	Qualifica	tion and
	Date:	Signatur	e:
	Stamp:		

(b) Chapter 3(D) is replaced by the following:

CHAPTER 3(D)

Health certificate

For raw petfood for direct sale or animal by-products to be fed to fur animals, intended for dispatch to or for transit through (2) the European Union

cou	JNTR	Υ										ν
	l.1.	Consignor Name					1.2.	Certifica	ite refe	rence No		1.2.
		Address					I.3. Central competent authority					
		Tel.					1.4.	Local co	ompete	nt authority		
ent	1.5.	I.5. Consignee				1.6.		respon	sible for the lo	ad ir	1 EU	
gnm		Name Address						Name Address				
nsi		Address						Address	•			
Part I: Details of dispatched consignment		Postcode						Postcod	le			
		Tel.						Tel.				
	1.7.	Country of origin ISO o	ode	1.8.	Region of origin	Code	1.9.	Country destinat		ISO code	1.1	0. R d
		Dia a a standada					140	D I				
	1.11.	Place of origin					1.12.	Place of	t destir			
	Name Address			Approval num	ber		Name Address			ustom oprova		
Pa		Name Address			Approval num	ber	Postcode					
		Name Address			Approval num	ber						
	I.13.	Place of loading					l.14.	Date of	depart	ure		
	I.15.	Means of transport					I.16.	Entry Bl	P in E	U		
		Aeroplane Ship Railwa			Railway v	wagon 🔲						
					ner 🗆							
		Identification					l.17.					
		Documentation references						-				
	I.18.	Description of commodity							I.19.	Commodity co	de (H	IS c
											1.20.	Qu
	1.21.	Temperature of product									1.22.	Nu
		Ambient			Chilled				Froze	n 🗆		
	1.23.	Seal/Container No									1.24.	Тур
	1.25.	Commodities certified for:										_
		Animal feedingstuff				Technical	use 🗌					
	1.26.	For transit through EU to	third c	ountry			1.27.	For impo	rt or a	dmission into	EU	
		Third country		I:	SO code							
	1.28.	Identification of the commo	odities									
		Species (Scientific name)	N	lature	of commodity	Appro		mber of our			Net v	weig
	I											

Raw petfood for direct sale or animal by COUNTRY II. Health information II.a. Certificate reference No IJЬ I, the undersigned official veterinarian, declare that I have read and understood Regulation (EC) No 1069/2009 and of the Council (¹a) and in particular Articles 10 thereof, and Commission Regulation (EU) No 142/2011 (¹b), of Annex XIII and Chapter II of Annex XIV thereto and certify that the raw petfood or animal by-products des Certification II.1. consist of animal by-products that satisfy the health requirements below; II.2. consist of animal by-products: (a) derived from meat which satisfies the relevant animal and public health requirements laid down in: Part Commission Regulation (EU) No 206/2010 (3) and provided the animals from which the meat is de countries, territories or parts thereof(ISO code in case of country or codes for territori that time (only as relevant for the susceptible species); (b) derived from animals that, at the slaughterhouse, have passed the ante-mortem health inspection durin slaughter and have shown no evidence of the diseases referred in the Regulations laid down in point (a susceptible; and (c) derived from animals that have been treated in the slaughterhouse before and at the time of slaughter or ki relevant provisions of Council Directive 93/119/EC (6) on the protection of animals at the time of slaughte (d) in the case of feed for fur animals derived from aquatic animals which satisfies the relevant animal and put down in Commission Decision 2006/766/EC (7), come from countries or territories thereof Annex II to that Decision: II.3.1. consist only of the following animal by-products: (a) carcases and parts of animals slaughtered or, in the case of game, bodies or parts of animals killed, a consumption in accordance with Union legislation, but are not intended for human consumption for comm (b) parts of slaughtered animals, which are rejected as unfit for human consumption but are not affected communicable to humans or animals and derive from carcases that are fit for human consumption in accord II.3.2. in the case of feed for fur animals in addition to II.3.1. consist also of the following animal by-products: (2) either [- animal by-products from poultry and lagomorphs slaughtered on the farm as referred to in Article No 853/2004, which did not show any signs of disease communicable to humans or animals; (²) and/or [- blood of animals which did not show any signs of disease communicable through blood to he from animals other than ruminants that have been slaughtered in a slaughterhouse after have slaughter for human consumption following an ante-mortem inspection in accordance with Uni (2) and/or [- animal by-products arising from the production of products intended for human consumption, greaves and centrifuge or separator sludge from milk processing;] (2) and/or [- products of animal origin, or foodstuffs containing products of animal origin, which are no

consumption for commercial reasons or due to problems of manufacturing or packaging def

aquatic animals, and parts of such animals, except sea mammals, which did not show any signs

(2) and/or [- petfood and feedingstuffs of animal origin, or feedingstuffs containing animal by-products or der longer intended for feeding for commercial reasons or due to problems of manufacturing or 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 from live animals any disease communicable through that product to humans or animals;]

which no risk to public or animal health arises;]

to humans or animals:1

(2) and/or [-

Raw petfood for direct sale or animal by animals

II.a. Certificate reference No

[- animal by-products from aquatic animals originating from plants or establishments manufa consumption;]

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Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNTRY

Health information

(2) and/or

	(2) and/or	[- the following material originating from animals which did not show any signs of disease material to humans or animals:
		(i) shells from shellfish with soft tissue or flesh;
		(ii) the following originating from terrestrial animals:
		 hatchery by-products,
		— eggs,
		 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 species pathogenic to
	(²) and/or	[- animals and parts thereof of the zoological orders of Rodentia and Lagomorpha, except Cate to in Article 8(a)(iii), (iv) and (v) of Regulation (EC) No 1069/2009 and Category 2 material as (g) of that Regulation;]
II.4.		ained and prepared without contact with other material not complying with the conditions laid dow d it has been handled so as to avoid contamination with pathogenic agents;
II.5.	BY-PRODUCT boxes/containe PET FOOD —	cked in final packaging which bear labels indicating "RAW PET FOOD — NOT FOR HUMAN CO S FOR FEED FOR FUR ANIMALS — NOT FOR HUMAN CONSUMPTION" and then in leak ers or in new packaging preventing any leakage and officially sealed boxes/containers which b NOT FOR HUMAN CONSUMPTION" or "ANIMAL BY-PRODUCTS FOR FEED FOR FUR ANIM DN", and the name and the address of the establishment of destination;
II.6.	in the case of	raw petfood:
		n prepared and stored in a plant approved and supervised by the competent authority in accord (EC) No 1069/2009 and
		nined by random sampling of at least five samples from each batch taken during storage (before ong standards (8):
	Salmonella	a: absence in 25 g: n=5, c=0, m=0, M=0
	Enterobac	teriaceae: n=5, c=2, m=10, M=5000 in 1 gram;
II.7.		
	(²) either	[the product does not contain and is not derived from specified risk material as defined in Anne 999/2001 of the European Parliament and of the Council (9) or mechanically separated meat obtation or caprine animals; and the animals from which this product is derived have not been slimeans of gas injected into the cranial cavity or killed by the same method or slaughtered by tessue by means of an elongated rod-shaped instrument introduced into the cranial cavity.]
	(²) or	[the product does not contain and is not derived from bovine, ovine or caprine materials oth animals born, continuously reared and slaughtered in a country or region classified as posing decision in accordance with Article 5(2) of Regulation (EC) No 999/2001;]
II.8.	in addition as	regards TSE:
	(²) either	[in case of animal by-products intended for feeding ruminants and containing milk or milk pr origin, the ovine and caprine animals from which these products are derived have been kept co the last three years on a holding where no official movement restriction is imposed due to a s has satisfied the following requirements for the last three years:
		(i) it has been subject to regular official veterinary checks;
		(ii) no classical scrapie case, as defined in point 2(g) of Annex I to Regulation (EC) No 999/20 following the confirmation of a classical scrapie case:

all animals in which classical scrapie was confirmed have been killed and destroyed, a
 all goats and sheep on the holding have been killed and destroyed, except for breed genotype and breeding ewes carrying at least one ARR allele and no VRQ allele;

(iii) ovine and caprine animals, with the exception of sheep of the ARR/ARR prion genotype, are only if they come from a holding which complies with the requirements set out in points (i)

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNTRY

Raw petfood for direct sale or animal by animals

II.	Health inforr	mation	II.a. Certificate reference No	II.b.
	(²) or	[in case of animal by-products intended for feeding and destined to a Member State listed in the Anne animals from which these products are derived holding where no official movement restriction is irrequirements for the last seven years:	x to Commission Regulation (EC) No save been kept continuously since bir	546/2006 th or for
		(i) it has been subject to regular official veterina	ry checks;	
		(ii) no classical scrapie case, as defined in point following the confirmation of a classical scrap		999/200

- all animals in which classical scrapie was confirmed have been killed and destroyed, a
- all goats and sheep on the holding have been killed and destroyed, except for breed genotype and breeding ewes carrying at least one ARR allele and no VRQ allele;
- (iii) ovine and caprine animals, with the exception of sheep of the ARR/ARR prion genotype, are only if they come from a holding which complies with the requirements set out in points (i)

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 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 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
 case of unloading and reloading, the consignor must inform the BIP of entry into the EU.

- Box reference I.23: for bulk containers, the container number and the seal number (if applicable) should be given.

- Box I.19: use the appropriate Harmonized System (HS) code under the following heading: 05.11.
- 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.
- Box reference I.28:

Nature of commodity: select raw petfood or animal by-product.

In case of raw material for manufacture of raw pet food indicate scientific name of the species.

In case of raw material for manufacture of feed for fur animals select from the following: Aves, Ruminantia, Mam Mollusca, Crustacea, Invertebrata.

