Changes to legislation: There are outstanding changes not yet made to Commission Regulation (EU) No 28/2012. Any changes that have already been made to the legislation appear in the content and are referenced with annotations. (See end of Document for details)

Commission Regulation (EU) No 28/2012 of 11 January 2012 laying down requirements for the certification for imports into and transit through the Union of certain composite products and amending Decision 2007/275/ EC and Regulation (EC) No 1162/2009 (Text with EEA relevance)

Changes to legislation: There are outstanding changes not yet made to Commission Regulation (EU) No 28/2012. Any changes that have already been made to the legislation appear in the content and are referenced with annotations. (See end of Document for details)

[F1ANNEX I

Model Health Certificate for import into the European Union of composite products intended for human consumption]

Textual Amendments

F1 Substituted by Commission Implementing Regulation (EU) No 468/2012 of 1 June 2012 amending Regulation (EU) No 28/2012 laying down requirements for the certification for imports into and transit through the Union of certain composite products (Text with EEA relevance).

COUNTRY Veterinary certificate to EU							
	l.1.	Consignor Name Address	Certificate reference No I.2.a. Central competent authority				
		Tel.	I.4. Local competent authority				
Ħ	1.5.	Consignee	I.6.				
ignme	1.0.	Name Address					
suoo pa		Postcode Tel.					
dispatched consignment	1.7.	Country of origin ISO code I.8. Region of origin Code	I.9. Country of destination ISO code I.10.				
ğ	1.11.	Place of origin	1.12.				
Part I: Details		Name Approval number Address					
Pg		Name Approval number Address					
		Name Approval number Address					
	I.13.	Place of loading	I.14. Date of departure				
	I.15.	Means of transport	I.16. Entry BIP in EU				
		Aeroplane Ship Railway wagon Road vehicle Other Ship Railway wagon Shi	1.17.				
		Identification Documentation references					
	I.18.	Description of commodity	I.19. Commodity code (HS code)				
			I.20. Quantity				
	1.21.	Temperature of product Ambient ☐ Chilled ☐	I.22. Number of packages				
	1.23.	Seal/Container No	I.24. Type of packaging				
	1.25.	Commodities certified for:					
		Human consumption					
	1.26.		I.27. For import or admission into EU				
	1.28. Identification of the commodities						
			ature of Net weight Batch number mmodity				

Changes to legislation: There are outstanding changes not yet made to Commission Regulation (EU) No 28/2012. Any changes that have already been made to the legislation appear in the content and are referenced with annotations. (See end of Document for details)

COUNTRY Composite products intended for human consumption Health information II.a. Certificate reference No I, the undersigned official veterinarian/official inspector hereby certify that Certification I am aware of the relevant provisions of Regulations (EC) No 178/2002, (EC) No 852/2004 and (EC) No 853/2004, in particular Article 6.1(b) on the origin of the products of animal origin used in the production of the composite products described above and certify that the composite 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: Part the composite products described above contain: (1) either [II.2.A Meat products, treated stomachs, bladders and intestines (2) in any quantity which meet the animal health requirements in Commission Decision 2007/777/EC and contain the following meat constituents which meet the criteria indicated below: Approved Establishment(s) (D) Species (A) Treatment (B) Origin (C) (A) Insert the code for the relevant species of meat product, treated stomachs, bladders and intestines where BOV = domestic bovine animals (Bos taurus, Bison bison, Bubalus bubalis and their crossbreds); OVI = domestic sheep (Ovis aries) and goats (Capra hircus); EQI = domestic equine animals (Equus caballus, Equus asinus and their crossbreds), POR = domestic porcine animals (Sus scrola); RM = domestic rabbits, PFG = domestic poultry and farmed feathered game, RUF farmed non-domestic animals other than suidae and solipeds; RUW = wild non-domestic animals other than suidae and solipeds; SUW = wild non-domestic suidae: EQW = wild non-domestic solipeds, WL = wild lagomorphs, WGB = wild game birds. (B) Insert A, B, C, D, E or F for the required treatment as specified and defined in Parts 2, 3 and 4 of Annex II to Decision 2007/777/EC. (C) Insert the ISO code of the country of origin of the meat product, treated stomachs, bladders and intestines as listed in Annex II, Part 2 to Decision 2007/777/EC and, in the case of regionalization by Union legislation for the relevant meat constituents, the region as indicated in Part 1 of Annex II to Decision 2007/777/EC or a Member State of the European Union. The country of origin of the meat products must be one the following: - the same as the country of export in box I.7. a Member State of the European Union. a third country or parts thereof authorised to export to the Union meat products treated with treatment A as set out in Annex II to Decision 2007/777/EC, where the third country where the composite product is produced is also authorised to export to the Union meat products treated with that treatment. (D) Insert EU approval number of the establishments of origin of the meat products, treated stomachs, bladders and intestines contained in the composite product. ▶⁽¹⁾ (E) If containing material from bovine, ovine or caprine animals, the fresh meat and/or intestines used in the preparation of the meat products and/or treated intestines shall be subject to the following conditions depending on the BSE risk category of the (1)[(E.1) for imports from a country or a 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 products of bovine, ovine and caprine animal origin are derived, have passed ante mortem and post mortem inspection: 2. the products of bovine, ovine and caprine animal origin do not contain and are 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 (11); 3. the products of bovine, ovine and caprine animal origin do not contain and are not derived from mechanically separated meat obtained from the bones of bovine, ovine or caprine animals, except for products of bovine, ovine and caprine animal origin 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

