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COMMISSION IMPLEMENTING REGULATION (EU) 2016/759

of 28 April 2016

drawing up lists of third countries, parts of third countries and territories from which Member States are to authorise the introduction into the Union of certain products of animal origin intended for human consumption, laying down certificate requirements, amending Regulation (EC) No 2074/2005 and repealing Decision 2003/812/EC

(Text with EEA relevance)

(OJ L 126, 14.5.2016, p. 13)

Amended by:

<u>B</u>

Official Journal

		No	page	date
<u>M1</u>	Commission Implementing Regulation (EU) 2016/1793 of 10 October 2016	L 274	48	11.10.2016
► <u>M2</u>	Commission Implementing Regulation (EU) 2017/731 of 25 April 2017	L 108	7	26.4.2017

COMMISSION IMPLEMENTING REGULATION (EU) 2016/759

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(Text with EEA relevance)

CHAPTER 1

IMPORTS OF CERTAIN PRODUCTS OF ANIMAL ORIGIN

Article 1

Lists of third countries, parts of third countries and territories

The third countries, parts of third countries and territories from which Member States are to authorise the import of the following products of animal origin intended for human consumption are set out in the relevant Parts of Annex I:

- (a) frogs' legs, Part I;
- (b) snails, Part II;
- (c) gelatine and collagen, Part III;
- (d) raw materials for the production of gelatine and collagen, Part IV;
- (e) treated raw materials for the production of gelatine and collagen, Part V;
- (f) honey, royal jelly and other products of apiculture, Part VI;
- (g) the following highly refined products, Part VII:
 - (i) chondroitin sulphate;
 - (ii) hyaluronic acid;
 - (iii) other hydrolysed cartilage products;
 - (iv) chitosan;
 - (v) glucosamine;
 - (vi) rennet;
 - (vii) isinglass;
 - (viii) amino acids that are authorised as food additives in accordance with Regulation (EC) No 1333/2008 of the European Parliament and of the Council (1).

Article 2

Model certificates

1. The model certificates for imports into the Union of the products referred to in Article 1 are set out in Annex II as follows:

Regulation (EC) No 1333/2008 of the European Parliament and of the Council of 16 December 2008 on food additives (OJ L 354, 31.12.2008, p. 16).

(a) frogs'	legs,	Part	I;

- (b) snails, Part II;
- (c) gelatine, Part III;
- (d) collagen, Part IV;
- (e) raw materials for the production of gelatine and collagen, Part V;
- (f) treated raw materials for the production of gelatine and collagen, Part VI;
- (g) honey, royal jelly and other products of apiculture, Part VII;
- (h) the following highly refined products, Part VIII:
 - (i) chondroitin sulphate;
 - (ii) hyaluronic acid;
 - (iii) other hydrolysed cartilage products;
 - (iv) chitosan;
 - (v) glucosamine;
 - (vi) rennet;
 - (vii) isinglass;
 - (viii) amino acids that are authorised as food additives in accordance with Regulation (EC) No 1333/2008.

Those certificates must be completed in accordance with the explanatory notes set out in Annex IV and the notes in the relevant certificate.

2. Electronic certification and other systems agreed between the Union and the third country concerned may be used.

CHAPTER 2

TRANSIT OF CERTAIN PRODUCTS OF ANIMAL ORIGIN

Article 3

Lists of third countries, parts of third countries and territories

The third countries, parts of third countries and territories from which Member States are to authorise the transit through the Union of raw materials and treated raw materials for the production of gelatine and collagen intended for human consumption bound for a third country, either by immediate transit or after storage in the Union in accordance with Article 12(4) and Article 13 of Council Directive 97/78/EC (1), are set out in Parts IV and V of Annex I to this Regulation, respectively.

⁽¹) Council Directive 97/78/EC of 18 December 1997 laying down the principles governing the organisation of veterinary checks on products entering the Community from third countries (OJ L 24, 30.1.1998, p. 9).

Article 4

Model certificate

1. The model certificate for the transit through the Union of the raw materials and treated raw materials referred to in Article 3 is set out in Annex III.

That certificate must be completed in accordance with the notes set out in Annex IV and in the relevant model certificate.

2. Electronic certification and other systems harmonised at Union level may be used.

Article 5

Derogation for transit through Latvia, Lithuania and Poland

- 1. By way of derogation from Article 3, transit by road or by rail between the specific, designated border inspection posts in Latvia, Lithuania and Poland, listed and marked with special remark 13 in Annex I to Commission Decision 2009/821/EC (¹), of consignments of the raw materials or treated raw materials referred to in Article 3 of this Regulation coming from and bound for Russia, directly or via another third country, shall be authorised where the following conditions are met:
- (a) the consignment is sealed with a serially numbered seal by the official veterinarian at the border inspection post of entry;
- (b) the documents accompanying the consignment, as provided for in Article 7 of Directive 97/78/EC, are stamped with the words 'Only for transit to Russia via the EU' on each page by the official veterinarian at the border inspection post of entry;
- (c) the procedural requirements provided for in Article 11 of Directive 97/78/EC are complied with;
- (d) the consignment is certified as acceptable for transit on the common veterinary entry document issued by the official veterinarian at the border inspection post of entry.
- 2. The consignments referred to in paragraph 1 shall not be unloaded or put into storage, as referred to in Article 12(4) or in Article 13 of Directive 97/78/EC, within the Union.
- 3. Regular audits shall be conducted by the competent authority to ensure that the number of consignments referred to in paragraph 1 and the corresponding quantities of products leaving the Union correspond with the number and quantities which have been introduced in the Union.

⁽¹) Commission Decision 2009/821/EC of 28 September 2009 drawing up a list of approved border inspection posts, laying down certain rules on the inspections carried out by Commission veterinary experts and laying down the veterinary units in Traces (OJ L 296, 12.11.2009, p. 1).

CHAPTER 3

FINAL PROVISIONS

Article 6

Amendment

Annex VI to Regulation (EC) No 2074/2005 is amended as follows:

- (1) in Section I, Chapters I, II, III and VI are deleted;
- (2) Appendices I, II, III and VI are deleted.

Article 7

Repeal

Decision 2003/812/EC is repealed.

Article 8

Transitional provisions

Consignments of products of animal origin in respect of which the relevant certificates have been issued in accordance with Regulation (EC) No 2074/2005 may continue to be introduced into the Union provided that the certificate was signed before 3 December 2016.

Article 9

Entry into force

This Regulation shall enter into force on the twentieth day following that of its publication in the Official Journal of the European Union.

This Regulation shall be binding in its entirety and directly applicable in all Member States.

ANNEX I

Lists of third countries, parts of third countries and territories as referred to in Article 1

PART I

FROGS' LEGS

Third countries and territories listed in the column 'Countries' of Annex II to Decision 2006/766/EC, except those for which a restriction is mentioned in the column 'Restrictions' of that Annex, and the following countries or territories:

COUNTRY ISO CODE	COUNTRY/TERRITORY
MK (*)	former Yugoslav Republic of Macedonia

^(*) The former Yugoslav Republic of Macedonia; provisional code that does not prejudge in any way the definitive nomenclature for this country, which will be agreed following the conclusion of negotiations currently taking place on this subject in the United Nations.

PART II

SNAILS

Third countries and territories listed in the column 'Countries' of Annex II to Decision 2006/766/EC, except those for which a restriction is mentioned in the column 'Restrictions' of that Annex, and the following countries/territories:

COUNTRY ISO CODE	COUNTRY/TERRITORY
MD	Moldova
MK (*)	former Yugoslav Republic of Macedonia
SY	Syria

^(*) The former Yugoslav Republic of Macedonia; provisional code that does not prejudge in any way the definitive nomenclature for this country, which will be agreed following the conclusion of negotiations currently taking place on this subject in the United Nations.

PART III

GELATINE AND COLLAGEN INTENDED FOR HUMAN CONSUMPTION

SECTION A

Gelatine and collagen derived from bovine, ovine, caprine, porcine and equine animals, both farmed and wild

Third countries and territories listed in column 1 of Part 1 of Annex II to Regulation (EU) No 206/2010 and the following countries or territories:

COUNTRY ISO CODE	COUNTRY/TERRITORY
KR	Republic of Korea
MY	Malaysia

▼B

COUNTRY ISO CODE	COUNTRY/TERRITORY
PK	Pakistan
TW	Taiwan

▼<u>M1</u>

SECTION B

Gelatine and collagen derived from poultry including ratites and feathered game

Third countries and territories listed in column 1 of Part 1 of Annex I to Regulation (EC) No 798/2008 and the following countries or territories:

COUNTRY ISO CODE	COUNTRY/TERRITORY
TW	Taiwan

▼B

SECTION C

Gelatine and collagen derived from fishery products

All third countries and territories listed in the column 'Countries' of Annex II to Decision 2006/766/EC, regardless of whether a restriction is mentioned in the column 'Restrictions' of that Annex.

$SECTION\ D$

Gelatine and collagen derived from leporidae and from wild land mammals not referred to in Section A

Third countries listed in column 1 of Part 1 of Annex I to Regulation (EC) No 119/2009.

PART IV

RAW MATERIALS FOR THE PRODUCTION OF GELATINE AND COLLAGEN INTENDED FOR HUMAN CONSUMPTION

SECTION A

Raw materials from bovine, ovine, caprine, porcine and equine animals, both farmed and wild

Third countries, territories and parts thereof listed in Part 1 of Annex II to Regulation (EU) No 206/2010 from which introduction into the Union of that category of fresh meat of the respective species is authorised as specified in that Part of that Annex, unless such introduction is limited by supplementary guarantees A or F as indicated in column 5.

SECTION B

Raw materials from poultry including ratites and feathered game

Third countries, parts of third countries and territories listed in Part 1 of Annex I to Regulation (EC) No 798/2008 from which imports of fresh poultry meat of the respective species is authorised as specified in that Part of that Annex.

SECTION C

Raw materials from fishery products

Third countries and territories listed in the column 'Countries' of Annex II to Decision 2006/766/EC, subject to the restrictions mentioned in the column 'Restrictions' of that Annex.

SECTION D

Raw materials from leporidae and from wild land mammals not referred to in Section A

Third countries listed in column 1 of Part 1 of Annex I to Regulation (EC) No 119/2009 from which imports of fresh meat of the respective species is authorised as specified in that Part of that Annex.