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 73, 20.3.2010, p. 1.
- (4) OJ L 226, 23.8.2008, p. 1.
- (5) OJ L 39, 10.2.2009, p. 12.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

cou	JNTRY		Raw petfood for direct sale or ani animals	mal by
II.	Н	ealth information	II.a. Certificate reference No	II.b.
(⁶)	OJ L	340, 31.12.1993, p. 21.		
(⁷)	OJ L	320, 18.11.2006, p. 53.		
(8)	Where	r:		
	n =	number of samples to be tested;		
		threshold value for the number of bacteria; the result is consid m;	ered satisfactory if the number of bacte	eria in a
	M =	maximum value for the number of bacteria; the result is consider more; and	lered unsatisfactory if the number of ba	acteria ir
		number of samples the bacterial count of which may be betwee count of the other samples is m or less.	een m and M, the sample still being co	nsidere
(9)	OJ L	147, 31.5.2001, p. 1.		
(10)	OJ L	94, 1.4.2006, p. 28.		
-	The sig	nature and the stamp must be in a different colour to that of	the printing.	
		or the person responsible for the consignment in the Europ cany the consignment until it reaches the border inspection po		r veteri
Offi	icial vet	erinarian/Official inspector		
	Name (in capital letters):	Qualifica	tion and
	Date:		Signature	e:
	Stamp:			

Chapter 4(A) is replaced by the following: (c)

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

CHAPTER 4(A)

Health certificate

For the import of blood and blood products from equidae to be used outside the feed chain, for dispatch to or for transit through (2) the European Union

cou	NTR	Y			, ,
	l.1.	Consignor		I.2. Certificate reference No	1.2.
		Name Address		I.3. Central competent authority	
		Tol		I de la collection de la contraction de la contr	
ment		Tel.		I.4. Local competent authority	
	1.5.	Consignee		I.6. Person responsible for the loa	d in El
igi		Name Address		Name Address	
S		Postcode		Postcode	
hed		Tel.		Tel.	
Part I: Details of dispatched consignment	1.7.	Country of origin ISO code	I.8. Region of origin Code	I.9. Country of ISO code destination	I.10. F
ls of	111	Place of origin		I.12. Place of destination	
)etai	1.11.	Name	Approval number		tom wa
=		Address	Approval Hambon		roval n
Pa		Name Address	Approval number	Postcode	
		Name Address	Approval number		
	I.13.	Place of loading		I.14. Date of departure	
	l.15.	Means of transport		I.16. Entry BIP in EU	
		Aeroplane Ship	☐ Railway wagon ☐		
		Road vehicle Othe	r 🗆	1.17.	
		Identification Documentation references			
	Ι 1Ω	Description of commodity		I.19. Commodity code	, (HS
	1.10.	Description of commodity		1.19. Commodity Code	(110 0
				1	.20. Qı
	1.21.	Temperature of product		ı	.22. Nu
		Ambient	Chilled	Frozen	
	1.23.	Seal/Container No		I	.24. Ty
	1.25.	Commodities certified for:			
		Technical use			
	1.26.	For transit through EU to third of	country	I.27. For import or admission into El	J
		Third country	ISO code		
	1.28.	Identification of the commodities			
		Species (Scientific name)		Approval number Manufactu	

Blood and blood products from equidae feed chain

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNTRY

Part II: Certification

NTRY			feed	d chain	
II.	Health inform	ation	II.a	. Certificate reference No	II.b.
	and of the Cou	ned official veterinarian, declare that I have read a uncil (^{1a}) and in particular Article 8(c) and (d) and <i>I</i> hapter IV of Annex XIII thereto, and certify that th	Article	e 10 thereof, and Commission Re	egulation
II.1.	consist of bloc	od or blood products from equidae that satisfy the	e hea	alth requirements below;	
II.2.	consist exclus	ively of blood or blood products of equidae not in	ntend	led for human or animal consum	ption;
II.3.	column "third of following disease	tained from animals that originate from the EU Micountries' lists" of row No 3 of Table 2 in Section uses are compulsorily notifiable: African horse sick g Venezuelan equine encephalomyelitis), equine in	1 of ness,	Chapter II of Annex XIV to Reg dourine, glanders (Burkholderia	ulation (E <i>mallei</i>), e
II.4.	accordance w supervised by	rived from blood from equidae, which was collecte ith Regulation (EC) No 853/2004 of the Europe the competent authority of the country of collect of collection for the purpose of collecting blood fr med animals;	an P ion a	'arliament and of the Council (3) and in facilities approved and su), in slaug pervised
II.5.	have been de	rived from blood which was collected from equida	ae:		
II.5.1.	I to Council D	action on the date of blood collection did not show irective 2009/156/EC (*), and of equine influenza, 4 of Article 1.2.3 of the Terrestrial Animal Health	equ	ine piroplasmosis, equine rhinop	neumonit
II.5.2.		een kept for at least 30 days prior to the date of a ect to a prohibition order pursuant to Article 4(5) /156/EC;			
II.5.3.		contact with equidae from a holding which was s	ubjec	ct to a prohibition order for anim	al health
II.5.4.	for which the	period for the prohibition order referred to in point	ts II.6	5.2. and II.5.3 has been determine	ned as fo
	(²) either	[not all the animals of species susceptible to the period of prohibition must be at least:	disea	ase located on the holding have	been slau
		 six months in the case of glanders (Burkholde disease are slaughtered, 	eria r	mallei), beginning on the date on	which th
		 six months in the case of equine encepha beginning on the date on which the equidae 			
		 in the case of equine infectious anaemia, until remaining animals have shown a negative re 			
		- six months from the date of the last recorded	d cas	se of vesicular stomatitis,	
		- one month from the date of the last recorded	d cas	e of rabies,	
		 15 days from the date of the last recorded c 	ase (of anthrax;]	
	(²) or	[all the animals of species susceptible to the dise disinfected, in which case the period of prohibit slaughtered and the premises disinfected, excep	ion n	nust be 30 days, beginning on t	the date of
II.6.		s come from an establishment or plant approved ions set out in Article 23 or 24 of Regulation (EC			ority of the
II.7.	blood products	s have been produced from blood which fulfils the	e cor	nditions referred in II.4 and II.5 a	ind
	(²) either	[has been collected from equidae which have be three months old, prior to the date of collection of during that period and the period of blood collections.]	on ho	oldings under veterinary supervision	
		(a) African horse sickness for two years;			

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNTI	RY			Blood and blood products from e feed chain	quidae
II.	Health inforr	mation		II.a. Certificate reference No	II.b.
		(b) Venezuelan	equine encephalomyelitis for a po	eriod of at least two years;	
		(c) glanders			
		(²) either	[for a period of three years;]		
		(²) or	slaughterhouse referred to in II.4	e the animals have passed the post-mo, including a careful examination of mu and their ramifications, after splitting	icous me
		(d) in the case	of blood products other than seru	um and plasma, vesicular stomatitis for	r six mo
	(²) or	possible causat	ive pathogens for African horse sid	ing treatments, followed by an effectiv ckness, equine encephalomyelitis of all sicular stomatitis and glanders (<i>Burkho</i>	types in
		(2) either	[heat treatment at a temperature	e of 65°C for at least three hours;]	
		(2) and/or	[irradiation at 25 kGy by gamma	ı rays;]	
		(2) and/or	[change in pH to pH 5 for two h	nours;]	
		(2) and/or	[heat treatment of at least 80°C	throughout their substance;]]	
II.8.	all precautions and packagin		n to avoid contamination of the blo	ood and blood products with pathogenic	agents
11.9.		olood products ION" and bearing		neable containers clearly labelled "N	NOT FO
	(a) in the cas	se of blood, the	approval number of the establishn	nent of collection;	
	(b) in the cas	se of blood produ	ucts, the approval number of the	establishment of production;	
II.10.	the products	were stored in e	nclosed storage.		
Notes					
Part I:					
			ble for the consignment in the Eu e certificate is for import commodi	ropean Union: this box is to be filled i ity.	n only if
	reference I.11 nority.	and I.12: Approv	val number: the registration number	er of the establishment or plant, which	has bee
			ation: this box is to be filled in only nouses and custom warehouses.	y if it is a certificate for transit commod	ity. The
			mber (railway wagons or containe ne consignor must inform the BIP	er and lorries), flight number (aircraft) of of entry into the EU.	r name
— Вох	I.19: use the	appropriate Harm	nonized System (HS) code under	the following heading: 30.02.	
— Вох	reference I.23	3: for bulk contain	ners, the container number and the	e seal number (if applicable) must be	included
— Вох	reference I.25	i: technical use: a	any use other than for animal con	sumption.	
— Вох	reference I.26	and I.27: fill in a	according to whether it is a transit	t or an import certificate.	
— Вох	reference I.28	3:			
(a)	Manufacturing	plant:			
	(i) in the case	of blood, provid	e the approval number of the reg	istered establishment of collection;	

(ii) in the case of blood products, provide the approval number of the establishment of production;(b) Species: select amongst the following: Equus cabalus, Equus asinus, Equus cabalus*asinus.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNTR	ıy	Blood and blood products from feed chain	equidae
II.	Health information	II.a. Certificate reference No	II.b.
Part II:			
(^{1a}) OJ	L 300, 14.11.2009, p. 1.		
(1b) OJ	L 54, 26.2.2011, p. 1		
(²) Del	ete as appropriate.		
(3) OJ	L 139, 30.4.2004, p. 55.		
(4) OJ	L 192, 23.7.2010, p. 1.		
— The	signature and the stamp must be in a different colour to that of	the printing.	
	for the person responsible for the consignment in the European consignment until it reaches the border inspection post.	Union: this certificate is only for vete	rinary purp
Official v	veterinarian/Official inspector		
Nam	e (in capital letters):	Qualifi	cation and
Date	:	Signat	ure:
Starr	p:'		

Chapter 4(C) is replaced by the following: (d)

CHAPTER 4(C)

Health certificate

For untreated blood products, excluding of equidae, for the manufacture of derived products for purposes outside the feed chain for farmed animals, intended for dispatch to or for transit through (2) the European Union

cou	JNTR'	Υ				٧
	l.1.	Consignor Name		I.2. Certifica	te reference No	1.2.
		Address		I.3. Central	competent authority	
		Tel.		I.4. Local co	ompetent authority	
Part I: Details of dispatched consignment	1.5.	Consignee Name Address Postcode Tel.		I.6. Person Name Address Postcoo Tel.		ad in EU
of dispate	1.7.	Country of origin ISO code	I.8. Region of origin Code	I.9. Country destinat		I.10. R
ils	1.11.	Place of origin		I.12. Place o	f destination	
l: Deta		Name Address	Approval number	Name Address		stom wa
Part		Name Address	Approval number	Postcoo		,
		Name Address	Approval number			
	I.13.	Place of loading		I.14. Date of	departure	
	l.15.	Means of transport		I.16. Entry B	IP in EU	
		Aeroplane Ship				
		Road vehicle Other [l.17.		
		Documentation references				
	I.18.	Description of commodity			I.19. Commodity co	de (HS c
						I.20. Qu
	I.21.	Temperature of product Ambient □	Chilled		Frozen 🗆	I.22. Nu
	1.23.	Seal/Container No				1.24. Тур
	1.25.	Commodities certified for:			ı	
		Technical use				
	1.26.	For transit through EU to third cou	untry	I.27. For impo	ort or admission into E	U
		Third country	ISO code			
	1.28.	Identification of the commodities				
		Species (Scientific name)	Approval numbe Manufac	er of establishn cturing plant	nents	
	I					