4. the animals, from which the products of bovine, ovine and caprine animal origin are 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;

in which there have been no BSE indigenous cases;

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COUNTRY

Health information

II.

Composite products intended for human consumption

II.b.

if the animals, from which the products of	bovine, ovine and caprine animal origin are derived, originate from a country
or region classified in accordance with D	ecision 2007/453/EC as posing an undetermined BSE risk, those 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, and the	products were produced and handled in a manner which ensures that it

(*) or [(E.2) for imports from a country or a region classified in accordance with Decision 2007/453/EC as a country or region posing a controlled BSE risk:

II.a. Certificate reference No

did not contain and was not contaminated with nervous and lymphatic tissues exposed during the deboning process.]

- the animals, from which the products of bovine, ovine and caprine animal origin are derived, have passed
 ante mortem and post mortem inspection and 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;
- the products of bovine, ovine and caprine animal origin do not contain and are not derived from specified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001, or mechanically separated meat obtained from the bones of bovine, ovine or caprine animals.
- (¹) (⁴) 3. In the case of intestines originally sourced from a country or a region with a negligible BSE risk, imports of treated intestines have been subject to the following conditions:
 - (a) the country or region was classified in accordance with Decision 2007/453/EC as a country or region posing a controlled BSE risk;
 - (b) the animals, from which the products of bovine, ovine and caprine animal origin are derived, were born, continuously reared and slaughtered in the country or region with a negligible BSE risk and have passed ante mortem and post mortem inspections;
 - (1) (c) if the intestines are sourced from a country or region where there have been BSE indigenous cases:
 - (i) the animals were born after the date from which the ban on the feeding of ruminants with meatand-bone meal and greaves derived from ruminants was enforced; or
 - (ii) the products of bovine, ovine and caprine animal origin do not contain and are not derived from specified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001.)
- (*) or [(E.3) for imports from a country or a region classified in accordance with Decision 2007/453/EC as a country or region with an undetermined BSE risk:
 - the animals, from which the products of bovine, ovine and caprine animal origin are derived, were not fed
 meat-and-bone meal or greaves derived from ruminants, as defined in the Terrestrial Animal health Code
 of the World Organisation for Animal Health, and have passed ante mortem and post mortem inspections;
 - the animals, from which the products of bovine, ovine and caprine animal origin 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 injected into the cranial cavity;
 - 3. the products of bovine, ovine and caprine animal origin are not derived from:
 - (a) specified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001;
 - (b) nervous and lymphatic tissues exposed during the deboning
 - (c) mechanically separated meat obtained from bones of bovine, ovine or caprine animals.
 - (¹) (⁴) 4. In the case of intestines originally sourced from a country or a region with a negligible BSE risk, imports of treated intestines have been subject to the following conditions:
 - (a) the country or region was classified in accordance with Decision 2007/453/EC as a country or region posing a undetermined BSE risk;
 - (b) the animals, from which the products of bovine, ovine and caprine animal origin are derived, were born, continuously reared and slaughtered in the country or region with a negligible BSE risk and have passed ante mortem and post mortem inspections;