PART V

TREATED RAW MATERIALS FOR THE PRODUCTION OF GELATINE AND COLLAGEN INTENDED FOR HUMAN CONSUMPTION

SECTION A

Treated raw materials from bovine, ovine, caprine, porcine and equine animals, both farmed and wild

Third countries and territories and parts thereof listed in column 1 of Part 1 of Annex II to Regulation (EU) No 206/2010 and the following countries or territories:

COUNTRY ISO CODE	COUNTRY/TERRITORY
KR	Republic of Korea
MY	Malaysia
PK	Pakistan
TW	Taiwan

▼<u>M1</u>

SECTION B

Treated raw materials from poultry including ratites and feathered game

Third countries and territories listed in column 1 of Part 1 of Annex I to Regulation (EC) No 798/2008 and the following countries or territories:

COUNTRY ISO CODE	COUNTRY/TERRITORY
TW	Taiwan

▼B

SECTION C

Treated raw materials from fishery products

All third countries and territories listed in the column 'Countries' of Annex II to Decision 2006/766/EC regardless of whether a restriction is mentioned in the column 'Restrictions' of that Annex.

SECTION D

Treated raw materials from leporidae and wild land mammals not referred to in Section A

Third countries listed in column 1 of Part 1 of Annex I to Regulation (EC) No 119/2009.

SECTION E

Treated raw materials referred to in Annex III to Regulation (EC) No 853/2004, Section XIV, Chapter I point 4(b)(iii) and Section XV, Chapter I, point 4(b)(iii)

Third countries, parts of third countries and territories referred to in Part IV of this Annex.

PART VI

HONEY, ROYAL JELLY AND OTHER PRODUCTS OF APICULTURE INTENDED FOR HUMAN CONSUMPTION

Third countries and territories listed in the column 'Country' in the Annex to Commission Decision 2011/163/EU (¹) and marked with an 'X' in the column 'Honey' in that Annex.

PART VII

HIGHLY REFINED CHONDROITIN SULPHATE, HYALURONIC ACID, OTHER HYDROLYSED CARTILAGE PRODUCTS, CHITOSAN, GLUCOSAMINE, RENNET, ISINGLASS AND AMINO ACIDS FOR HUMAN CONSUMPTION

(a) In the case of raw materials derived from ungulates including equidae, third countries and territories listed in column 1 of Part 1 of Annex II to Regulation (EU) No 206/2010 and the following countries or territories:

COUNTRY ISO CODE	COUNTRY/TERRITORY
KR	Republic of Korea
MY	Malaysia
PK	Pakistan
TW	Taiwan

- (b) In the case of the raw materials derived from fishery products, all third countries and territories listed in the column 'Countries' in Annex II to Decision 2006/766/EC, regardless of whether a restriction is mentioned in the column 'Restrictions' of that Annex.
- (c) In the case of raw materials derived from poultry, third countries and territories listed in column 1 of Part 1 of Annex I to Regulation (EC) No 798/2008.

⁽¹) Commission Decision 2011/163/EU of 16 March 2011 on the approval of plans submitted by third countries in accordance with Article 29 of Council Directive 96/23/EC (OJ L 70, 17.3.2011, p. 40).

ANNEX II

Model certificates as referred to in Article 2

PART I

COL	JNTRY	':	Veterinary certificate to EU			
	l.1.	Consignor	I.2. Certificate reference No I.2.a.			
		Name	I.3. Central competent authority			
		Address	I.4. Local competent authority			
		Tel.				
	1.5.	Consignee	1.6.			
nent		Name				
signr		Address				
con						
ched		Postcode Tel.				
spat						
of di	1.7.	Country of ISO code I.8. origin	I.9. Country of ISO code I.10. destination			
stails						
Part I: Details of dispatched consignment	111	Place of origin	I.12.			
Part	1.11.	Flace of origin	1.12.			
		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				
		Aeroplane □ Ship □ Railway wagon □				
		Road vehicle Other O	1.17.			
		Identification Documentation references				
	I.18.	Description of commodity	1.19. Commodity code (HS code) 02.08.90			
			I.20. Quantity			
	I.21.	Temperature of product Ambient □ Chilled □	I.22. Number of packages			
	1.23.	Seal/Container No	I.24. Type of packaging			
	1.23.		Frozen 🗆			

I.25. Commodities certified for:							
	Human consump	tion 🗖					
1.26.				1.27.	For impor	t or admission into EU	
1.28.	Identification of th	ne commodities					
Treatment type e		Approval r establish Manufactu	nments		Number of packages	Net weight	

COUNTRY Model FRG Frogs' legs

II. **Health information** II.a. II.b. Certificate reference No II.1. **Public Health Attestation** I, the undersigned, declare that I am aware of the relevant provisions of Regulation (EC) No 178/2002 of the European Parliament and of the Council of 28 January 2002 laying down the general principles and requirements of food law, establishing the European Food Safety Authority and laying down procedures in matters of food safety (OJ L 31, 1.2.2002, p. 1), Regulation (EC) No 852/2004 of the European Parliament and of the Council of 29 April 2004 on the hygiene of foodstuffs (OJ L 139, 30.4.2004, p. 1) and Regulation (EC) No 853/2004 of the European Parliament and of the Council of 29 April 2004 laying Part II: Certification down specific hygiene rules for food of animal origin (OJ L 139, 30.4.2004, p. 55) and certify that the frogs' legs described above were produced in accordance with those requirements, in particular that they: come from (an) establishment(s) implementing a programme based on the HACCP principles in accordance with Article 5 of Regulation (EC) No 852/2004; and originate from frogs that have been bled, prepared and, where appropriate, chilled, frozen or processed, packaged and stored in a hygienic manner in accordance with the requirements of Section XI of Annex III to Regulation (EC) No 853/2004. **Notes** Part I: Box reference I.11: Place of origin: name and address of the dispatch establishment. Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship). Separate information is to be provided in the event of unloading and reloading. Box reference I.20: Indicate total gross weight and total net weight. Box reference I.23: Identification of container/Seal number: only where applicable. Box reference I.28: Treatment type: fresh, treated. Part II: The colour of the stamp and signature must be different from that of the other particulars in the certificate. Official inspector Name (in capital letters): Qualification and title: Date: Signature: Stamp:

PART II

MODEL CERTIFICATE FOR IMPORTS OF CHILLED, FROZEN, SHELLED, COOKED, PREPARED OR PRESERVED SNAILS INTENDED FOR HUMAN CONSUMPTION

COL	JNTRY	/ :	Veterinary certificate to EU			
	l.1.	Consignor	I.2. Certificate reference No I.2.a.			
		Name Address	I.3. Central competent authority			
		Address	I.4. Local competent authority			
		Tel.				
	1.5.	Consignee	1.6.			
nen		Name				
ısignr		Address				
loo pa		Postcode				
patch		Tel.				
Part I: Details of dispatched consignment	1.7.	Country of ISO code I.8. origin	I.9. Country of ISO code I.10. destination			
: Deta						
Part	l.11.	Place of origin	I.12.			
		Name Approval number				
		Address				
	I.13.	Place of loading	I.14. Date of departure			
	I.15.	Means of transport	I.16. Entry BIP in EU			
		Aeroplane ☐ Ship ☐ Railway wagon ☐				
		Road vehicle ☐ Other ☐	1.17.			
		Identification				
		Documentation references				
	I.18.	Description of commodity	I.19. Commodity code (HS code)			
			I.20. Quantity			
	I.21.	Temperature of product Ambient ☐ Chilled ☐	I.22. Number of packages			
	1.23.	Seal/Container No	I.24. Type of packaging			

I.25. Commodities certified for:			
Human consumption \square			
1.26.		I.27. For import or admission into EU	
I.28. Identification of the commoditi	es		
Species (scientific name)	Treatment type	Approval number of establish	ments
Number of packages	Net weight		
Manufacturing plant			

COUNTRY Model SNS Snails

	II.	Health information	II.a.	Certificate reference No	II.b.				
	II.1.	Public Health Attestation							
Part II: Certification		I, the undersigned, declare that I am aware of the relevant provisions of Regulation (EC) No 178/2002 of the European Parliament and of the Council of 28 January 2002 laying down the general principles and requirements of food law, establishing the European Food Safety Authority and laying down procedures in matters of food safety (OJ L 31, 1.2.2002, p. 1), Regulation (EC) No 852/2004 of the European Parliament and of the Council of 29 April 2004 on the hygiene of foodstuffs (OJ L 139, 30.4.2004, p. 1) and Regulation (EC) No 853/2004 of the European Parliament and of the Council of 29 April 2004 laying down specific hygiene rules for food of animal origin (OJ L 139, 30.4.2004, p. 55) and certify that the snails described above were produced in accordance with those requirements, in particular that they: — come from (an) establishment(s) implementing a programme based on the HACCP principles in							
Cer		accordance with Article 5 o			ed on the HACCE philoples in				
Ë		and							
Pa			hygienic n	nanner in accordance with th	prepared, preserved, frozen, e requirements of Section XI of				
	Note	es							
	Part	l:							
	_	Box reference I.11: Place of origin: r	ame and a	ddress of the dispatch establi	shment.				
		Box reference I.15: Registration nur name (ship). Separate information is							
		Box reference I.19: Use the appropr 16.05.	iate Harmo	onised System (HS) code und	er the following headings: 03.07,				
	-	Box reference I.20: Indicate total gro	ss weight a	and total net weight.					
	-	Box reference I.23: Identification of c	ontainer/S	eal number: only where applic	able.				
	_	Box reference I.28: Treatment type:	fresh, treat	ed.					
	Part	II:							
	-	The colour of the stamp and signatu	re must be	different from that of the other	r particulars in the certificate.				
	Offic	ial inspector							
		Name (in capital letters):			Qualification and title:				
		Date:			Signature:				
		Stamp:			-				
	1								