Part II: Certification

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OUNTRY			Untreated blood products, excludi facture of derived products for pu for farmed animals	
II.	Health informa	tion	II.a. Certificate reference No	II.b.
	Parliament ar	igned official veterinarian, declare that I ha nd of the Council (¹ a) and in particular Article , and in particular Annex XIV, Chapter II ther	8(c) and (d) and Article 10 thereof, and	
II.1.	the blood pro	ducts described above consist of blood prod	lucts that satisfy the health requirements b	elow;
II.2.	they consist e	exclusively of blood products not intended for	r human or animal consumption;	
II.3.		en prepared and stored in a plant supervised wing animal by-products:	by the competent authority or in the estab	olishme
	(²) either [-	blood of slaughtered animals, which is fit for for human consumption for commercial reasons.		nion le
	(²) and/or [-	blood of slaughtered animals, which is reject which did not show any signs of diseases of slaughtered in a slaughterhouse and were of accordance with Union legislation;]	communicable to humans or animals, deriv	ed fro
	(²) and/or [-	blood of slaughtered animals, which did not from animals that have been slaughtered ir following an ante-mortem inspection in according	n a slaughterhouse after having been cons	
	(²) and/or [-	blood and blood products derived from the	production of products intended for human	n cons
	(²) and/or [-	blood and blood products originating from lin product to humans or animals;]	ve animals that did not show signs of any di	isease
	(²) and/or [-	animal by-products derived from animals w Directive 96/22/EC or Article 2(b) of Directive		ent as
	(²) and/or [-	animal by-products containing residues of Annex I to Directive 96/23/EC, if such res absence thereof, in national legislation;]		
II.4.	legislation, in	m which such products are manufactured h slaughterhouses approved and supervised l oproved and supervised by the competent au	by the competent authority of the country	
(²) [II.5.		f blood products derived from animals belong the products come:	ging to the taxa Artiodactyla, Perissodactyla	a and I
II.5.1.		y where no case of rinderpest, peste des pe ation has not been carried out against those		en rec
(²) [II.5.2	c. either	[from the third countries, territories or parts tories or parts thereof) (3) where no case vaccination has not been carried out again	of foot-and-mouth disease has been recor	
	or	[from the countries, territories or parts there parts thereof) (*) where no case of foot-and programmes against foot-and-mouth diseas for at least 12 months (*);]]	I-mouth disease has been recorded for 12	month
(²) [II.5.3	. In addition, in	case of animals other than Suidae and Tay	assuidae:	
	(²) either	[in the country or region of origin no ca seropositive animals) has been recorded those diseases for at least 12 months;]		
	(2) or	[in the country or region of origin vesicular	stomatitis and bluetongue (2) seropositive	anima
(²) [II.5.4	. In addition, in	case of Suidae and Tayassuidae:		
II.5.4.1.		or region of origin no case of swine vesicular onths and vaccination has not been carried of		

COUNTRY			Untreated blood products, exclu facture of derived products for professionals							
II. Hea	lth information	on	II.a. Certificate reference No	II.b.						
(²) [II.5.4.2.	either	[in the country or region of origin no case of been recorded for 12 months and in which months;]								
(²) [II.5.4.2.	or	[in the country or region of origin vesicular s	stomatitis seropositive animals are pre	sent (4);						
(²) [II.6.	in the case of blood products derived from poultry or other avian species the animals and the products of country or region with code $^{(5)}$									
	which has been free from Newcastle disease and highly pathogenic avian influenza as defined in the Ter of the OIE,									
	which for at least 12 months has not carried out vaccination against avian influenza,									
	where the animals from which the products derive have not been vaccinated against to a Newcastle disease master strain showing a higher pathogenicity than lentogenic to									
II.7.	II.7. the products were:									
	(²) either	[packed in new or sterilised bags or bottles,	1							
	(²) or	[transported in bulk in containers or other in disinfectant approved by the competent authors]		hly clea						
	the outer packaging or containers bear labels indicating "NOT FOR HUMAN OR ANIMAL CONSUMPT									
II.8.	the products were stored in enclosed storage;									
II.9.	all precautions were taken to avoid contamination of the products with pathogenic agents during trans									
II.10.										
	(²) either	[the product does not contain and is not deri No 999/2001 of the European Parliament and bovine, ovine or caprine animals; and the ar stunning by means of gas injected into the of central nervous tissue by means of an elong	I of the Council (6) or mechanically sep nimals from which the product is deriver anial cavity or killed by the same met	arated ned have thod or						

Notes

Part I:

(2) or

Box reference I.6: Person responsible for the consignment in the European Union: this box is to be filled in only if commodity; it may be filled in if the certificate is for import commodity.

[the product does not contain and is not derived from bovine, ovine or caprine materials ot animals born, continuously reared and slaughtered in a country or region classified as posin decision in accordance with Article 5(2) of Regulation (EC) No 999/2001.]

- Box reference I.11 and I.12: Approval number: the registration number of the establishment or plant, which has been
- Box reference I.12: Place of destination: this box is to be filled in only if it is a certificate for transit commodity. The 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 case of unloading and reloading, the consignor must inform the border inspection post of entry into the EU.
- Box I.19: use the appropriate Harmonized System (HS) code under the following headings: 30.02 or 35.02.
- Box reference I.23: for bulk containers, the container number and the seal number (if applicable) should be include
- 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.
 - Box reference I.28 Species: select from the following: Aves, Bovidae, Suidae, Otra Mammalia, Pesca, Reptilia.

Untreated blood products, excluding of

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Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

facture of derived products for purpose COUNTRY for farmed animals II. Health information II.a. Certificate reference No 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) Code of the territory as it appears in Part 1 of Annex II to Regulation (EU) No 206/2010. In this case following the border check provided for in Directive 97/78/EC, and in accordance with the conditions laid Directive, the products must be transported directly to the plant of destination. (5) Code of the territory as it appears in Part 1 of Annex I to Regulation (EC) No 798/2008. (6) OJ L 147, 31.5.2001, p. 1. - 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 purporting the consignment until it reaches the border inspection post. Official veterinarian/Official inspector Name (in capital letters): Qualification and Signature: Stamp:

(e) Chapter 4(D) is replaced by the following:

CHAPTER 4(D)

Health certificate

For treated blood products, excluding of equidae, for the manufacture of derived products for purposes outside the feed chain for farmed animals, intended for dispatch to or for transit through (2) the European Union

cou	INTR	1					٧			
	l.1.	Consignor Name	1.2.	Certifica	ite refer	ence No	1.2.			
		Address	I.3. Central competent authority							
Part I: Details of dispatched consignment		Tel.	I.4. Local competent authority							
	1.5.	Consignee Name Address Postcode Tel.	I.6. Person responsible for the load in Et Name Address Postcode Tel.							
of dispatc	1.7.	Country of origin ISO code I.8. Region of origin Code	1.9.	Country destinat		ISO code	I.10. R			
Details o	l.11.	Place of origin Name Approval number	I.12.	Place of	f destina		ustom wa			
Part I:		Address Name Approval number Address		Address		A	oproval nu			
		Name Approval number Address								
	I.13.	Place of loading	l.14.	Date of	departu	ire				
	I.15.	Means of transport	I.16.	Entry BI	P in El	J				
		Aeroplane Ship Railway wagon Road vehicle Other Other								
		Identification Documentation references	1.17.							
	I.18.	Description of commodity			I.19. C	Commodity co	de (HS c			
				,			I.20. Qu			
	I.21.	Temperature of product Ambient ☐ Chilled ☐			Frozen		1.22. Nu			
	1.23.	Seal/Container No					1.24. Тур			
	1.25.	Commodities certified for:								
		Technical use								
	1.26.	For transit through EU to third country Third country ISO code	1.27.	For impo	rt or ad	Imission into	EU			
	1.28.	28. Identification of the commodities								
		Species Approval number of establishments (Scientific name) Manufacturing plant								

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

Treated blood products, excluding of equ of derived products for purposes outside animals

cou	INTRY			of derived products for purposes animals	outside					
	II.	Health infor	mation	II.a. Certificate reference No	II.b.					
		and of the C	igned official veterinarian, declare that I have read a ouncil (^{1a}) and in particular Article 8(c) and (d) and Annex XIV, Chapter II thereof, and certify that:							
cation	II.1.	the blood pr	oducts described above consist of blood products	that satisfy the requirements below;						
Certific	II.2.	they consist exclusively of blood products not intended for human or animal consumption;								
Part II: Certification	II.3.	they have be	been prepared and stored in a plant supervised by the competent authority, exclusively with the							
		(²) either	[- blood of slaughtered animals, which is fit for his for human consumption for commercial reason		Union le					
		(²) and/or	[- blood of slaughtered animals, which is rejecter which did not show any signs of diseases con slaughtered in a slaughterhouse and were con accordance with Union legislation;]	mmunicable to humans or animals, der	rived fro					
		how any signs of diseases communica a slaughterhouse after having been col dance with Union legislation;]								
		(2) and/or [- blood and blood products originating from live animals that did not show clinical s through these products to humans or animals;]								
		(²) and/or	[- animal by-products which have been derived a Article 1(2)(d) of Directive 96/22/EC or Article		ed to ille					
		(²) and/or	[- animal by-products containing residues of ot Annex I to Directive 96/23/EC, if such residu absence thereof, in national legislation;]							
	II.4.	legislation, in	om which such products are manufactured has a slaughterhouses approved and supervised by the roved and supervised by the competent authority of	e competent authority of the country of collect						
	(²) [II.5.	Tayassuidae	lood products derived from Artiodactyla, Perissoda , the products have undergone one of the followin icular stomatitis, rinderpest, peste des petits rumin	ng treatments, guaranteeing the absen	ice of pa					
		(²) either	[heat treatment at a temperature of 65 °C for at	least three hours, followed by an effect	tiveness					
		(²) and/or	[irradiation at 25 kGy by gamma rays, followed by	by an effectiveness check;]						
		(²) and/or	[change in pH to pH 5 for two hours, followed b	y an effectiveness check;]						
		(²) and/or	[heat treatment of at least 80 °C throughout their	substance, followed by an effectivene	ess chec					
	(²) [II.6.	following treat	of blood products derived from Suidae, Tayassuidae atments guaranteeing the absence of pathogens of ease, classical swine fever, African swine fever, No	the following diseases: foot-and-mouth	disease					
		(²) either	[heat treatment at a temperature of 65 °C for at	least three hours, followed by an effect	tiveness					
		(2) and/or	[irradiation at 25 kGy by gamma rays, followed by	by an effectiveness check;]						
		(²) and/or	[heat treatment of at least 80 °C for Suidae/Ta throughout their substance, followed by an effect		poultry a					

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNTRY

(1b) OJ L 54, 26,2,2011, p. 1.

Treated blood products, excluding of equ of derived products for purposes outside animals

II. Health information II.a. Certificate reference No (2) [II.7 In the case of blood products derived from species other than listed in points II.5 or II.6 the protection the following treatment (please specify): II.8. The products were: (2) either [packed in new or sterilised bags or bottles;] (2) or [transported in bulk in containers or other means of transport that were thoroughly cleaned and di approved by the competent authority before use;] and the outer packaging or containers bear labels indicating "NOT FOR HUMAN OR ANIMAL CONSUMPTION"; II.9. the products were stored in enclosed storage; II.10. all precautions were taken to avoid contamination of the products with pathogenic agents after treatment: II.11. (2) either If the product does not contain and is not derived from specified risk material as defined in Anne [the product does not contain and is not derived from specified risk material as defined in Anni 999/2001 of the European Parliament and of the Council (3) or mechanically separated meat obtation or caprine animals; and the animals from which the product is derived have not been slimeans of gas injected into the cranial cavity or killed by the same method or slaughtered by latissue by means of an elongated rod-shaped instrument introduced into the cranial cavity.] [the product does not contain and is not derived from bovine, ovine or caprine materials oth animals born, continuously reared and slaughtered in a country or region classified as posing decision in accordance with Article 5(2) of Regulation (EC) No 999/2001.] (2) or 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 commodity; it may be filled in if the certificate is for import commodity. Box reference I.11 and I.12: Approval number: the registration number of the establishment or plant, which has been Box reference I.12: Place of destination: this box is to be filled in only if it is a certificate for transit commodity. The 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 In case of unloading and reloading, the consignor must inform the BIP of entry into the EU. Box I.19: use the appropriate Harmonized System (HS) code under the following headings: 05.11, 30.02 or 35.02 Box reference I.23: for bulk containers, the container number and the seal number (if applicable) should be included. 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. Box reference I.28 in case of Species: select from the following: Aves, Bovidae, Suidae, Otra Mammalia, Pesca, I (1a) OJ L 300, 14.11.2009, p. 1.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

CO	UNTRY	Treated blood products, excluding of eq of derived products for purposes outside animals								
II.	Health information	II.a. Certificate reference No	II.b.							
(²)	Delete as appropriate.									
(3)	OJ L 147, 31.5.2001, p. 1.									
_	— 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 purp the consignment until it reaches the border inspection post. 										
Of	ficial veterinarian/Official inspector									
	Name (in capital letters):	Qual	ification and							
	Date:	Sign	ature:							
	Stamp:'									

(f) Chapter 6(A) is replaced by the following:

CHAPTER 6(A)

Health certificate

For treated game trophies and other preparations of birds and ungulates, consisting only of bones, horns, hooves, claws, antlers, teeth, hides or skins, for dispatch to or for transit through (2) the European Union

;OU	INTR	Y											١	
	l.1.	Consignor						1.2.	Certifica	ite refe	rence No		1.2.	
		Name Address						1.3.	I.3. Central competent authority					
		Tel.							I.4. Local competent authority					
dispatched consignment	1.5.	Consignee Name Address Postcode Tel.		1.6.	I.6. Person responsible for the load in E Name Address Postcode Tel.									
of dispatcl	1.7.	Country of origin	ISO code	1.8.	Region of ori	gin	Code	1.9.	Country		ISO code	1.10	0. F	
ils	1.11.	Place of origin						I.12.	Place o	f destin	nation			
Part I: Details of		Name Address				Approval number			Name Address	Custom wa				
Part		Name Approval number Address					nber		Postcoo					
		Name Approval number Address												
	l.13.	3. Place of loading							I.14. Date of departure					
	I.15.	5. Means of transport						I.16. Entry BIP in EU						
		Aeroplane												
		Identification							I.17. Number(s) of CITES					
	140	Documentation references								140	O	-1- 4	10	
	1.18.	3. Description of commodity								1.19. (Commodity co	oae (F	15 0	
												1.20.	Qu	
	I.21.	1.										1.22.	Nu	
	1.23.	Seal/Container No										1.24.	Ту	
	1.25.	25. Commodities certified for:												
		Technical use												
	1.26.	6. For transit through EU to third country						1.27.	For impo	ort or a	dmission into	EU		
		Third country			ISO code									
	1.28.	Identification of the	commodities											
		Species (Scientific name)			Nature of commodity									

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

(1a) OJ L 300, 14.11.2009, p. 1.

II.

Certification

Part II:

Treated game trophies and other prepara lates, consisting only bones, horns, hoo COUNTRY hides or skins Health information II.a. Certificate reference No I, the undersigned official veterinarian, declare that I have read and understood Regulation (European Parliament and of the Council (1a) and Commission Regulation (EU) No 142/2011 (1 XIV, Chapter II thereof, and certify that the game trophies described above: have been packaged, immediately after treatment, without being in contact with other product contaminate them, in individual, transparent and closed packages so as to avoid any subsequent (2) either [II.2.1] in the case of game trophies or other preparations consisting only of hides or skin: (2) either [have been dried;] (2) and/or [have been dry-salted or wet-salted for a minimum of 14 days before dispatch;] (2) and/or minimum of 14 days salting before they reach the EU border inspection post;]] (2) and/or [II.2.2 in the case of game trophies or other preparations consisting only of bone, horns, hooves, claws (a) have been immersed in boiling water for an appropriate time so as to ensure that any mat hooves, claws, antlers or teeth is removed, and (b) have been disinfected with a product authorised by the competent authority, in particular with parts consisting of bone are concerned.] 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 commodity; it may be filled in if the certificate is for import commodity. Box reference I.11 and I.12: Approval number: the registration number of the establishment or plant, which has been Box reference I.12: Place of destination: this box is to be filled in only if it is a certificate for transit commodity. The 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 case of unloading and reloading, the consignor must inform the BIP of entry into the EU. Box I.19: use the appropriate Harmonized System (HS) code under the following headings: 05.05, 05.06, 05.07 or - Box reference I.23: for bulk containers, the container number and the seal number (if applicable) should be include 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. Box reference I.28: (a) for nature of commodity, select one or more of the following: [bones], [horns], [hooves], [claws], [antlers], [te (b) in case of Species: select from the following: Aves, Equidae, Tapiridae, Rhinoceritidae, Antilocaparidae, Bov Giraffidae, Hippopotamindae, Moschidae Suidae, Tayassuidae, Tragulidae and Elephantidae. Part II:

Treated game trophies and other prepar

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNTRY	hides or skins	norns, noc
II. Health information	II.a. Certificate reference No	II.b.
(^{1b}) OJ L 54, 26.2.2011, p. 1		
(²) Delete as appropriate.		
- The signature and the stamp must be in a different colour to the	at of the printing.	
 Note for the person responsible for the consignment in the Europe the consignment until it reaches the border inspection post. 	ean Union: this certificate is only for vete	rinary purp
Official veterinarian/Official inspector		
Name (in capital letters):	Qualit	fication and
Date:	Signa	iture:
Stamp:'		

(g) Chapter 8 is replaced by the following:

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

οu	INTR	1										٧
	l.1.	Consignor Name					1.2.	Certificat	te refer	ence No		1.2.
		Address					1.3.	Central	compet	ent authority		
		Tel.					1.4.	Local co	mpeter	nt authority		
Part I: Details of dispatched consignment	I.5.	Consignee Name Address					I.6.	Person r Name Address	espons	sible for the lo	oad in	EU
cuea col		Postcode Tel.						Postcode Tel.	В			
or dispar	1.7.	Country of origin	ISO code		egion of igin	Code	1.9.	Country destinati		ISO code	1.10	0. R d
2	l.11.	Place of origin					1.12.	Place of	destin	ation		
: Det		Name Address			Approval nu	mber		Name Address			ustom	
Fan		Name Address			Approval nu	mber		Postcode		,,,	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	
		Name Address			Approval nu	mber						
	I.13.	Place of loading					l.14.	Date of	departu	ıre		
	I.15.	Means of transport	:				I.16.	Entry BII	P in El	J		
		Aeroplane	Ship		Railway	wagon 🗌						
		Road vehicle Identification	Other				1.17.					
		Documentation refe	erences									
	I.18.	Description of com	modity						I.19. C	Commodity co	de (H	IS c
											1.20.	Qu
	I.21.	Temperature of pro	oduct		Chilled				Frozer		1.22.	Nu
	1.23.	Seal/Container No									1.24.	Тур
	1.25.	Commodities certifi	ied for:									
		Technical use										
	1.26.	For transit through	EU to third o				1.27.	For impor	t or ac	Imission into	EU	
		Third country		ISO	code							
	1.28.	Identification of the	commodities									
		Species (Scientific name)	Nature commo		Approv	al number of Manufacturin				Number of packages		N

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

cou	NTRY				Animal by-products to be used for or for trade samples (2)	r purpo
	II.	Health informa	tion		II.a. Certificate reference No	II.b.
		Parliamen	t and	ned official veterinarian, declare that I hav of the Council (^{1a}) and Commission Regula the animal by-products described above:		
cation	(²) II.1.			es which consist of animal by-products inte julation (EU) No 142/2011, that are bearing		
Part II: Certification	(²) II.2.	satisfy the	anim	nal health requirements below;		
art	II.2.1.	have been	n			
		(²) either	[(a)	obtained from materials imported from the to export fresh meat of the species to the		
		(²) and/or	[(b)	obtained in the exporting country, territor	y or part thereof:	
Ч				either		
				(i) That have remained in this territory of birth or for at least the last three mo		neat of t
				(ii) Killed in the wild in this territory (4);]		
		(²) and/or	[(c)	are derived from eggs, milk, rodents, lag	gomorphs, or aquatic animals or terres	trial or a
	II.2.2.			of materials other than derived from egga ave been obtained from animals:	s, milk, rodents, lagomorphs, or aqua	itic anim
		(²) either	[(a)	coming from holdings:		
					Newcastle disease or highly pathogenic e fever during the prior 40 days; nor in	c avian i
				(ii) where there has been neither case/holdings situated in their vicinity with	outbreak of foot-and-mouth disease d in 25 km, during the prior 30 days; ar	
			(b)	which:		
				(i) were not killed to eradicate any epiz	ootic disease;	
				 (ii) have remained in their holdings of originate directly to the slaughterhouse without conditions; 	gin for at least 40 days before departu ut contact with other animals which di	
				(iii) at the slaughterhouse, have passed to and have shown no evidence of the	he ante-mortem health inspection during diseases referred to above for which	
				(iv) have been treated in the slaughterhorelevant provisions of Council Directive killing;]	ouse before and at the time of slaugh ve 93/119/EC) (5) on the protection of	
		(²) or	[(a)	captured and killed in the wild in an area	a:	
					n no case/outbreak of any of the follow ase, rinderpest, Newcastle disease o cical or African swine fever during the	r highly
				(ii) that is situated at a distance that exc part thereof, which is not authorised	ceeds 20 km from the borders separal d at these dates for exporting this ma	
			(b)	which after killing were transported with afterwards to a game establishment, or or		collecti