PART III

MODEL CERTIFICATE FOR IMPORTS OF GELATINE INTENDED FOR HUMAN CONSUMPTION

COL	COUNTRY:			Veterinary certificate to EU			
			1.2.	Certificate ref	erence No	I.2.a.	
		Name Address		Central comp	etent authority	/	
		Addiess	1.4.	Local compet	ent authority		
		Tel.					
	1.5.	Consignee	1.6.				
nen		Name					
nsignı		Address					
oo pe		Postcode					
patche		Tel.					
Part I: Details of dispatched consignment	1.7.	Country of ISO code I.8. origin	I.9.	Country of destination	ISO co	ode I.10.	
Part I: D	I.11.	I. Place of origin					
		Name Approval number					
		Address					
	I.13.	Place of loading	I.14.	Date of depar	ture		
	l.15.	Means of transport	I.16.	Entry BIP in E	EU		
		Aeroplane ☐ Ship ☐ Railway wagon ☐					
		Road vehicle Other	l.17.				
		Identification					
		Documentation references					
	I.18.	Description of commodity			I.19. Comm	nodity code (HS code)	
						I.20. Quantity	
	I.21.	Temperature of product Ambient □ Chilled □	F	Frozen 🗆		I.22. Number of packages	
	1.23.	Seal/Container No				I.24. Type of packaging	

1.25.	Commodities cert	ified for:				
	Human consumpt	ion 🗖				
1.26.				I.27. For	import or admission into EU	
1.28.	Identification of th	e commodities				
(so	Species cientific name)	Date production (dd/mm/yyyy)	Approval r establish Manufactu	nments	Number of packages	Net weight

Part II: Certification

Model GEL Gelatine intended for human consumption

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

II.1. Public Health Attestation

- I, the undersigned, declare that I am aware of the relevant provisions of Regulation (EC) No 178/2002 of the European Parliament and of the Council of 28 January 2002 laying down the general principles and requirements of food law, establishing the European Food Safety Authority and laying down procedures in matters of food safety (OJ L 31, 1.2.2002, p. 1), Regulation (EC) No 852/2004 of the European Parliament and of the Council of 29 April 2004 on the hygiene of foodstuffs (OJ L 139, 30.4.2004, p. 1) and Regulation (EC) No 853/2004 of the European Parliament and of the Council of 29 April 2004 laying down specific hygiene rules for food of animal origin (OJ L 139, 30.4.2004, p. 55) and certify that the gelatine described above was produced in accordance with those requirements, in particular that:
 - it comes from (an) establishment(s) implementing a programme based on the Hazard analysis and critical control points (HACCP) principles in accordance with Article 5 of Regulation (EC) No 852/2004,
 - it has been produced from raw materials that met the requirements of Chapters I and II of Section XIV of Annex III to Regulation (EC) No 853/2004,
 - it has been manufactured in compliance with the conditions set out in Chapter III of Section XIV of Annex III to Regulation (EC) No 853/2004,
 - it satisfies the criteria of Chapter IV of Section XIV of Annex III to Regulation (EC) No 853/2004 and of Commission Regulation (EC) No 2073/2005 of 15 November 2005 on microbiological criteria for foodstuffs (OJ L 338, 22.12.2005, p. 1);

and, if of bovine, ovine and caprine animal origin,

it has been derived from animals which have passed ante mortem and post mortem inspections,

- (1) and, except for gelatine derived from hides and skins,
- (¹) either [it comes from a country or a region classified in accordance with Commission Decision 2007/453/EC of 29 June 2007 establishing the BSE status of Member States or third countries or regions thereof according to their BSE risk (OJ L 172, 30.6.2007, p. 84) as a country or region posing a negligible BSE risk,
 - the gelatine does not contain and is not derived from specified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001 of the European Parliament and of the Council of 22 May 2001 laying down rules for the prevention, control and eradication of certain transmissible spongiform encephalopathies (OJ L 147, 31.5.2001, p. 1) (2),
 - the gelatine does not contain and is not derived from mechanically separated meat obtained from bones of bovine, ovine or caprine animals, except for gelatine derived from animals that were born, continuously reared and slaughtered in a country or region classified in accordance with Decision 2007/453/EC as a country or region posing a negligible BSE risk in which there have been no BSE indigenous cases.
 - the animals, from which the gelatine is derived were not slaughtered after stunning by means of gas injected into the cranial cavity or killed by the same method or slaughtered by laceration after stunning of central nervous tissue by means of an elongated rod-shaped instrument introduced into the cranial cavity, except if the animals were born, continuously reared and slaughtered in a country or region classified in accordance with Decision 2007/453/EC as a country or region posing a negligible BSE risk,
 - (¹) [the animals, from which the gelatine is derived, originate from a country or region classified in accordance with Decision 2007/453/EC as a country or region posing an undetermined BSE risk, and the animals were not fed with meat-and-bone meal or greaves, as defined in the Terrestrial Animal Health Code of the World Organisation for Animal Health],
 - (¹) [the animals, from which the gelatine is derived, originate from a country or region classified in accordance with Decision 2007/453/EC as a country or region posing an undetermined BSE risk, and the gelatine was produced and handled in a manner which ensures that it did not contain and were not contaminated with nervous and lymphatic tissues exposed during the deboning process.]]

Model GEL Gelatine intended for human consumption

II.	Health information		II.a.	Certificate reference No	II.b.			
	June 2007 establishir			region classified in accordance witl E status of Member States or third o . 172, 30.6.2007, p. 84) as a countr	countries or regions thereof ac-			
		central nervous tissue	e by mea	pelatine is derived, were not killed and some some some some instructions are some some some some some some some som				
			ation (E0	and is not derived from specified risl C) No 999/2001, or mechanically s rine animals.]				
	or	 [it comes from a country or a region classified in accordance with Decision 2007/453/EC of 2 June 2007 establishing the BSE status of Member States or third countries or regions thereof ac cording to their BSE risk (OJ L 172, 30.6.2007, p. 84) as a country or region posing a negligibl BSE risk, 						
		— the gelatine is derived both from animals born, continuously reared and slaughtered in a country or region with negligible BSE risk in accordance with Decision 2007/453/EC, and from animals born in a country or region classified in accordance with Decision 2007/453/EC as a country or region posing a controlled BSE risk, and which were not killed after stunning by laceration of central nervous tissue by means of an elongated rod-shaped instrument introduced into the cranial cavity, or by means of gas injected into the cranial cavity,						
			ation (E	and is not derived from specified ris C) No 999/2001, or mechanically s rine animals.]				
	or	June 2007 establishir	ng the BS	region classified in accordance wit E status of Member States or third o L 172, 30.6.2007, p. 84) as a cour	countries or regions thereof ac-			
				elatine is derived, were not fed meat- ed in the Terrestrial Animal health C				
		central nervous tissue	e by mea	gelatine is derived, were not killed on ns of an elongated rod-shaped instr s injected into the cranial cavity,				
		— the gelatine is not de	rived fror	ived from:				
		(i) specified risk ma	terial as	defined in point 1 of Annex V to Re	gulation (EC) No 999/2001;			
		(ii) nervous and lym	phatic tissues exposed during the deboning process;					
		(iii) mechanically sep	parated r	neat obtained from bones of bovine	, ovine or caprine animals.] ◀			
Notes	Notes							
Part I:								
— Во	ox referenc	ce I.11: Place of origin: na	me and	address of the dispatch establishn	nent.			

- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship). Separate information is to be provided in case of unloading and reloading.
- Box reference I.19: Use the appropriate Harmonised System (HS) code under the heading of 35.03.
- Box reference I.20: Indicate total gross weight and total net weight.
- Box reference I.23: Identification of container/Seal number: only where applicable.

COUNTRY

Model GEL Gelatine intended for human consumption

II. Health information	II.a. Certificate reference No	II.b.				
Part II:						
(1) Delete as appropriate.						
► ⁽¹⁾ (²) The removal of specified risk material is not required if the gelatine derives from animals born, continuously reared and slaughtered in a third country or region of a third country classified in accordance with Decision 2007/453/EC as posing a negligible BSE risk.						
The colour of the stamp and signature	e must be different from that of the other pa	rticulars in the certificate.				
Official veterinarian Name (in capital letters):						
Qualification and title:						
Date:	Sig	nature:				
Stamp:						

PART IV

MODEL CERTIFICATE FOR IMPORTS OF COLLAGEN INTENDED FOR HUMAN CONSUMPTION

COL	JNTRY	′ :	Veterinary certificate to EU			
	l.1.	Consignor	I.2. Certificate reference No I.2.a.			
		Name	I.3. Central competent authority			
		Address	I.4. Local competent authority			
		Tel.				
	1.5.	Consignee	1.6.			
nent		Name				
nsignr		Address				
d cor		Postcode				
chec		Tel.				
spat		161.				
Part I: Details of dispatched consignment	1.7.	Country of ISO code I.8. origin	I.9. Country of ISO code I.10. destination			
: Detai						
Part	l.11.	Place of origin	I.12.			
		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 ☐				
			1.17.			
		Road vehicle Other Identification				
		Documentation references				
	I.18	Description of commodity	I.19. Commodity code (HS code)			
		,	,,,			
			I.20. Quantity			
	l.21.	Temperature of product	I.22. Number of			
		Ambient ☐ Chilled ☐	Frozen packages			
	1.23.	Seal/Container No	I.24. Type of packaging			

1.25.	Commodities cert	ified for:				
	Human consumpt	ion 🗖				
1.26.				I.27. For	import or admission into EU	
1.28.	Identification of th	e commodities				
(so	Species cientific name)	Date production (dd/mm/yyyy)	Approval r establish Manufactu	nments	Number of packages	Net weight

Part II: Certification

Model COL Collagen intended for human consumption

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

II.1. Public Health Attestation

- I, the undersigned, declare that I am aware of the relevant provisions of Regulation (EC) No 178/2002 of the European Parliament and of the Council of 28 January 2002 laying down the general principles and requirements of food law, establishing the European Food Safety Authority and laying down procedures in matters of food safety (OJ L 31, 1.2.2002, p. 1), Regulation (EC) No 852/2004 of the European Parliament and of the Council of 29 April 2004 on the hygiene of foodstuffs (OJ L 139, 30.4.2004, p. 1) and Regulation (EC) No 853/2004 of the European Parliament and of the Council of 29 April 2004 laying down specific hygiene rules for food of animal origin (OJ L 139, 30.4.2004, p. 55) and certify that the collagen described above was produced in accordance with those requirements, in particular that:
 - it comes from (an) establishment(s) implementing a programme based on the Hazard analysis and critical control points (HACCP) principles in accordance with Article 5 of Regulation (EC) No 852/2004,
 - it has been produced from raw materials that met the requirements of Chapters I and II of Section XV of Annex III to Regulation (EC) No 853/2004,
 - it has been manufactured in compliance with the conditions set out in Chapter III of Section XV of Annex III to Regulation (EC) No 853/2004,
 - it satisfies the criteria of Chapter IV of Section XIV of Annex III to Regulation (EC) No 853/2004 and of Commission Regulation (EC) No 2073/2005 of 15 November 2005 on microbiological criteria for foodstuffs (OJ L 338, 22.12.2005, p. 1);
 - (1) and, if of bovine, ovine and caprine animal origin,
 - it has been derived from animals which have passed ante mortem and post mortem inspections,
 - (1) and, except for collagen derived from hides and skins,
 - (¹) either [it comes from a country or a region classified in accordance with Commission Decision 2007/453/EC of 29 June 2007 establishing the BSE status of Member States or third countries or regions thereof according to their BSE risk (OJ L 172, 30.6.2007, p. 84) as a country or region posing a negligible BSE risk,
 - the collagen does not contain and is not derived from specified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001 of the European Parliament and of the Council of 22 May 2001 laying down rules for the prevention, control and eradication of certain transmissible spongiform encephalopathies (OJ L 147, 31.5.2001, p. 1) (²),
 - the collagen does not contain and is not derived from mechanically separated meat obtained from bones of bovine, ovine or caprine animals, except for collagen derived from animals that were born, continuously reared and slaughtered in a country or region classified in accordance with Decision 2007/453/EC as a country or region posing a negligible BSE risk in which there have been no BSE indigenous cases,
 - the animals from which the collagen was derived have not been slaughtered after stunning by means of gas injected into the cranial cavity or killed by the same method or slaughtered by laceration after stunning of central nervous tissue by means of an elongated rod-shaped instrument introduced into the cranial cavity, except if the animals were born, continuously reared and slaughtered in a country or region classified in accordance with Decision 2007/453/EC as a country or region posing a negligible BSE risk,
 - (¹) [the animals, from which the collagen is derived, originate from a country or region classified in accordance with Decision 2007/453/EC as a country or region posing an undetermined BSE risk, and the animals were not fed with meat-and-bone meal or greaves, as defined in the OIE Terrestrial Animal Health Code],
 - (¹) [the animals, from which the collagen is derived, originate from a country or region classified
 in accordance with Decision 2007/453/EC as a country or region posing an undetermined BSE
 risk, and the collagen was produced and handled in a manner which ensures that it did not contain and were not contaminated with nervous and lymphatic tissues exposed during the deboning process.]]

Model COL Collagen intended for human consumption

II.	Health information			II.a.	Certificate reference No	II.b.
	(¹) or	_	June 2007 establishi	ng the BSE	egion classified in accordance wit E status of Member States or third 172, 30.6.2007, p. 84) as a countr	countries or regions thereof ac-
		_	central nervous tissu	e by mean	llagen is derived, were not killed, is of an elongated rod-shaped inst injected into the cranial cavity,	
		_		lation (EC	nd is not derived from specified ris) No 999/2001, or mechanically s ne animals.]	
	(¹) or	_	June 2007 establishi	ng the BSE	egion classified in accordance wit E status of Member States or third . 172, 30.6.2007, p. 84) as a coul	countries or regions thereof ac-
		_			agen is derived, were not fed meat d in the Terrestrial Animal health C	
		_	central nervous tissu	e by mean	ollagen is derived, were not killed is of an elongated rod-shaped inst injected into the cranial cavity,	
		_	the collagen is not de	erived from	1:	
			(i) specified risk ma	aterial as c	lefined in point 1 of Annex V to Re	gulation (EC) No 999/2001,
			(ii) nervous and lym	nphatic tiss	ues exposed during the deboning	process,
			(iii) mechanically se	parated m	eat obtained from bones of bovine	e, ovine or caprine animals.] ◀

Notes

Part I:

- Box reference I.11: Place of origin: name and address of the dispatch establishment.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship). Separate information is to be provided in case of unloading and reloading.
- Box reference I.18: This certificate may also be used for import of collagen casings.
- Box reference I.19: Use the appropriate Harmonised System (HS) code under the heading of 35.04 or 39.17.
- Box reference I.20: Indicate total gross weight and total net weight.
- Box reference I.23: Identification of container/Seal number: only where applicable.

Model COL Collagen intended for human consumption

II.	Health information		Certificate reference No	II.b.					
Part II:									
(1) D	(¹) Delete as appropriate.								
, the state of the	 (²) The removal of specified risk material is not required if the collagen derives from animals born, continuously reared and slaughtered in a third country or region of a third country classified in accordance with Decision 2007/453/EC as posing a negligible BSE risk. ◀ The colour of the stamp and signature must be different from that of the other particulars in the certificate. 								
Officia	l veterinarian								
	Name (in capital letters):		C	Qualification and title:					
	Date: Signature:								
	Stamp:								

PART V

MODEL CERTIFICATE FOR IMPORTS OF RAW MATERIALS FOR THE PRODUCTION OF GELATINE/COLLAGEN INTENDED FOR HUMAN CONSUMPTION $(\sline{1})$

COL	JNTR	/ :							Ve	terinary certificate to EU
	l.1.	Consignor					1.2.	Certificate i	reference No	I.2.a.
		Name					I.3. Central competent authority			
		Address					1.4.	Local comp	etent authority	1
		Tel.								
	1.5.	Consignee)				1.6.			
nent		Name								
signn		Address								
d con		Postcode								
che		Tel.								
ispa										
s of d	1.7.	Country of origin	ISO code	I.8. F	Region of origin	Code	1.9.	Country of destination	ISO code	I.10.
Part I: Details of dispatched consignment										
art I:	l.11.	Place of or	rigin				I.12.			
<u> </u>										
		Name		Аррі	roval nur	nber				
		Address								
	I.13.	Place of lo	ading				1.14.	Date of dep	parture	
	I.15.	Means of t	transport				I.16.	Entry BIP in	n EU	
		Aeroplane		Ship [∃ R	ailway				
		wagon 🗖			_		1.17.			
		Road vehic		Other	Ц					
		Identification Documents	on ation refere	nces						
	1.18.	Description	n of commo	aity					1.19. Comm	odity code (HS code)
										I.20. Quantity
	L21	Temperatu	are of produ	ct						I.22. Number of
		Ambient C			Chilled]		Frozen		packages
	1.23.	Seal/Conta	ainer No							I.24. Type of packaging

⁽¹⁾ Unless covered by Part VI.

1.25.	I.25. Commodities certified for:								
	Production of gelatine / collagen for human consumption \square								
1.26.				I.27. For impo	rt or admission into EU				
1.28.	Identification of the	commodities							
	Species entific name)	Nature of commodity	Approval n establish Manufactu	nments	Number of packages	Net weight			

	II.	Health	information	II.a.	Certificate reference No	II.b.						
	II.1.	Public	Health Attestation									
Part II: Certification	▶ ⁽¹⁾	pean Pa law, esta 31, 1.2.2 the hygie of the Co 55) and specific (OJ L 13	I, the undersigned, declare that I am aware of the relevant provisions of Regulation (EC) No 178/2002 of the European Parliament and of the Council of 28 January 2002 laying down the general principles and requirements of food law, establishing the European Food Safety Authority and laying down procedures in matters of food safety (OJ L 31, 1.2.2002, p. 1), Regulation (EC) No 852/2004 of the European Parliament and of the Council of 29 April 2004 on the hygiene of foodstuffs (OJ L 139, 30.4.2004, p. 1), Regulation (EC) No 853/2004 of the European Parliament and of the Council of 29 April 2004 laying down specific hygiene rules for food of animal origin (OJ L 139, 30.4.2004, p. 55) and Regulation (EC) No 854/2004 of the European Parliament and of the Council of 29 April 2004 laying down specific rules for the organisation of official controls on products of animal origin intended for human consumption (OJ L 139, 30.4.2004, p. 206) and certify that the raw materials described above comply with those requirements, in particular that:									
Part		(1) [bones, hides and skins of domestic and farmed ruminant animals, pigs and poultry and tendons and sinews described above are derived from animals which were slaughtered in a slaughterhouse and the carcasses of which were found fit for human consumption following ante and post mortem inspection,]										
		and	/or									
		bee	n found fit for human cons		cribed above are derived from killed owing post mortem inspection,]	I animals whose carcasses have						
		and										
			fish skins and bones desc sumption authorised for ex		are derived from plants manufactu	ring fishery products for human						
		(¹) a	(1) and, if of bovine, ovine and caprine animal origin,									
		 they have been derived from animals which passed ante mortem and post mortem inspections, 										
		(1) and, except for hides and skins of ruminants,										
		(1) either — [they come from a country or a region classified in accordance with Decision 2007/453/EC of 29 June 2007 establishing the BSE status of Member States or third countries or regions thereof according to their BSE risk (OJ L 172, 30.6.2007, p. 84) as a country or region posing a negligible BSE risk,										
			to Regulation (EC	No 999/20 or the preve	t derived from specified risk materia 01 of the European Parliament and ntion, control and eradication of cert 5.2001, p. 1) (⁶),	of the Council of 22 May 2001						
			bovine, ovine or c continuously reare	aprine anim d and slaug	t derived from mechanically separa als, except for raw materials derive htered in a country or region classil egion posing a negligible BSE risk in	ed from animals that were born, ied in accordance with Decision						
			means of gas injection after stunning duced into the cra	ted into the of central ne nial cavity, e ion classifie	raw materials are derived, were no cranial cavity or killed by the same re- virvous tissue by means of an elonga except if the animals were born, cont d in accordance with Decision 200	nethod or slaughtered by lacera- ted rod-shaped instrument intro- inuously reared and slaughtered						
			fied in accordance risk, and the anima	with Decision	e raw materials are derived, originat on 2007/453/EC as a country or regi fed with meat-and-bone meal or gre rld Organisation for Animal Health],	on posing an undetermined BSE						
			fied in accordance risk, and the raw m	with Decision aterials were	e raw materials are derived, originate on 2007/453/EC as a country or regi e produced and handled in a manne ated with nervous and lymphatic tis	on posing an undetermined BSE r which ensures that they did not						