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNT	RY		Animal by-products to be used for or for trade samples (2)	r purpo
II.	Health information		II.a. Certificate reference No	II.b.
II.2.3.	lishment arour animals are so to the Europea	e of materials other than materials derived nd which, within a radius of 10 km, there has usceptible during the prior 30 days or, in the an Union has been authorised only after remotrol of an official veterinarian;	been no case/outbreak of diseases re- event of a case of disease, the prepar	ferred to ation of
II.2.4.		tained and prepared without contact with ot so as to avoid contamination with pathoge		nditions
II.2.5.	and, in the ca authority, bear	cked in new packaging preventing any leak se of consignments shipped other than via p ring the label indicating "ANIMAL BY-PRODI DE THE FEED CHAIN" and the name and	parcel post, in containers sealed under UCTS ONLY FOR THE MANUFACTUR	the res
II.2.6.	consist only o	f the following animal by-products:		
	(²) either [-	carcases and parts of animals slaughtered for human consumption in accordance w commercial reasons;]		
	(²) and/or [-	carcases and the following parts originating were considered fit for slaughter for hum following parts of animals from game killed	an consumption following an ante-mo	rtem in:
		(i) carcases or bodies and parts of anima Union legislation, but which did not sh	als which are rejected as unfit for humanow any signs of disease communicab	
		(ii) heads of poultry;		
		(iii) hides and skins, including trimmings are and metacarpus bones, tarsus and tarsus bones, tarsus and tarsus bones, tarsus and tarsus and tarsus bones, tarsus and tarsus bones, tarsus and tarsus bones, tarsus and tarsus bones, tar		uding th
		(iv) pig bristles;		
		(v) feathers;]		
	(²) and/or [-	animal by-products from poultry and lagome (EC) No 853/2004, which did not show an		
	(²) and/or [-	blood of animals which did not show any obtained from animals other than ruminal considered fit for slaughter for human conlegislation;]	nts that have been slaughtered in a	slaught
	(²) and/or [-	animal by-products arising from the produ bone, greaves and centrifuge or separator		consur
	(²) and/or [-	products of animal origin, or foodstuffs cor consumption for commercial reasons or du which no risk to public or animal health ar	e to problems of manufacturing or pack	
	(²) and/or [-	petfood and feedingstuffs of animal origin, are no longer intended for feeding for common other defects from which no risk to pub.	mercial reasons or due to problems of	
	(²) and/or [-	blood, placenta, wool, feathers, hair, hornsigns of any disease communicable through		
	(²) and/or [-	aquatic animals, and parts of such anim communicable to humans or animals;]	als, except sea mammals, which did	not sh

(²) and/or [- animal by-products from aquatic animals originating from establishments or plants manufaconsumption;]

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNTRY

II.

Health information

Animal by-products to be used for purpo or for trade samples (2)

II.b.

II.a. Certificate reference No

	(²) and/or	[-	the following material originating from animals which did not show any signs of disease material to humans or animals:
			(i) shells from shellfish with soft tissue or flesh;
			(ii) the following originating from terrestrial animals:
			 hatchery by-products,
			— eggs,
			 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 species pathoge
	(²) and/or	[-	animals and parts thereof of the zoological orders of Rodentia and Lagomorpha, excereferred to in Article 8(a)(iii), (iv) and (v) of Regulation (EC) No 1069/2009 and Categor in Article 9(a) to (g) of that Regulation;]
	(²) and/or	[-	fur originating from dead animals that did not show clinical signs of any disease communic humans or animals;]
II.2.7.			ep-frozen at the plant of origin or have been preserved in accordance with EU legislation i een dispatch and delivery to the plant of destination.
(²) (⁶) [II.2.8.	Specific I	requ	irements
(²) (⁷) II.2.8.1.			ots in this consignment come from animals that have been obtained in the territory ment ogrammes against foot-and-mouth disease are being regularly carried out and officially co
(²) (8) II.2.8.2.	The by-pr	oduc	ots in this consignment consist of animal by-products derived from offal or deboned meat.
II.2.9.			
	(²) either	999 bov stur	e product does not contain and is not derived from specified risk material as defined in Ann ### product of the European Parliament and of the Council (*) or mechanically separated me rine, ovine or caprine animals; and the animals from which this product is derived have nning by means of gas injected into the cranial cavity or killed by the same method or s tral nervous tissue by means of an elongated rod-shaped instrument introduced into the
	(²) either (²) or	999 bov stur cen [the	N/2001 of the European Parliament and of the Council (9) or mechanically separated me rine, ovine or caprine animals; and the animals from which this product is derived have nning by means of gas injected into the cranial cavity or killed by the same method or s
II.2.10.	(²) or	999 bov stur cen [the anir dec	M2001 of the European Parliament and of the Council (9) or mechanically separated me rine, ovine or caprine animals; and the animals from which this product is derived have ning by means of gas injected into the cranial cavity or killed by the same method or so strain nervous tissue by means of an elongated rod-shaped instrument introduced into the perproduct does not contain and is not derived from bovine, ovine or caprine materials oth mals born, continuously reared and slaughtered in a country or region classified as posing
II.2.10.	(²) or	999 bov stur cen [the anir dec n as [in orig the	wine, ovine or caprine animals; and the animals from which this product is derived have ning by owner or caprine animals; and the animals from which this product is derived have ning by means of gas injected into the cranial cavity or killed by the same method or starl nervous tissue by means of an elongated rod-shaped instrument introduced into the product does not contain and is not derived from bovine, ovine or caprine materials oth mals born, continuously reared and slaughtered in a country or region classified as posing ision in accordance with Article 5(2) of Regulation (EC) No 999/2001;]
II.2.10.	(²) or in addition	ges bov stur cen [the anir dec orig the has	pi2001 of the European Parliament and of the Council (9) or mechanically separated me vine, ovine or caprine animals; and the animals from which this product is derived have entral nervous tissue by means of an elongated rod-shaped instrument introduced into the product does not contain and is not derived from bovine, ovine or caprine materials oth mals born, continuously reared and slaughtered in a country or region classified as posing ision in accordance with Article 5(2) of Regulation (EC) No 999/2001;] regards TSE: case of animal by-products intended for feeding ruminants and containing milk or milk pr jin, the ovine and caprine animals from which these products are derived have been kept or last three years on a holding where no official movement restriction is imposed due to a s
II.2.10.	(²) or in addition	good sturr cen [the anim dec or as [in origothe has (i)]	wine, ovine or caprine animals; and the animals from which this product is derived have ning by means of gas injected into the cranial cavity or killed by the same method or starl nervous tissue by means of an elongated rod-shaped instrument introduced into the product does not contain and is not derived from bovine, ovine or caprine materials oth mals born, continuously reared and slaughtered in a country or region classified as posing cision in accordance with Article 5(2) of Regulation (EC) No 999/2001;] regards TSE: case of animal by-products intended for feeding ruminants and containing milk or milk prin, the ovine and caprine animals from which these products are derived have been kept collast three years on a holding where no official movement restriction is imposed due to a se satisfied the following requirements for the last three years:

 all goats and sheep on the holding have been killed and destroyed, except for breed genotype and breeding ewes carrying at least one ARR allele and no VRQ allele;

(iii) ovine and caprine animals, with the exception of sheep of the ARR/ARR prion genoty holding only if they come from a holding which complies with the requirements set out i

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

> Animal by-products to be used for purpo COUNTRY or for trade samples (2) Health information II.a. Certificate reference No II. II.b. [in case of animal by-products intended for feeding ruminants and containing milk or milk p origin, and destined to a Member State listed in the Annex to Commission Regulation (EC) and caprine animals from which these products are derived have been kept continuously since years on a holding where no official movement restriction is imposed due to a suspicion of T the following requirements for the last seven years: (i) it has been subject to regular official veterinary checks; (ii) no classical scrapie case, as defined in point 2(g) of Annex I to Regulation (EC) No 999/20 following the confirmation of a classical scrapie case: all animals in which classical scrapie was confirmed have been killed and destroyed, all goats and sheep on the holding have been killed and destroyed, except for breed genotype and breeding ewes carrying at least one ARR allele and no VRQ allele; (iii) ovine and caprine animals, with the exception of sheep of the ARR/ARR prion genoty holding only if they come from a holding which complies with the requirements set out in 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 commodity; it may be filled in if the certificate is for import commodity.
- Box reference I.11: In case of consignments for the particular technological studies or analyses: indicate name an
- Box reference I.11 and I.12: Approval number: the registration number of the establishment or plant, which has been 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 trans 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
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name case of unloading and reloading, the consignor must inform the BIP of entry into the EU.
- Box reference I.19: use the appropriate Harmonized System (HS) code under the following headings: 05.11.91; 05
- Box reference I.23: for bulk containers, the container number and the seal number (if applicable) should be include
- Box reference I.25: technical use: any use other than for animal consumption.
- Box reference I.25: 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
- Box reference I.28:
- products for the manufacture of derived products for uses outside the feed chain: Manufacturing plant: provide th of the approved establishment
- products for the particular technological studies or analyses: the EU plant indicated in authorisation of competent
- Species: select from the following: Aves, Ruminantia, Mammalia Ruminantia, Pesca, Mollusca, Crustacea, Inv

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNTRY

ou	INTRY	Animal by-products to be used for for trade samples (2)	or purpo
II.	Health information	II.a. Certificate reference No	II.b.
Par	t II:		
(^{1a})	OJ L 300, 14.11.2009, p. 1.		
(^{1b})	OJ L 54, 26.2.2011, p. 1.		
(²)	Delete as appropriate.		
(³)	The name and ISO code number of the exporting country as laid de	own in:	
	- Part 1 of Annex II to Regulation (EU) No 206/2010,		
	- the Annex to Regulation (EC) No 798/2008, and		
	— the Annex to Regulation (EC) No 119/2009.		
	In addition the ISO code of territories and parts thereof referred susceptible species concerned) should be included.	to in Regulations mentioned in this	footnote
(⁴)	Only for countries from where game meat intended for human consulturopean Union.	umption of the same animal species	is authori
(⁵)	OJ L 340, 31.12.1993, p. 21.		
(⁶)	Supplementary guarantees to be provided when the material of don African country or part thereof from where only maturated and debon for exportation to the European Union. The whole masseter muscles of I, Part B(1) of Regulation (EC) No 854/2004 of the European Parliar	ed fresh meat of domestic ruminants of bovine animals, incised in accordar	for humance with A
(⁷)	Only for certain South American countries.		
(⁸)	Only for certain South American and South African countries.		
(⁹)	OJ L 147, 31.5.2001, p. 1.		
(¹⁰)	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 U the consignment until it reaches the border inspection post.	Inion: this certificate is only for vetering	nary purpo
Offi	cial veterinarian/Official inspector		
	Name (in capital letters):	Qualific	ation and
	Date:	Signatu	re:
	Stamp:'		