II.	Health information		II.a.	. Certificate reference No II.b.						
	(¹) or	June 2007 establishi	ng the BS	a region classified in accordance wit SE status of Member States or third o L 172, 30.6.2007, p. 84) as a country	countries or regions thereof ac-					
		export are derived, w of an elongated rod	the animals, from which the raw materials of bovine, ovine and caprine animal origin intended for export are derived, were not killed, after stunning, by laceration of central nervous tissue by means of an elongated rod-shaped instrument introduced into the cranial cavity, or by means of gas njected into the cranial cavity,							
		specified risk materia	aw materials of bovine, ovine and caprine animal origin do not contain and are not derived from ified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001, or mechan-separated meat obtained from bones of bovine, ovine or caprine animals;							
	(¹) or	 (they come from a country or a region classified in accordance with Decision 2007, June 2007 establishing the BSE status of Member States or third countries or reaccording to their BSE risk (OJ L 172, 30.6.2007, p. 84) as a country or region with mined BSE risk, 								
			n rumina	raw materials are derived, were nonts, as defined in the Terrestrial Anima,						
		were not killed, after	which the raw materials of bovine, ovine and caprine animal origin are de fter stunning, by laceration of central nervous tissue by means of an elon ment introduced into the cranial cavity, or by means of gas injected into the							
		— the raw materials are	not deriv	red from:						
		(i) specified risk ma	aterial as	defined in point 1 of Annex V to Reg	ulation (EC) No 999/2001;					
		(ii) nervous and lym	phatic tis	ssues exposed during the de-boning	process;					
		(iii) mechanically se	parated r	meat obtained from bones of bovine,	ovine or caprine animals.] ◀					

II.	Health information		II.a.	Certificate reference No	II.b.						
(¹) [II.2.	Animal H	lealth Attestation									
	I, the und	, the undersigned official veterinarian, certify that the raw materials described above:									
II.2.1.	consist of	f animal products that	satisfy the	animal health requirements belo	w;						
II.2.2.	have bee from:	n obtained in the territ	ory of (1) e	either [:] (1) or	[] (²) (³) (⁴)						
(1) either	[II.2.2.1	animals that come fr the last 3 months be		gs and have remained in that terr nter; and	itory since birth or for at least						
	(¹) either	that are of the species referred to in Commission Regulation (EU) No 206/2010 of 12 March 2010 laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements (OJ L 73, 20.3.2010, p. 1), fulfilling all the relevant animal health import requirements laid down in that Regulation, and that were slaughtered for human consumption on a date for which import into the European Union of fresh meat from animals of those species was authorised from the country or territory thereof in accordance with Column 8 of Part 1 of Annex II to that Regulation;]									
,	(1) or [(ii) that are of the species referred to in Commission Regulation (EC) No 119/2009 of 9 February 2009 laying down a list of third countries or parts thereof, for imports into, or transit through, the Community of meat of wild leporidae, of certain wild land mammals and of farmed rabbits and the veterinary certification requirements (OJ L 39, 10.2.2009, p. 12), fulfilling all the relevant animal health import requirements laid down in that Regulation.]]										
(¹) or	[II.2.2.1	[II.2.2.1 poultry that have remained in that territory since hatching or have been imported as day-old chicks or slaughter poultry from (a) third country(ies) listed for that commodity in Part 1 of Annex I to Commission Regulation (EC) No 798/2008 of 8 August 2008 laying down a list of third countries, territories, zones or compartments from which poultry and poultry products may be imported into and transit through the Community and the veterinary certification requirements (OJ L 226, 23.8.2008, p. 1) under conditions at least equivalent to those in that Regulation, and consist of species referred to in that Regulation, fulfilling all the relevant animal health import requirements laid down in that Regulation, and were slaughtered for human consumption on a date for which import into the European Union of meat from animals of those species was authorised from the country or territory thereof in accordance with Column 6 B of Part 1 to Annex I to that Regulation.]									
(¹) or	[II.2.2.1	animals that have barea:	een killed	in the wild in that territory(5); a	nd captured and killed in an						
		(i) in which within 25 km there has been no case/outbreak of any of the following diseases for which the animals are susceptible: foot and mouth disease, rinderpest, Newcastle disease or highly pathogenic avian influenza during the prior 30 days, nor of classical or African swine fever during the prior 40 days and									
		territory of a	country o	nce that exceeds 20 km from the r part thereof, which is not au rials to the European Union, and	thorised at these dates for						
			e and imr	ey were transported within 12 h nediately afterwards to a game e							

Health information

II.

Model RCG Raw materials for the production of gelatine / collagen intended for human consumption

II.b.

II.2.3.	have been obtained in an establishment around which, within a radius of 10 km, there has been no case/outbreak of the following diseases for which the animals are susceptible: foot and mouth disease, rinderpest, Newcastle disease or highly pathogenic avian influenza, classical or African swine fever during the prior 30 days or, in the event of a case of one of those diseases, the preparation of raw materials for exportation to the European Union has been authorised only after the removal of all meat and the total cleaning and disinfection of the establishment under the control of an official veterinarian; and:

Certificate reference No

II.a.

- II.2.4. have been obtained and prepared without contact with other materials not complying with the conditions required above, and it has been handled so as to avoid contamination with pathogenic agents; and
- II.2.5. have been transported in clean and sealed containers or lorries.]

Notes

Part I:

- Box reference I.8: Provide the code of territory as appearing in Annex II to Commission Decision 2006/766/EC of 6 November 2006 establishing the lists of third countries and territories from which imports of bivalve molluscs, echinoderms, tunicates, marine gastropods and fishery products are permitted (OJ L 320, 18.11.2006, p. 53) and/or Part 1 of Annex I to Regulation (EC) 798/2008 and/or in Part 1 of Annex I to Regulation (EC) No 119/2009 and/or Part 1 of Annex II to Regulation (EU) No 206/2010.
- Box reference I.11: Place of origin: name and address of the dispatch establishment; registration or approval number as appropriate.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship). Separate information is to be provided in case of unloading and reloading.
- Box reference I.19: Use the appropriate Harmonised System (HS) code under the following headings: 02.08, 03.05, 05.05, 05.06, 05.11.91, 05.11.99, 41.01, 41.02, 41.03.
- Box reference I.20: Indicate total gross weight and total net weight.
- Box reference I.23: Identification of container/Seal number: only where applicable.
- Box reference I.28: Nature of commodity: hides, skins, bones, tendons and sinews;

Approval number of establishments: registration or approval number as appropriate;

Manufacturing plant: includes slaughterhouse, factory vessel, cutting plant, game handling establishment and processing plant.

Part II:

- Delete as appropriate. In case of products derived from fishery products, the whole section II.2 should be deleted.
- (2) The name and ISO code number of the exporting country or territory or zone as laid down in:
 - the Annexes to Decision 2006/766/EC;
 - Annex I to Regulation (EC) 798/2008;

COUNTRY

 Part 1 of Annex II to Regulation (EC) No 119/2009; Part 1 of Annex II to Regulation (EU) No 206/2010. If parts of the materials were derived from animals originating from an(other) third country(ies) listed in Annex II to Regulation (EU) No 206/2010 for import of that commodity into the Union, then the code(s) of country(ies) or territory(ies) and of the third country slaughtering the animals shall be indicated (the material cannot come from a country or territory that has supplementary guarantees A or F as indicated in column 5 of that Annex). If the materials were derived from slaughter poultry originating from an(other) third country(ies) listed in Part 1 of Annex I to Regulation (EC) No 798/2008 for imports of that commodity into the Union, then the code(s) of country(ies) or territory(ies) and of the third country slaughtering the poultry shall be indicated. Only for countries from where game meat intended for human consumption of the same animal species is authorised for importation into the European Union. If the removal of specified risk material is not required if the raw materials derive from animals born
 (3) If parts of the materials were derived from animals originating from an(other) third country(ies) listed in Annex II to Regulation (EU) No 206/2010 for import of that commodity into the Union, then the code(s) of country(ies) or territory(ies) and of the third country slaughtering the animals shall be indicated (the material cannot come from a country or territory that has supplementary guarantees A or F as indicated in column softhat Annex). (4) If the materials were derived from slaughter poultry originating from an(other) third country(ies) listed in Part 1 of Annex I to Regulation (EC) No 798/2008 for imports of that commodity into the Union, then the code(s) of country(ies) or territory(ies) and of the third country slaughtering the poultry shall be indicated. (5) Only for countries from where game meat intended for human consumption of the same animal species is authorised for importation into the European Union. ▶(1) (6) The removal of specified risk material is not required if the raw materials derive from animals born
Annex II to Regulation (EU) No 206/2010 for import of that commodity into the Union, then the code(s) or country(ies) or territory(ies) and of the third country slaughtering the animals shall be indicated (the material cannot come from a country or territory that has supplementary guarantees A or F as indicated in column 5 of that Annex). (4) If the materials were derived from slaughter poultry originating from an(other) third country(ies) listed in Part 1 of Annex I to Regulation (EC) No 798/2008 for imports of that commodity into the Union, then the code(s) of country(ies) or territory(ies) and of the third country slaughtering the poultry shall be indicated. (5) Only for countries from where game meat intended for human consumption of the same animal species is authorised for importation into the European Union. •• (6) The removal of specified risk material is not required if the raw materials derive from animals born
Part 1 of Annex I to Regulation (EC) No 798/2008 for imports of that commodity into the Union, then the code(s) of country(ies) or territory(ies) and of the third country slaughtering the poultry shall be indicated. (5) Only for countries from where game meat intended for human consumption of the same animal species is authorised for importation into the European Union. • (6) The removal of specified risk material is not required if the raw materials derive from animals born
authorised for importation into the European Union. • (6) The removal of specified risk material is not required if the raw materials derive from animals born
() The removal of specified lisk material is not required if the raw materials derive from animals born
continuously reared and slaughtered in a third country or region of a third country classified in accordance with Decision 2007/453/EC as posing a negligible BSE risk. ◀
 The signature and the stamp must be in a different colour to that of the printing.
NB Note for the person responsible for the consignment in the EU: this certificate is only for veterinary purposes and has to accompany the consignment until it reaches the border inspection post. The consignment must be transported directly to the manufacturing plant of destination.
Official veterinarian
Name (in capital letters): Qualification and title:
Date: Signature:
Stamp:

PART VI

MODEL CERTIFICATE FOR IMPORTS OF TREATED RAW MATERIALS FOR THE PRODUCTION OF GELATINE/COLLAGEN INTENDED FOR HUMAN CONSUMPTION

COL	JNTRY	' :							Ve	terinary certificate to EU
	l.1.	Consignor					1.2.	Certificate r	eference No	1.2.a.
		Name					I.3. Central competent authority			
		Address					I.4. Local competent authority			
		Tel.								
	1.5.	Consignee)				1.6.			
nent		Name								
signr		Address								
ed cor		Postcode								
patche		Tel.								
Part I: Details of dispatched consignment	1.7.	Country of origin	ISO code	1.8.	Region of origin	Code	1.9.	Country of destination	ISO code	1.10.
: Detai										
Part	l.11.	I.11. Place of origin			l.12.					
	Name Approval number									
		Address		, .h.						
	I.13.	Place of lo	ading				1.14.	Date of dep	parture	
	l.15.	Means of t	transport				I.16.	Entry BIP in	n EU	
		Aeroplane wagon		Ship	. □ R	ailway				
		Road vehic	cle \square	Oth∉	er 🗖		1.17.			
		Identification		Our	J. L.					
			ation refere	nces						
	I.18.	Description	n of commo	dity					I.19. Comm	nodity code (HS code)
										I.20. Quantity
	I.21.	Temperatu	ure of produ	ct						I.22. Number of
		Ambient 🗖]		Chilled]		Frozen \square		packages
	1.23.	Seal/Conta	ainer No							I.24. Type of packaging

1.25.	.25. Commodities certified for:								
	Production of gelatine / collagen for human consumption \square								
1.26.				I.27. For impo	rt or admission into EU				
1.28.	Identification of the	commodities							
(50	8. Identification of the commodities Species Nature of establish commodity Manufactu		nments	Number of packages	Net weight				

Model TCG Treated raw materials for the production of gelatine and collagen

	II.	Hea	alth info	ormation	II.a.	Certificate reference No	II.b.					
	II.1.	Pul	olic Hea	alth Attestation								
	▶ ⁽¹⁾	I, th	ne unde uiremer	ersigned, certify that the treated raw materials described above comply with the following ints:								
u		_	they ha		om establi	shments under the control of a	and listed by the competent					
icatic			and									
Part II: Certification			derived	from animals which w	ere slaugh	nd farmed ruminant animals, pigs a tered in a slaughterhouse and the e and <i>post mortem</i> inspection,]						
Par			(1) and/	or or								
		 [wild game hides, skins and bones described above are derived from killed animals whose care found fit for human consumption following post mortem inspection,] 										
			(¹) and/or									
		 [fish skins and bones described above are derived from plants manufacturing fishery product for human consumption authorised for export,] and 										
	(1) either — [they are dried bones of species from bovine, ovine, caprine, porcine and equine animal cluding farmed and wild animals, poultry including ratites and feathered game for the pition of collagen or gelatine, they derived from healthy animals slaughtered slaughterhouse, and they have been treated as follows:											
		(¹)	either	— [crushed to pieces of approximately 15 mm and degreased with hot water at a temperature of minimum 70 °C for at least 30 minutes, minimum 80 °C for at least 15 minutes, or minimum 90 °C for at least 10 minutes, and then separated and subsequently washed and dried for at least 20 minutes in a stream of hot air with an initial temperature of minimum 350 °C, or for 15 minutes in a stream of hot air with an initial temperature of more than 700 °C.]								
		(¹)	or	[sun-dried for a min	inimum of 42 days at an average temperature of at least 20 °C.]							
		(1)	or	[acid treatment such before drying.]]	nt such that the pH is maintained at less than 6 to the core for at least one hour]]							
		(1)	or	they are hides and skins of farmed ruminant animals, pig skins, poultry skins or wild game hid and skins, they derived from healthy animals and they:								
		(1)	either	— [have undergor lowed by salting the control of		cali treatment which ensures a east 7 days]	a pH > 12 to the core fol-					
		(¹)	or	[were dried for at le	east 42 da	ays at a temperature of at leas	t 20 °C.]					
		(¹)	or	[have undergone to the core for a		treatment that provides at le of 1 hour.]	east a pH of less than 5					
		(¹)	or	[have undergone a 8 hours.]]	an alkali ti	reatment which ensures a pH	> 12 to the core for at least					
		(1)	or	skins, fish skins third countries and plementing Regul countries, parts of authorise the introtended for human ulation (EC) No 2 14.5.2016, p. 13) above, and that c	and wild d territorication (EU third conduction consump 074/2005) that hacome from	skins of farmed ruminant a game hides and skins from es referred to in Part IV of Ar J) 2016/759 of 28 April 20 untries and territories from whito the Union of certain protion, laying down certificate recand repealing Decision 2 and repealing Decision 2 are undergone any other tree establishments registered to 852/2004 or in accordance	third countries, parts of nex I to Commission Im- Is drawing up lists of third ich Member States are to ducts of animal origin in- puirements, amending Reg- 003/812/EC (OJ L 126, atment than those listed or approved in accordance					
						caprine animal origin,						
						als which passed ante mortem and	d post mortem inspections,					
				(1) and, except for h	ides and s	kins of ruminants,						

Model TCG Treated raw materials for the production of gelatine and collagen

II.	Health infor	mation	II.a.	Certificate r	eference No	II.b.			
	(¹) either —	2007/453/EC of 2	29 June : ons there	2007 establisl of according to	hing the BSE statu their BSE risk (O	n accordance with Decision s of Member States or third J L 172, 30.6.2007, p. 84) as			
		1 of Annex V to Council of 22 Ma	Regulation y 2001 la	on (EC) No 9 aying down ru	99/2001 of the Eur	k material as defined in point ropean Parliament and of the on, control and eradication of 31.5.2001, p. 1) (4),			
	_	from bones of bo from animals tha classified in acco	they do not contain and are not derived from mechanically separated meat obtained from bones of bovine, ovine or caprine animals, except for treated raw materials derived from animals that were born, continuously reared and slaughtered in a country or region classified in accordance with Decision 2007/453/EC as a country or region posing a negligible BSE risk in which there have been no BSE indigenous cases,						
	_	ning by means of slaughtered by la gated rod-shape born, continuously	f gas injected and the following for the followi	ected into the after stunning ent introduced nd slaughtere	cranial cavity or k of central nervous into the cranial cav	were not slaughtered after stun- illed by the same method or tissue by means of an elon- ity, except if the animals were egion classified in accordance egligible BSE risk,			
	_	try or region clas posing an unde	sified in a termined meal or	accordance w BSE risk, greaves, as d	ith Decision 2007/49 and the animals	derived, originate from a coun- 53/EC as a country or region have not been fed with rial Animal Health Code of the			
	_	try or region clas posing an undet	sified in a ermined E It they did	accordance w BSE risk, the not contain a	ith Decision 2007/4: products were prodund and were not contan	derived, originate from a coun- 53/EC as a country or region uced and handled in a manner ninated with nervous and lym-			
	(¹) or —	2007/453/EC of 2	29 June : ons there	2007 establish of according to	hing the BSE statu their BSE risk (O	n accordance with Decision s of Member States or third J L 172, 30.6.2007, p. 84) as			
	_	origin destined for central nervous to	or export	are derived, means of an	were not killed, aff	ne, ovine and caprine animal ter stunning, by laceration of led instrument introduced into all cavity,			
	_	are not derived for	rom speci /2001, or	fied risk mate	rial as defined in p	mal origin do not contain and oint 1 of Annex V to Regula- tained from bones of bovine,			
	(¹) or —	2007/453/EC of 2	29 June 2 ons there	2007 establisl of according to	hing the BSE statu their BSE risk (O	n accordance with Decision s of Member States or third J L 172, 30.6.2007, p. 84) as			
	_	meat-and-bone m	eal or g	reaves derive		derived have not been fed as defined in the Terrestrial ealth,			

Model TCG Treated raw materials for the production of gelatine and collagen

II.	Health Information	II.a.	Certificate reference No	II.D.
	origin are derived by means of an	d, were no elongated	e treated raw materials of bovi of killed after stunning by lacera rod-shaped instrument introduce the cranial cavity,	ation of central nervous tissue
	 the treated raw m 	aterials are	e not derived from:	
	(i) specified risk	material a	s defined in point 1 of Annex V of F	Regulation (EC) No 999/2001;
	(ii) nervous and I	ymphatic t	ssues exposed during the de-boni	ng process;
	(iii) mechanically	separated	meat obtained from bones of bovi	ne, ovine or caprine animals.]]∢
(¹) [II.2.	Animal Health Attestation			
	I, the undersigned official veteri	inarian, ce	rtify that the treated raw materia	Is described above:
II.2.1.	consist of animal products that	satisfy the	animal health requirements bel	ow;
II.2.2.	have been obtained in the territ	ory(ies) of	: (¹) [] (¹) or [] (2) (3)
II.2.3.			hout contact with other mater een handled so as to avoid o	
II.2.4.	have been transported in clean	and seale	ed containers or lorries.]	
Notes				
Part I:				
2006 biva	reference I.8: Provide the co 6/766/EC of 6 November 2006 es Ive molluscs, echinoderms, tunic 1 2006, p. 53) or in Part 1 of Apr	tablishing ates, mari	the lists of third countries and te ne gastropods and fishery prod	rritories from which imports of ucts are permitted (OJ L 320,

- 2006/766/EC of 6 November 2006 establishing the lists of third countries and territories from which imports of bivalve molluscs, echinoderms, tunicates, marine gastropods and fishery products are permitted (OJ L 320, 18.11.2006, p. 53) or in Part 1 of Annex I to Commission Regulation (EC) 798/2008 of 8 August 2008 laying down a list of third countries, territories, zones or compartments from which poultry and poultry products may be imported into and transit through the Community and the veterinary certification requirements (OJ L 226, 23.8.2008, p. 1) or in Part 1 of Annex I to Commission Regulation (EC) No 119/2009 of 9 February 2009 laying down a list of third countries or parts thereof, for imports into, or transit through, the Community of meat of wild leporidae, of certain wild land mammals and of farmed rabbits and the veterinary certification requirements (OJ L 39, 10.2.2009, p. 12) or in Part 1 of Annex II to Commission Regulation (EU) No 206/2010 of 12 March 2010 laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements (OJ L 73, 20.3.2010, p. 1).
- Box reference I.11: Place of origin: name and address of the dispatch establishment and approval number or competent authority identification number as appropriate.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship). Separate information is to be provided in case of unloading and reloading.