(h) Chapter 10(B) is replaced by the following:

CHAPTER 10(B)

Health certificate

For rendered fats not intended for human consumption to be used for certain purposes outside the feed chain, intended for dispatch to or for transit through (2) the European Union

ou	NTR	1			١
	l.1.	Consignor Name	I.2. Certific	cate reference No	1.2.
		Address	I.3. Centra	al competent authority	
		Tel.	I.4. Local	competent authority	
Part I: Details of dispatched consignment	1.5.	Consignee Name Address Postcode Tel.	I.6. Person Name Addre Postco Tel.	ss	oad in EU
f dispatcl	1.7.	Country of origin ISO code I.8. Region of origin Code	I.9. Count destin		I.10. F
ils o	l.11.	Place of origin	I.12. Place	of destination	
l: Deta		Name Approval number Address	Name Addre		Custom wa
Part		Name Approval number Address	Postco		
		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 O	1.17.		
		Identification Documentation references			
	I 18	Description of commodity		I.19. Commodity o	ode (HS c
	1.10.	Description of commonly		1.19. Commodity C	
					1.20. Qu
	I.21.	Temperature of product Ambient ☐ Chilled ☐		Frozen	1.22. Nu
	1.23.	Seal/Container No			I.24. Ty
	1.25.	Commodities certified for:			
		Technical use ☐			
	1.26.	For transit through EU to third country	I.27. For imp	port or admission into	EU
		Third country ISO code			
	1.28.	Identification of the commodities			
		Species Approval number of establishments Ni (Scientific name) Manufacturing plant	umber of pack	ages Net v	weight

Rendered fats not intended for human

c	ou	NTRY				purposes outside the feed chain	numan
		II.	Health infor	mat	ion	II.a. Certificate reference No	II.b.
			and of the Co	oun	d official veterinarian, declare that I have read a cil (^{1a}) and in particular Articles 8, 9 and 10 the of, and certify that the rendered fats described a	reof, and Regulation (EU) No 142/201	
l	<u>5</u>	II.1.	consist of rer	nde	red fats not intended for human consumption th	nat satisfy the health requirements bel	ow;
	rtificat	II.2.	have been p	repa	ared exclusively with the following animal by-pro	oducts:	
	Part II: Certification	II.2.1.			aterials destined for the production of biodiesel ation (EC) No 1069/2009;	or oleochemical products, animal by-	products
	<u>.</u>	II.2.2.			aterials destined for the production of renewab No 142/2011, animal by-products referred to in		
		II.2.3.	in the case of	f m	aterials destined for purposes other than cosmo	etics, pharmaceuticals or medical dev	ices:
L	\dashv		(²) either	[-	animal by-products containing residues of author to in Article 15(3) of Directive 96/23/EC;]	orised substances or contaminants exc	eeding t
			(²) and/or	[-	products of animal origin which have been declithose products;]	ared unfit for human consumption due	to the pr
			(²) and/or	[-	animals and parts of animals, other than those other than being slaughtered or killed for hum		
			(²) and/or	[-	carcases and parts of animals slaughtered or, i human consumption in accordance with Union reasons;]		
			(²) and/or	[-	carcases and the following parts originating eith considered fit for slaughter for human consump of animals from game killed for human consum	tion following an ante-mortem inspection	on or bo
					(i) carcases or bodies and parts of animals who legislation, but which did not show any significant control of the control of		
					(ii) heads of poultry;		
					(iii) hides and skins, including trimmings and sp metacarpus bones, tarsus and metatarsus		g the pha
					(iv) pig bristles;		
					(v) feathers;]		
			(²) and/or	[-	blood of animals which did not show any signs from animals other than ruminants that have b slaughter for human consumption following an	een slaughtered in a slaughterhouse	after hav
			(²) and/or	[-	animal by-products arising from the production greaves and centrifuge or separator sludge fro		umption,
			(²) and/or	[-	products of animal origin, or foodstuffs contain consumption for commercial reasons or due to which no risk to public or animal health arises	o problems of manufacturing or packa	
			(²) and/or	[-	petfood and feeding stuffs of animal origin, or fe no longer intended for feeding for commercial re defects from which no risk to public or animal	easons or due to problems of manufac	
			(²) and/or	[-	blood, placenta, wool, feathers, hair, horns, hoo any disease communicable through that produ		e animal
			(²) and/or	[-	aquatic animals, and parts of such animals communicable to humans or animals;]	s, except sea mammals, which did	not sho
			(²) and/or	[-	animal by-products from aquatic animals origi consumption;]	nating from plants or establishments	manufac

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNTRY

COUNT	RY				fats not inte		human
II.	Health inform	mat	ion	II.a. Certific	cate reference	No	II.b.
	(²) and/or	[-	the following material originating from animals material to humans or animals:	which did	not show any	signs of d	isease o
			(i) shells from shellfish with soft tissue or fles	h;			
			(ii) the following originating from terrestrial ani	mals:			
			 hatchery by-products, 				
			— eggs,				
			 egg by-products, including egg shells; 				
			(iii) day-old chicks killed for commercial reason	ns;]			
	(²) and/or	[-	aquatic and terrestrial invertebrates other than	species path	hogenic to hur	mans or ani	mals;]
	(²) and/or	[-	animals and parts thereof of the zoological ordeto in Article 8(a)(iii), (iv) and (v) of Regulation (f. (g) of that Regulation;]				
	(²) and/or	[-	hides and skins, hooves, feathers, wool, horns, disease communicable through that product to			m dead ani	mals tha
	(²) and/or	[-	adipose tissue from animals which did not show animals, which were slaughtered in a slaughter following an ante-mortem inspection in accordant	nouse and w	hich were con:	sidered fit fo	
II.2.4.			aterials destined for purposes other than the pro- es or renewable fuels referred to in point J of				
	(²) either	[-	specified risk material as defined in Article 3(1)(Council $(^3)$;]	g) of Regula	tion (EC) No 9	99/2001 of	the Euro
	(²) and/or	[-	entire bodies or parts of dead animals containi No 999/2001 at the time of disposal;]	ng specified	risk material a	as defined i	n Article
	(²) and/or	[-	animal by-products which have been derived fr Article 1(2)(d) of Directive 96/22/EC or Article 1				ed to ille
	(²) and/or	[-	animal by-products containing residues of oth Annex I to Directive 96/23/EC, if such residue absence thereof, by legislation of the Member	s exceed th	e permitted le		
II.3.	the rendered	fats	s:				
			ubjected to processing in accordance with method /2011, in order to kill pathogenic agents,	nod	as laid	down in Ch	apter III
			marked before shipment to the European Uni n of at least 250 mg GTH per kilogram fat is ac		ceroltriheptano	ate (GTH),	so that
	(c) in the cas	se o	of rendered fats of ruminant origin, insoluble imp	ourities in ex	cess of 0.15%	6 in weight	have be
	(d) have been	n tr	ansported under conditions which prevent their	contamination	on, and		

(e) bear labels on the packaging or container indicating "NOT FOR HUMAN OR ANIMAL CONSUMPTION";

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

Signature:

COUNT	'RY	▶ [™]	Rendered fats not intended for huma outside the feed chain ◀	n consu
II.	Health infor	mation	II.a. Certificate reference No	II.b.
II.4.		of materials destined for organic fertilisers, cosmet n point J of Section 2 of Chapter IV of Annex IV to		, soil im
	(²) either	[the product does not contain and is not derived in 999/2001 or mechanically separated meat obtains which this product is derived have not been slauk illed by the same method or slaughtered by lack instrument introduced into the cranial cavity.]	ed from bones of bovine, ovine or cap ghtered after stunning by means of ga	orine ani as inject
	(²) or	[the product does not contain and is not derived animals born, continuously reared and slaughtere decision in accordance with Article 5(2) of Regula	ed in a country or region classified as	
Notes				
Part I:	!			
		Person responsible for the consignment in EU: this lificate is for import commodity.	box is to be filled in only if it is a certific	cate for t
	x reference I.11 hority.	1 and I.12: Approval number: the registration number	er of the establishment or plant, which	has bee
		2: Place of destination: this box is to be filled in only zones, free warehouses and custom warehouses.	if it is a certificate for a transit commod	dity. The
		5: Registration number (railway wagons or containe and reloading, the consignor must inform the BIP		r name
	x I.19: use the 17 or 15.18.	appropriate Harmonized System (HS) code under	the following headings: 15.01, 15.02;	15.03;
— Вох	k reference I.23	3: for bulk containers, the container number and the	e seal number (if applicable) should b	e includ
— Вох	k reference I.25	5: technical use: any use other than for animal con	sumption.	
— Вох	k reference I.26	6 and I.27: fill in according to whether it is a transit	t or an import certificate.	
— Вох	x reference I.28	В:		
— Spe	ecies: select fro	om the following: Ruminantia, Other		
— Ma	nufacturing pla	nt: provide the registration number of the treatment	t/processing establishment.	
Part II	:			
(^{1a}) O	J L 300, 14.11.	.2009, p. 1.		
(^{1b}) O	J L 54, 26.2.20	011, p. 1.		
(²) De	elete as approp	priate.		
(3) O	J L 147, 31.5.2	2001, p. 1.		
— The	signature and	the stamp must be in a different colour to that of	the printing.	
		son responsible for the consignment in EU: this it reaches the border inspection post.	certificate is only for veterinary purp	oses a
Official	veterinarian/O	official inspector		
Nar	me (in capital I	etters):	Qualifica	tion and

Chapter 11 is replaced by the following: (i)

Date: Stamp:

CHAPTER 11

Health certificate

For gelatine and collagen not intended for human consumption to be used as feed material or for purposes outside the feed chain, intended for dispatch to or for transit through (2) the European Union

cou	INTR	1		٧
	l.1.	Consignor Name	I.2. Certificate reference No	1.2.8
		Address	I.3. Central competent authority	
		Tel.	I.4. Local competent authority	
Part I: Details of dispatched consignment	1.5.	Consignee Name Address Postcode Tel.	I.6. Person responsible for the lo Name Address Postcode Tel.	pad in EU
ispatche	1.7.	Country of origin ISO code I.8. Region of origin Code	I.9. Country of ISO code destination	I.10. R
ð				
Part I: Details	l.11.	Place of origin	I.12. Place of destination	
		Name Approval number Address		ustom wa pproval nu
		Name Approval number Address	Postcode	
		Name Approval number Address		
	I.13.	Place of loading	I.14. Date of departure	
	l.15.	Means of transport	I.16. Entry BIP in EU	
		Aeroplane Ship Railway wagon Railway Railway wagon		
		Road vehicle Other I	I.17.	
		Documentation references		
	I.18.	Description of commodity	I.19. Commodity co	de (HS c
				I.20. Qu
	I.21.	Temperature of product Ambient ☐ Chilled ☐	Frozen	1.22. Nu
	1.23.	Seal/Container No		1.24. Тур
	1.25.	Commodities certified for:		
		Animal feedingstuff Technical	use 🗆	
	1.26.	For transit through EU to third country	I.27. For import or admission into	EU
		Third country ISO code		
	1.28.	Identification of the commodities		
		Species Approval number of establishments (Scientific name) Manufacturing plant	Number of packages	Net weigh

Health information

Gelatine and collagen not intended for hused as feed material or for purposes out

II.a. Certificate reference No

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Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

> COUNTRY II.