COUNTRY

Model TCG Treated raw materials for the production of gelatine and collagen

II.	Health informat	ion	II.a.	Certificate reference	No	II.b.		
_	Box reference I.19: Use the appropriate Harmonised System (HS) code under the following headings: 03.05, 05.05, 05.06, 05.11.91, 05.11.99, 41.01, 41.02, 41.03.							
_	Box reference I.20: Indicate total gross weight and total net weight.							
_	Box reference I.23: Ide	ntification of co	ntainer/Se	eal number: only wher	e applicabl	e.		
_	Box reference I.28:	Nature of con	nmodity: h	ides, skins, bones, ter	idons and	sinews;		
		Approval nui			oval numl	per or competent authority		
				ncludes slaughterhous and processing plant;		vessel, cutting plant, game		
		Approval num	<i>ber:</i> wher	n applicable.				
Par	t II:							
(¹)	Delete as appropriate. deleted.	In case of pro	ducts de	rived from fishery pro-	ducts, the	whole section II.2 should be		
(²)	The name and ISO coo	le number of th	e exportir	g country or territory of	or zone as	laid down in:		
	 Part 1 of Annex II to Regulation (EU) No 206/2010; 							
	Annex I to Regulation (EC) 798/2008;							
	— Part 1 of Annex II	to Regulation ((EC) No 1	19/2009.				
(3)	If parts of the materials were derived from animals originating from an(other) third country(ies) listed in Annex I to Commission Implementing Regulation (EU) 2016/759 of 28 April 2016 drawing up lists of third countries, parts of third countries and territories from which Member States are to authorise the introduction into the Union of certain products of animal origin intended for human consumption, laying down certificate requirements, amending Regulation (EC) No 2074/2005 and repealing Decision 2003/812/EC (OJ L 126, 14.5.2016, p. 13), the code(s) of country(ies) or territory(ies) shall be indicated.							
▶ ⁽¹⁾	born, continuously	/ reared and s	laughtere		r region of	aterials derive from animals a third country classified in		
_	The signature and the	stamp must be	in a differ	ent colour to that of th	e printing.			
NB		consignment	until it rea	aches the border insp		y for veterinary purposes and t. The consignment must be		
_	The time of transportat	ion may be incl	uded in th	e duration of treatmer	ıt.			
Offic	cial veterinarian							
	Name (in capital let	ters):			C	Qualification and title:		
	Date:				S	ignature:		
	Stamp:							

PART VII

COL	JNTRY	f :	Veterinary certificate to EU			
	l.1.	Consignor	I.2. Certificate reference No I.2.a.			
		Name	I.3. Central competent authority			
		Address	I.4. Local competent authority			
		Tel.				
	1.5.	Consignee	1.6.			
ent	1.5.	Name	1.0.			
ignm		Address				
cons						
Part I: Details of dispatched consignment		Postcode				
spato		Tel.				
of dis	1.7.	Country of ISO code I.8. origin	I.9. Country of ISO code I.10. destination			
stails						
.: D						
Par	l.11.	Place of origin	1.12.			
		Name Approval number				
		Address				
	I.13.	Place of loading	I.14. Date of departure			
	I.15.	Means of transport	I.16. Entry BIP in EU			
		Aeroplane ☐ Ship ☐ Railway wagon ☐				
		Road vehicle ☐ Other ☐	1.17.			
		Identification				
		Documentation references				
	I.18.	Description of commodity	I.19. Commodity code (HS code)			
			I.20. Quantity			
	I.21.	Temperature of product	I.22. Number of			
		Ambient ☐ Chilled ☐	Frozen ☐ packages			
	1.23.	Seal/Container No	I.24. Type of packaging			

1.25.	Commodities cert	ified for:				
	Human consumpt	ion 🗖				
1.26.				I.27. For impo	rt or admission into EU	
1.28.	Identification of th	e commodities				
(sc	Species ientific name)	Treatment type	Approval r establish Manufactu	nments	Number of packages	Net weight

Part II: Certification

Model HON Honey, royal jelly and other apiculture products

		Tioney, it	by a jeny and other apiculture products				
II.	Health information	II.a. Certificate reference	e No II.b.				
II.1	. Public Health Attestation						
	I, the undersigned, declare that I am aware of the relevant provisions of Regulation (EC) No 178/2002 of the European Parliament and of the Council of 28 January 2002 laying down the general principles and requirements of food law, establishing the European Food Safety Authority and laying down procedures in matters of food safety (OJ L 31, 1.2.2002, p. 1), Regulation (EC) No 852/2004 of the European Parliament and of the Council of 29 April 2004 on the hygiene of foodstuffs (OJ L 139, 30.4.2004, p. 1) and Regulation (EC) No 853/2004 of the European Parliament and of the Council of 29 April 2004 laying down specific hygiene rules for food of animal origin (OJ L 139, 30.4.2004, p. 55) and certify that honey, royal jelly and other apiculture products described above were produced in accordance with those requirements, in particular that they:						
	 come from (an) establishme accordance with Regulation 	() ()	nme based on the HACCP principles in				
		ere appropriate, prepared, pac ments of Annex II to Regulation	kaged and stored in a hygienic manner in (EC) No 852/2004;				
	and						
-	— the guarantees covering live animals and products thereof provided by the residue plans submitted in accordance with Council Directive 96/23/EC of 29 April 1996 on measures to monitor certain substances and residues thereof in live animals and animal products and repealing Directives 85/358/EEC and 86/469/EEC and Decisions 89/187/EEC and 91/664/EEC (OJ L 125, 23.5.1996, p. 10), and in particular Article 29 thereof, are fulfilled.						
Not	tes						
Par	rt I:						
_	Box reference I.11: Place of origin: means registration number.	name and address of the d	ispatch establishment. Approval number				
_	Box reference I.15: Registration numl name (ship). Separate information is to		ner and lorries), flight number (aircraft) or ding and reloading.				
_	Box reference I.19: Use the appropria 04.10.	ate Harmonised System (HS) o	code under the following headings: 04.09,				
—	Box reference I.20: Indicate total gros	s weight and total net weight.					
—	Box reference I.23: Identification of co	ontainer/Seal number: only whe	ere applicable.				
_		<i>type</i> : Indicate 'ultrasonica on', 'no thermal treatment'.	tion', 'homogenisation', ultrafiltration',				
		mber of establishments: ap number as appropriate	proval number or competent authority				
Pai	rt II:						
_	The colour of the stamp and signature must be different from that of the other particulars in the certificate.						
Offi	icial inspector						
	Name (in capital letters):		Qualification and title:				
	Date:		Signature:				
	Stamp:						

PART VIII

MODEL CERTIFICATE FOR IMPORTS OF HIGHLY REFINED CHONDROITIN SULPHATE, HYALURONIC ACID, OTHER HYDROLYSED CARTILAGE PRODUCTS, CHITOSAN, GLUCOSAMINE, RENNET, ISINGLASS AND AMINO ACIDS INTENDED FOR HUMAN CONSUMPTION

COL	JNTRY	′ :	Vet	erinary certificate to EU	
	l.1.	Consignor	I.2. Certificate reference No	I.2.a.	
		Name Address	I.3. Central competent authority		
		Addiess	I.4. Local competent authority		
		Tel.			
=	1.5.	Consignee	1.6.		
mer		Name			
nsign		Address			
oo p		Postcode			
che		Tel.			
spat		161.			
Part I: Details of dispatched consignment	1.7.	Country of ISO code I.8. origin	I.9. Country of ISO cool destination	de I.10.	
: Detail					
Part	l.11.	Place of origin	1.12.		
		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		
		Aeropla <u>ne</u> ☐ Ship ☐ Railway			
		wagon			
		Road vehicle Other	I.17.		
		Identification			
		Documentation references			
	I.18.	Description of commodity	I.19. Commo	odity code (HS code)	
		,			
				I.20. Quantity	
	I.21.	Temperature of product		I.22. Number of	
		Ambient ☐ Chilled ☐	Frozen 🗖	packages	
	1.23.	Seal/Container No		I.24. Type of packaging	

1.25.	Commodities cert	ified for:				
	Human consumpt	ion 🗖				
1.26.				I.27. For	import or admission into EU	
1.28.	Identification of th	e commodities				
(so	Species Date production Approval nu establish (scientific name) (dd/mm/yyyy) Manufactur		nments	Number of packages	Net weight	

Model HRP
Highly refined chondroitin sulphate, hyaluronic acid,
other hydrolysed cartilage products, chitosan,
glucosamine, rennet, isinglass and amino acids for
human consumption

II. I	Health information	II.a.	Certificate reference No	II.b.
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II.1. Public Health Attestation

I, the undersigned, declare that I am aware of the relevant provisions of Regulation (EC) No 178/2002 of the European Parliament and of the Council of 28 January 2002 laying down the general principles and requirements of food law, establishing the European Food Safety Authority and laying down procedures in matters of food safety (OJ L 31, 1.2.2002, p. 1), Regulation (EC) No 852/2004 of the European Parliament and of the Council of 29 April 2004 on the hygiene of foodstuffs (OJ L 139, 30.4.2004, p. 1) and Regulation (EC) No 853/2004 of the European Parliament and of the Council of 29 April 2004 laying down specific hygiene rules for food of animal origin (OJ L 139, 30.4.2004, p. 55) and certify that the highly refined products described above were produced in accordance with those requirements, in particular:

- that they come from (an) establishment(s) implementing a programme based on the HACCP principles in accordance with Regulation (EC) No 852/2004;
- that they have been handled and, where appropriate, prepared, packaged and stored in a hygienic manner in accordance with the requirements of Annex II to Regulation (EC) No 852/2004;
- that they comply with the requirements of Section XVI of Annex III to Regulation (EC) No 853/2004;

and

- (¹) [if amino acids, that
 - (i) human hair was not used as a source for their manufacture; and
 - (ii) they comply with Regulation (EC) No 1333/2008 of the European Parliament and of the Council of 16 December 2008 on food additives ((OJ L 354, 31.12.2008, p. 16)]

Notes

Part I:

- Box reference I.11: Place of origin: name and address of the dispatch establishment.
- Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship). Separate information is to be provided in case of unloading and reloading.
- Box reference I.19: Use the appropriate Harmonised System (HS) code under the heading of 21.06.90, 29.22, 29.30, 29.32, 35.07, 35.03 or 39.13.
- Box reference I.20: Indicate total gross weight and total net weight.
- Box reference I.23: Identification of container/Seal number: only where applicable.