	II.	Health inf	orm	ation	II.a. Certificate reference No	II.b.		
		and of the	Cou	ned official veterinarian, declare that I have read a noil (^{1a}) and in particular Article 10 thereof, and Co reto, and certify that the gelatine/collagen (²) desc	ommission Regulation (EU) No 142/20	69/2009 11 (^{1b}), a		
ation	II.1.	consists of	f gel	atine/collagen (2) that satisfy the health requireme	ents below;			
ertifica	II.2.	consist ex	clusi	vely of gelatine/collagen (2) not intended for huma	an consumption;			
Part II: Certification	II.3.			ared and stored in a plant approved, validated ar c) No 1069/2009, in order to kill pathogenic agen	 d, validated and supervised by the competent authority in ac thogenic agents; 			
-	II.4.	has been	prep	ared exclusively with the following animal by-produced	l by-products:			
		(²) either	[-	carcases and parts of animals slaughtered or, in human consumption in accordance with Union reasons;]				
		(²) and/or	[-	carcases and the following parts originating eithe considered fit for slaughter for human consumptio animals from game killed for human consumption	on following an ante-mortem inspection	or bodie		
				(i) carcases or bodies and parts of animals whi legislation, but which did not show any signs				
				(ii) heads of poultry;				
				(iii) hides and skins, including trimmings and spli metacarpus bones, tarsus and metatarsus b		the pha		
				(iv) pig bristles;				
				(v) feathers;]				
		(²) and/or	[-	animal by-products arising from the production of greaves and centrifuge or separator sludge from		ımption,		
		(²) and/or	[-	products of animal origin, or foodstuffs containiconsumption for commercial reasons or due to which no risk to public or animal health arises;]				
		(²) and/or	[-	petfood and feedingstuffs of animal origin, or feed longer intended for feeding for commercial reas defects from which no risk to public or animal his	ons or due to problems of manufactu			
		(²) and/or	[-	aquatic animals, and parts of such animals, except to humans or animals;]	pt sea mammals, which did not show a	ny signs		
		(²) and/or	[-	animal by-products from aquatic animals origin consumption;]	ating from plants or establishments	manufac		
	II.5.	the gelatin	e/co	llagen (²):				
				 (a) was wrapped, packaged, stored and transport packaging took place in a dedicated room Wrappings and packages containing gelated FOR ANIMAL CONSUMPTION"; and 	n, and only preservatives permitted u	ınder Ur		
		(²) either		[(b) in the case of gelatine, has been produced subjected to a treatment with acid or alkali, heating one or several times in succession, kill pathogenic agents;]	, followed by one or more rinses, inve	olving ph		
		(²) or		(b) in the case of collagen, has been produce subjected to a treatment involving washing, g and extrusion, in order to kill pathogenic ag	oH adjustment using acid or alkali follo	nprocess wed by o		

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNTRY

Gelatine and collagen not intended for h used as feed material or for purposes out

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

(2) either

[the product does not contain and is not derived from specified risk material as defined in Anne 999/2001 of the European Parliament and of the Council (3) or mechanically separated meat obtovine or caprine animals; and the animals from which this product is derived have not been sli means of gas injected into the cranial cavity or killed by the same method or slaughtered by la tissue by means of an elongated rod-shaped instrument introduced into the cranial cavity;]

[the product does not contain and is not derived from bovine, ovine or caprine materials oth animals born, continuously reared and slaughtered in a country or region classified as posing decision in accordance with Article 5(2) of Regulation (EC) No 999/2001;] in the case of gelatine/collagen (2) from materials other than hides and skins: in addition as regards TSE:

in the case of gelatine/collagen (2) from materials other than hides and skins:

(2) either

(2) or

II.7.

[in case of animal by-products intended for feeding ruminants and containing milk or milk pr origin, the ovine and caprine animals from which these products are derived have been kept of the last three years on a holding where no official movement restriction is imposed due to a s has satisfied the following requirements for the last three years:

- (i) it has been subject to regular official veterinary checks:
- (ii) no classical scrapie case, as defined in point 2(g) of Annex I to Regulation (EC) No 999/20 following the confirmation of a classical scrapie case:
 - all animals in which classical scrapie was confirmed have been killed and destroyed, a
 - all goats and sheep on the holding have been killed and destroyed, except for breed genotype and breeding ewes carrying at least one ARR allele and no VRQ allele;
- (iii) ovine and caprine animals, with the exception of sheep of the ARR/ARR prion genotype, are only if they come from a holding which complies with the requirements set out in points (i

(2) or

[in case of animal by-products intended for feeding ruminants and containing milk or milk prorigin, and destined to a Member State listed in the Annex to Commission Regulation (EC) No caprine animals from which these products are derived have been kept continuously since birth on a holding where no official movement restriction is imposed due to a suspicion of TSE a following requirements for the last seven years:

- (i) it has been subject to regular official veterinary checks;
- (ii) no classical scrapie case, as defined in point 2(g) of Annex I to Regulation (EC) No 999/20 following the confirmation of a classical scrapie case:
 - all animals in which classical scrapie was confirmed have been killed and destroyed.
 - all goats and sheep on the holding have been killed and destroyed, except for breed genotype and breeding ewes carrying at least one ARR allele and no VRQ allele;
- (iii) ovine and caprine animals, with the exception of sheep of the ARR/ARR prion genotype, are only if they come from a holding which complies with the requirements set out in points (i

Notes

- Box reference I.6: Person responsible for the consignment in the European Union: this box is to be filled in only if 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 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 case of unloading and reloading, the consignor must inform the BIP of entry into the EU.
- Box I.19: use the appropriate Harmonized System (HS) code under the following headings: 35.03 or 35.04.
- Box reference I.23: for bulk containers, the container number and the seal number (if applicable) should be included
- Box reference I.26 and I.27: fill in according to whether it is a transit or an import certificate.

Box reference I.25: technical use: any use other than for animal consumption.

Box reference I.28: Species: select from the following: Aves, Ruminantia, Mammalia - Ruminantia, Pesca.

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

col	UNTRY	Gelatine and collagen not intended for hused as feed material or for purposes out
II.	Health information	II.a. Certificate reference No II.b.
Pa	t II:	<u> </u>
(^{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 the consignment until it reaches the border inspection post.	Union: this certificate is only for veterinary purp
Off	icial veterinarian/Official inspector	
	Name (in capital letters):	Qualification and
	Date:	Signature:
	Stamp:	

(j) Chapter 14(A) is replaced by the following:

CHAPTER 14(A)

Health certificate

For fat derivatives not intended for human consumption to be used outside the feed chain, intended for dispatch to or for transit through $\binom{2}{1}$ the European Union

cou	INTR	Y		٧					
	l.1.	Consignor Name	I.2. Certificate reference No I.2	2.					
		Address	I.3. Central competent authority						
		Tel.	I.4. Local competent authority						
Part I: Details of dispatched consignment	1.5.	Consignee Name Address Postcode Tel.	I.6. Person responsible for the load in E Name Address Postcode Tel.	Ū					
of dispatcl	1.7.	Country of origin ISO code I.8. Region of origin Code	I.9. Country of ISO code I.10. destination	R					
sils	1.11.	Place of origin	I.12. Place of destination	_					
l: Det		Name Approval number Address	Name Custom w Address Approval						
Part		Name Approval number Address	Postcode						
		Name Approval number Address							
	I.13.	Place of loading	I.14. Date of departure	_					
	l.15.	Means of transport	I.16. Entry BIP in EU	_					
		Aeroplane Ship Railway wagon Road vehicle Other							
		Identification Documentation references	1.17.	_					
	I.18.	Description of commodity	I.19. Commodity code (HS	С					
			I.20. G	ìu					
	I.21.	Temperature of product Ambient Chilled Chilled	Frozen I.22. N	lu					
	1.23.	Seal/Container No	I.24. T	У					
	1.25.	Commodities certified for:		_					
		Technical use							
	1.26.	For transit through EU to third country	I.27. For import or admission into EU						
		Third country ISO code							
	1.28.	28. Identification of the commodities							
		Species Approval number of establishments (Scientific name) Manufacturing plant	Number of packages Net weight						
	I								

Fat derivatives not intended for human

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c	OÚN	NTRY				outside the feed chain			
	Ī	II.	Health inforr	nat	ion	II.a. Certificate reference No	II.b.		
Part II: Certification			and of the C	oun	d official veterinarian, declare that I have read an icil (^{1a}) and in particular Article 10 thereof, and C ereto, and certify that the fat derivatives describ	Commission Regulation (EU) No 142/20			
	Ē 1	II.1.	consist of fat derivatives that satisfy the health requirements below;						
	a l	11.2.	consist of fat derivatives intended for purposes outside the feed chain, other than in cosmetics, pharmaceut						
	3 ı ≅ ı	II.3.		een prepared and stored in a plant approved, validated and supervised by the competent authority in action (EC) No 1069/2009, in order to kill pathogenic agents;					
٤	ا <u>۽</u>	II.4.	have been pr	have been prepared from rendered fats exclusively produced from the following materials:					
	ŀ	II.4.1.		the fat derivatives are intended for uses outside the feed chain, other than in organic fertilisers, ceuticals and medical devices, the following Category 1 materials:					
			(²) either	[-	the following material:				
					(i) specified risk material;				
					(ii) entire bodies or parts of dead animals cont	taining specified risk material at the tire	me of di		
			(²) and/or	[-	animal by-products which have been derived fr Article 1(2)(d) of Directive 96/22/EC or Article 2		ed to ille		
			(²) and/or	[-	animal by-products containing residues of othe Annex I to Directive 96/23/EC, if such residue absence thereof, by legislation of the Member	es exceed the permitted levels laid de			
	ľ	11.4.2.			erivatives are intended for use in organic fertilis naceuticals and medical devices, the following of		utside th		
			(²) either	[-	animal by-products containing residues of authoto in Article 15(3) of Directive 96/23/EC;]	rised substances or contaminants exc	eeding t		
			(²) and/or	[-	products of animal origin which have been declaration products;]	ared unfit for human consumption due	to the pr		
			(²) and/or	[-	animals and parts of animals, other than those rother than being slaughtered or killed for hum				
	ŀ	II.4.3.	the following	Cat	egory 3 materials:				
			(²) either	[-	carcases and parts of animals slaughtered or, in human consumption in accordance with Union reasons;]				
			(²) and/or	[-	carcases and the following parts originating eith considered fit for slaughter for human consumpl of animals from game killed for human consum	tion following an ante-mortem inspection	on or bo		
					(i) carcases or bodies and parts of animals whe legislation, but which did not show any significant controls.				
					(ii) heads of poultry;				
					(iii) hides and skins, including trimmings and sp metacarpus bones, tarsus and metatarsus		g the pha		
					(iv) pig bristles;				
					(v) feathers;]				
			(²) and/or	[-	blood of animals which did not show any signs from animals that have been slaughtered in a consumption following an ante-mortem inspection	slaughterhouse after having been cor	nsidered		
			(²) and/or	[-	animal by-products arising from the production greaves and centrifuge or separator sludge fro		umption,		