Part II:

- (1) Delete as appropriate.
- The colour of the stamp and signature must be different from that of the other particulars in the certificate.

Official veterinarian		Name (in capital letters):
	Qualification and title:	
	Date:	Signature:
	Stamp:	

Part II: Certification

ANNEX III

MODEL CERTIFICATE FOR THE TRANSIT THROUGH THE UNION, IMMEDIATE TRANSIT OR AFTER STORAGE, FOR RAW MATERIALS OR TREATED RAW MATERIALS FOR THE PRODUCTION OF GELATINE/COLLAGEN FOR HUMAN CONSUMPTION

COL	COUNTRY:				Veterinary certificate to EU					
	l.1.	Consignor					1.2.	Certificate i	reference No	I.2.a.
	Name Address			1.3.	Central cor	npetent autho	rity			
		Address					1.4.	Local comp	etent authorit	у
		Tel.								
	1.5.	Consignee)				1.6.	Person res	ponsible for th	ne consignment in EU
Ħ		Name						Name		
gnme		Address						Address		
onsi		Postcode						Postcode		
ped c		Tel.						Tel.		
Part I: Details of dispatched consignment	1.7.	Country of origin	ISO code	1.8.	Region of origin	Code	1.9.	Country of destination	ISO code	I.10.
ls of										
: I: Detai	l.11.	. Place of origin				I.12.	Place of de	stination		
Part		Name						Customs w	arehouse 🗖	Ship supplier 🗖
		Address								
								Name		Approval number
								Address		
								Postal code	e	
	113	Place of lo	ading				114	Date of dep		
	1.10.	riace of to	ading				1.14.	Date of dep	Janui e	
	I.15.	Means of t	ransport				I.16.	Entry BIP in	n EU	
		Aeroplane wagon 🗖		Ship	□ R	ailway				
		Road vehic	cle 🗖	Othe	r 🗖		1.17.			
		Identification	on							
		Document	ation referer	nces						
	I.18.	Description	n of commo	dity					I.19. Comn	nodity code (HS code)
										I.20. Quantity
	I.21.	Temperatu		ct	Chilled]		Frozen 🗆		I.22. Number of packages
	1.23.	Seal/Conta	ainer No							I.24. Type of packaging

1.25.	Commodities certified for:						
	Production of gelatine / collagen for human consumption \square						
1.26.	For transit through EU to third country	1.27.					
l	Third country ISO code						
1.28.	Identification of the commodities						
	Species Manufacturing plant (scientific name)	Number of packages	Net weight				

COUNTRY Model TRANSIT/STORAGE

II. Health information II.a. Certificate reference number II.b.

II.1. Public Health Attestation

I, the undersigned official veterinarian, hereby certify, that the raw materials or treated raw materials described in Part I:

- II.1.1. come from a country or region authorised for imports into the EU as laid down in Part 1 of Annex I to Commission Regulation (EC) 798/2008 of 8 August 2008 laying down a list of third countries, territories, zones or compartments from which poultry and poultry products may be imported into and transit through the Community and the veterinary certification requirements (OJ L 226, 23.8.2008, p. 1) or in Commission Regulation (EC) No 119/2009 of 9 February 2009 laying down a list of third countries or parts thereof, for imports into, or transit through, the Community of meat of wild leporidae, of certain wild land mammals and of farmed rabbits and the veterinary certification requirements (OJ L 39, 10.2.2009, p. 12) or in Part 1 of Annex II to Commission Regulation (EU) No 206/2010 of 12 March 2010 laying down lists of third countries, territories or parts thereof authorised for the introduction into the European Union of certain animals and fresh meat and the veterinary certification requirements (OJ L 73, 20.3.2010, p. 1), and
- II.1.2. comply with the relevant animal health conditions as laid down in the animal health attestation in the model certificate in Part V or VI of Annex II to Commission Implementing Regulation (EU) 2016/759 of 28 April 2016 drawing up lists of third countries, parts of third countries and territories from which Member States are to authorise the introduction into the Union of certain products of animal origin intended for human consumption, laying down certificate requirements, amending Regulation (EC) No 2074/2005 and repealing Decision 2003/812/EC (OJ L 126, 14.5.2016, p. 13).

Notes

Part II: Certification

This certificate is meant for transit and storage in accordance with Article 12(4) or Article 13 of Council Directive 97/78/EC of 18 December 1997 laying down the principles governing the organisation of veterinary checks on products entering the Community from third countries (OJ L 24, 30.1.1998, p. 9) of raw materials or treated raw materials for the production of gelatine/collagen for human consumption of:

- (1) domestic bovine animals (including Bubalus and Bison species and their cross-breeds);
- (2) domestic ovine animals (Ovis aries) or domestic caprine animals (Capra hircus);
- (3) domestic porcine animals (Sus scrofa);
- (4) domestic solipeds (Equus caballus, Equus asinus and their cross-breeds);
- (5) farmed non-domestic animals of the order Artiodactyla (excluding bovine animals (including Bison and Bubalus species and their cross-breeds), Ovis aries, Capra hircus, Suidae and Tayassuidae), and of the families Rhinocerotidae and Elephantidae;
- (6) wild non-domestic animals of the order Artiodactyla (excluding bovine animals (including Bison and Bubalus species and their cross-breeds), Ovis aries, Capra hircus, Suidae and Tayassuidae), and of the families Rhinocerotidae and Elephantidae;
- (7) farmed non-domestic animals belonging to the Suidae, Tayassuidae, or Tapiridae families;
- (8) wild non-domestic animals belonging to the Suidae, Tayassuidae, or Tapiridae families;
- (9) wild solipeds belonging to the subgenus Hippotigris (zebra);
- (10) wild leporidae (rabbits and hares);
- (11) wild land mammals other than ungulates and leporidae;
- (12) farmed rabbits;
- (13) poultry;
- (14) farmed ratites;
- (15) wild game;
- (16) fish.

Model TRANSIT/STORAGE

II.	Health information	II.a.	Certificate reference number	II.b.				
Par	Part I:							
_	Box reference I.8: Provide the code of territory as appearing in Part 1 of Annex II to Regulation (EU) No 206/2010 or in Part 1 of Annex I to Regulation (EC) No 119/2009 or Part 1 of Annex I to Regulation (EC) No 798/2008 or Annex II to Commission Decision 2006/766/EC of 6 November 2006 establishing the lists of third countries and territories from which imports of bivalve molluscs, echinoderms, tunicates, marine gastropods and fishery products are permitted (OJ L 320, 18.11.2006, p. 53).							
_	Box reference I.11: Place of origin: na	ime and a	ddress of the dispatch establish	nment.				
_	Box reference I.12: Address (and warehouse, customs warehouse or sh			rehouse in a free zone, free				
_	 Box reference I.15: Registration number (railway wagons or container and lorries), flight number (aircraft) or name (ship) is to be provided. In case of unloading and reloading, the consignor must inform the BIP of entry into the EU. 							
_	Box reference I.19: Use the appropri 05.04, 05.05, 05.06, 05.11.91, 05.11.91			er the heading of 02.08, 03.05,				
_	Box reference I.20: Indicate total gros	s weight a	nd total net weight.					
_	Box reference I.23: For containers or boxes, the container number and the seal number (if applicable) should be included.							
_	 Box reference I.28: Manufacturing plant: provide the registration number, approval number or competent authority identification number of establishment as appropriate. It includes slaughterhouse, factory vessel, cutting plant, game handling establishment and processing plant. 							
Offi	Official veterinarian or Official inspector							
	Name (in capital letters): Qualification and title:							
	Date: Signature:							
	Stamp:							

ANNEX IV

EXPLANATORY NOTES FOR COMPLETING THE CERTIFICATES

(referred to in Articles 2(1) and 4(1))

(a) Certificates shall be issued by the exporting third country, based on the models set out in Annexes II and III according to the layout of the model that corresponds to the products of animal origin concerned.

They shall contain, in the numbered order that appears in the model, the attestations that are required for any third country and, as the case may be, those supplementary guarantees that are required for the exporting third country or part thereof.

If the Member State of destination imposes, for the products of animal origin concerned, additional certification requirements, attestations to certify that those requirements are fulfilled shall also be incorporated in the original form of the certificate.

- (b) Where the model certificate states that certain statements shall be kept as appropriate, statements which are not relevant, may be crossed out nd initialled and stamped by the certifying officer, or completely deleted from the certificate.
- (c) A separate and unique certificate must be provided for the products of animal origin that are exported from a territory or territories or zone or zones of the same exporting country listed or referred to in Annex I which are consigned to the same destination and transported in the same railway wagon, lorry, aircraft or ship.
- (d) The original of each certificate shall consist of a single sheet of paper, or, where more text is required it must be in such a form that all sheets of paper required are part of an integrated whole and indivisible.
- (e) The certificate shall be drawn up in at least one of the official languages of the Member State of the border inspection post of entry of the consignment into the EU and of the Member State of destination. However, those Member States may authorise the certificate to be drawn up in the official language of another Member State, and accompanied, if necessary, by an official translation.
- (f) If for reasons of identification of the items of the consignment (schedule in point I.28 of the model certificate), additional sheets of paper are attached to the certificate, those sheets of paper shall also be considered as forming part of the original of the certificate by the application of the signature and stamp of the certifying officer, on each of the pages.
- (g) When the certificate, including additional sheets of paper referred to in (f), comprises more than one page, each page shall be numbered, (page number) of (total number of pages), at the end of the page and shall bear the certificate reference number that has been designated by the competent authority at the top of the pages.
- (h) The original of the certificate must be completed and signed by an official veterinarian or by another designated official inspector where this is provided for in the model certificate. The competent authorities of the exporting third country shall ensure that rules of certification equivalent to those laid down in Council Directive 96/93/EC (¹) are followed.

The colour of the signature shall be different from that of the printing. This requirement also applies to stamps other than those embossed or water-marked.

 The certificate reference number referred to in boxes I.2 and II.a. must be issued by the competent authority.

Council Directive 96/93/EC of 17 December 1996 on the certification of animals and animal products (OJ L 13, 16.1.1997, p. 28).