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNT	RY				Fat derivatives not intended for loutside the feed chain	human
II.	Healti	h informat	ion		II.a. Certificate reference No	II.b.
	(²) and	d/or [-	consumption		ining products of animal origin, which to problems of manufacturing or packa s;]	
	(²) and	d/or [-	no longer inte		eedingstuffs containing animal by-prod al reasons or due to problems of man animal health arises;]	
	(²) and	d/or [-		ta, wool, feathers, hair, horns, ho e communicable through that pr	oof cuts and raw milk originating from loduct to humans or animals;	ive anim
	(²) and	d/or [-		als, and parts of such animals to humans or animals;]	s, except sea mammals, which did	not sho
	(²) and	d/or [-	animal by-pro consumption;]		inating from plants or establishments	manufa
	(²) and	d/or [-		material originating from animal mans or animals:	s which did not show any signs of d	isease o
			(i) shells from	m shellfish with soft tissue or fle	esh;	
			(ii) the follow	ing originating from terrestrial ar	nimals:	
			— hatche	ery by-products,		
			— eggs,			
			— egg b	y-products, including egg shells;		
			(iii) day-old c	hicks killed for commercial reason	ons;]	
II.5.	in cas	e of fat de	rivatives produc	ced from animal by-products refe	erred to in point II.4.1 and point II.4.2:	
	(a) ha	ve been p	roduced using	the following methods:		
	(2)	either	[transesterifica acids and est		C, under corresponding appropriate pre-	ssure, fo
	(²)	or	[saponification	with NaOH 12M (glycerol and	soap):	
			(²) either	[in a batch process at 95 °C fo	or three hours;]	
			(2) or	[in a continuous process at 14	0 °C, 2 bars (2000 hPa) for eight minu	ıtes;]]
	(2)	or	[hydrogenation	n at 160 °C at 12 bars (12000 h	Pa) pressure for 20 minutes;]	
				ners or in containers that have "NOT FOR HUMAN OR ANIMA	been cleaned, and all precautions are AL CONSUMPTION";	taken t

Notes

II.6.

Part I:

 Box reference I.6: Person responsible for the consignment in the European Union: this box is to be filled in only if 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 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 case of unloading and reloading, the consignor must inform the BIP of entry into the EU.
- Box I.19: use the appropriate Harmonized System (HS) code under the following headings: 15.16 or 15.08.

Fat derivatives not intended for human

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNTRY outside the feed chain Health information II.a. Certificate reference No II. Box reference I.23: for bulk containers, the container number and the seal number (if applicable) should be included. 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. Box reference I.28: Species: select from the following: Ruminantia, Other; Manufacturing plant: provide the registration number of treatment/processing establishment. Part II: (1a) OJ L 300, 14.11.2009, p. 1. (1b) OJ L 54, 26.2.2011, p. 1. (2) Delete as appropriate. - 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 purporting the consignment until it reaches the border inspection post. Official veterinarian/Official inspector Name (in capital letters): Qualification and Date: Signature: Stamp:

(k) Chapter 15 is replaced by the following:

CHAPTER 15

Health certificate

For egg products not intended for human consumption that could be used as feed material, intended for dispatch to or for transit through $\binom{2}{1}$ the European Union

cou	INTR	1		٧	
	l.1.	Consignor Name	I.2. Certificate reference No	1.2.	
		Address	I.3. Central competent authority		
		Tel.	I.4. Local competent authority		
Part I: Details of dispatched consignment	I.5.	Consignee Name Address Postcode Tel.	Person responsible for the load in E Name Address Postcode Tel.		
f dispatc	1.7.	Country of origin ISO code I.8. Region of origin Code	I.9. Country of destination	l.10. R d	
ils o	1.11.	Place of origin	I.12. Place of destination		
l: Deta		Name Approval number Address	I	om wai	
Part		Name Approval number Address	Postcode	7701 110	
		Name Approval number Address			
	l.13.	Place of loading	I.14. Date of departure		
	l.15.	Means of transport	I.16. Entry BIP in EU		
		Aeroplane Ship Railway wagon Railway Railway Railway Railway wagon			
		Road vehicle Other I	1.17.		
		Documentation references			
	I.18.	Description of commodity	I.19. Commodity code	(HS c	
			1.2	20. Qu	
	I.21.	Temperature of product Ambient Chilled	Frozen	22. Nu	
	1.23.	Seal/Container No	1.2	24. Тур	
	1.25.	Commodities certified for:	•		
		Animal feedingstuff Technical	use		
	1.26.	For transit through EU to third country Third country ISO code	I.27. For import or admission into EU		
	1.28.	Identification of the commodities Approval number of establishments Manufacturing plant Number of positions of the commodities	ackages Net weight		

Egg products not intended for human coused as feed

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Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

COUNTRY

II.						
	Health info	rmation	II.a. Certificate reference No	II.b.		
	and of the C	signed official veterinarian, declare that I have reac ouncil (^{1a}) and in particular Article 10 thereof, and hereto, and certify that the egg products describe	Commission Regulation (EU) No 142/20	069/2009 011 (^{1b}), a		
II.1.	consist of egg products that satisfy the health requirements below;					
II.2.	consist exclusively of egg products not intended for human consumption;					
II.3.		orepared and stored in a plant, approved, validate EC) No 1069/2009 or Article 4(2) of Regulation (Ed plic agents;				
II.4.	have been p	prepared (derived) exclusively with the following a	animal by-products:			
	(²) either	[- animal by-products arising from the product	ion of products intended for human cor	nsumption		
	(²) and/or	[- products of animal origin, or foodstuffs con consumption for commercial reasons or due which no risk to public or animal health aris	e to problems of manufacturing or pack			
	(²) and/or	[- the following material originating from terrest that material to humans or animals:	trial animals which did not show any sig	ns of dise		
		 hatchery by-products, 				
		— eggs,				
		egg by-products, including egg shells;]				
II.5.	have been	subjected to processing:				
	(²) either	[in accordance with processing method No 142/2011;]	(4) as set out in Chapt	ter III of A		
	(²) or	[in accordance to a method and parameters wout in Chapter I of Annex X, to Regulation (EU		ith the m		
	(²) or	[in accordance with Section X, Chapters I and	II of Annex III to Regulation (EC) No 8	53/2004;]		
II.6.	have been of following sta	examined by the competent authority taking a randards $(^5)$:	ndom sample immediately prior to dispa	atch and		
	Salmonella:	absence in 25g: $n = 5$, $c = 0$, $m = 0$, N	$\Lambda = 0$,			
	Enterobacte	riaceae: n = 5, c = 2, m = 10, M = 300 in 1 gra	am;			
II.7.		standards on residues of substances that are hard dangerous or harmful to animal health;	mful or might alter the organoleptic char	acteristic		
II.8.	the end pro	duct was:				
	(²) either	[packed in new or sterilised bags,]				
	(²) or	[transported in bulk in containers or other means approved by the competent authority before us		ed and di		
	and which b	ear labels indicating "NOT FOR HUMAN CONSU	JMPTION";			
II.9.	the end pro	duct was stored in enclosed storage;				
1	the product	has undergone all precautions to avoid contamin	ation with pathogenic agents after treat	ment.		
II.10.	us .					
II.10.						

Egg products not intended for human co

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Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

cou	NTRY	Egg products not intended for huused as feed	ıman c
II.	Health information	II.a. Certificate reference No	II.b.
	Box reference I.12: Place of destination: this box is to be filled in onlibe stored in free zones, free warehouses and custom warehouses.	y if it is a certificate for transit commod	lity. The
	Box reference I.15: Registration number (railway wagons or contain case of unloading and reloading, the consignor must inform the BIP		or name
— I	Box I.19: use the appropriate Harmonized System (HS) code under	the following headings: 04.08, 23.09	or 35.02
— I	Box reference I.23: for bulk containers, the container number and the	ne seal number (if applicable) should b	e includ
— I	Box reference I.25: technical use: any use other than for animal con	nsumption.	
— I	Box reference I.26 and I.27: fill in according to whether it is a trans	it or an import certificate.	
Par	t 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 139, 30.4.2004, p. 55.		
(⁴)	Insert method 1 to 5 or 7 as applicable.		
(⁵)	Where:		
	n = number of samples to be tested;		
	$m=% \frac{1}{2}m^{2}$ threshold value for the number of bacteria; the result is consider $m;$	dered satisfactory if the number of bact	eria in a
	$M = \mbox{maximum value}$ for the number of bacteria; the result is consider or more; and	dered unsatisfactory if the number of ba	acteria ir
	c = number of samples the bacterial count of which may be betw count of the other samples is m or less.	een m and M, the sample still being co	onsidere
	The signature and the stamp must be in a different colour to that o	f the printing.	
	Note for the person responsible for the consignment in the European the consignment until it reaches the border inspection post.	Union: this certificate is only for vetering	ary purp
Offic	cial veterinarian/Official inspector		
1	Name (in capital letters):	Qualifica	ation and
	Date:	Signatur	e:
,	Stamp:		

in Annex XVI, Chapter III, Section 6 is replaced by the following:

Section 6

Official controls regarding the feeding of wild animals and certain zoo animals with Category 1 material

The competent authority shall monitor the health status of the farmed animals in the region where feeding is carried out as referred to in Sections 2, 3 and 4 of Chapter II of Annex VI and shall carry out appropriate TSE surveillance involving regular sampling and laboratory examination for TSEs.

Those samples shall include samples taken from suspected animals and from older breeding animals..

Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX. (See end of Document for details)

(1) OJ L 135, 30.5.1991, p. 40.';

Changes to legislation:

There are currently no known outstanding effects for the Commission Regulation (EU) No 294/2013, ANNEX.