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COUNTRY

Composite products intended for human consumption

II. Health information				II.a. Certificate reference No	II.b.		
	Ticulti	mormation					
(') (c) if the intestines are sourced from a country or region where there have been BSE indige							
				fter the date from which the ban on the feedir derived from ruminants was enforced; or	ng of ruminants with meat-and-		
(ii) the products of bovine, ovine and caprine animal origin do not contain and are not d specified risk material as defined in point 1 of Annex V to Regulation (EC) No 999/2001							
(¹) and/or	and/or [II.2.B Processed dairy products (6) in an amount of half or more of the substance of the composite product or not shelf stable deproducts in any quantity that:						
	(a) have been produced in the country						
		— the sam	e as the country of export in box I.	7,			
		— a Memb	per State of the European Union,				
	 a third country authorised to export to the Union milk and dairy products in Column A or B of Annex I to Regulation No 605/2010, where the third country where the composite product is produced is also authorised, under the conditions, to export to the Union milk and dairy products. 						
	The country of origin indicated in box I.7 must be listed in Annex I to Regulation (EU) No 605/2010 and the treatment apmust be conform to the treatment provided for in that list for the relevant country;						
	(b) have been produced from milk obtained from animals:						
	(i) under the control of the official veterinary service;						
(ii) belonging to holdings which were not under restrictions due to foot-and-mouth disease or rinderpest; and							
				ensure that they satisfy the animal health cont No 853/2004 and in Directive 2002/99/EC;	ditions laid down in Chapter I of		
		(c) are dairy pr	oducts made from raw milk obtaine	d from:			
(1) either [cows, ewes, goats or buffaloes and produced from raw milk which has und					Union have undergone or been		
		(¹) eithei	achieved by a pasteurisation prod	ng a single heat treatment with a heating eless of at least 72 °C for 15 seconds and alkaline phosphatase test applied immedia	where applicable, sufficient to		
		(¹) or	[a sterilisation process, to achieve	an F ₀ value equal to or greater than three;]			
		(¹) or	[an ultra high temperature (UHT) tr	reatment at not less than 135 °C in combination	on with a suitable holding time;]		
		(¹) or		teurisation treatment (HTST) at 72 °C for 15 sapplied to milk with a pH lower than 7,0 anosphatase test			
		(¹) or	equivalent pasteurisation effect, ap	teurisation treatment (HTST) at 72 °C for 15 soplied twice to milk with a pH equal to or grean alkaline phosphatase test, immediately for	eater than 7,0 achieving, where		
	(1) either [lowering the pH below 6 for one hour;]						
		C,	or [additional heating equal to	or greater than 72 °C, combined with desico	cation;]]		
			als other than cows, ewes, goats or gone or been produced from raw r	r buffaloes and prior to import into the territo nilk which has undergone	ry of the European Union have		

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COUNTRY	,						Composite products into	ended for human consumptio
II.	Health	infor	mat	ion			II.a. Certificate reference No	II.b.
			(1)	eithe	r [a ste	rilisation process, to achieve	an F ₀ value equal to or greater than three;]	
			(1)	or	[an ul	tra high temperature (UHT) tre	eatment at not less than 135 °C in combination	on with a suitable holding time;]]
							or between	
(1) and/or	III.2.C						e approved establishment No (8)	situated
()	į <u>.</u>]	(,	
(1) and/or	[II.2.D	Proce	esse	ed eg	g prod	ucts that originate from the a	pproved country (9)]
		(EC)	No	853/2	2004 w		ment which satisfies the requirements of Sec the certificate is free from highly pathogeni	
		eithei	r					
	(1)) II.2.D					where appropriate, the territory of a neighbounza or Newcastle disease for at least the pro-	
		or						
	(1)) II.2.D	.2	[the e	gg pro	ducts were processed:		
			(1)	either	[liqui	d egg white was treated:		
				(¹) eithe	r [with 55,6 °C for 870 secon	nds.]	
				(¹) or	[with 56,7 °C for 232 secon	nds.]	
			(1)	or	[10 9	6 salted yolk was treated with	n 62,2 °C for 138 seconds.]	
			(1)	or	[drie	d egg white was treated:		
				(¹) eithe	r [with 67 °C for 20 hours.]		
				(1) or	[with 54,4 °C for 513 hours	.]	
			(1)	or	[who	le eggs were at least treated:		
				(¹) eithe	r [with 60 °C for 188 second	s.]	
				(1) or	[completely cooked.]		
					[who	le egg blends were at least tr	reated]:	
				(¹) eithe	r [with 60 °C for 188 second	s.]	
				(1) or	[with 61,1 °C for 94 second	ds.]	
Notes								
Part I:								
intesti No 60	nes as li 5/2010 a	isted ir and/or	n Ar for	nnex II proces	l, Part 2 ssed fis	to Decision 2007/777/EC and	composite product containing meat product, d/or for processed dairy products in Annex I I to Commission Decision 2006/766/EC and/	to Commission Regulation (EU)
	 Box reference I.11: Name, address and registration/approval number if available of the establishments of production of the composite product(s) Name of the country of origin which must be the same as the country of origin in box I.7. 							
transp	ort in co	ontaine	rs,	the to	tal num	ber of containers and their re	ner and road vehicles), flight number (aircraf gistration number and where there is a seria gnor must inform the border inspection post o	al number of the seal it must be
						te Harmonised System (HS) o	code of the World Customs Organisation sucl 06.	h as: 16.01; 16.02; 16.03; 16.04;

- Box reference I.20: Indicate total gross weight and total net weight.

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cou	INTRY	Composite products inte	nded for human consumptio					
II.	Health information	II.a. Certificate reference No	II.b.					
_	Box reference I.23: For containers or boxes, the container number	and the seal number (if applicable) must be	e included.					
	— Box reference I.28: Manufacturing plant: insert the name and approval number if available of the establishments of production of the compoproduct(s). Nature of commodity: in case of composite products containing meat products, treated stomachs, bladders and intestines indic "meat product", "treated stomachs", "bladders" or "intestines". In case of composite product containing dairy product indicate "dairy product case of composite product containing processed fishery products specify whether aquaculture or wild origin. In case of composite product containing egg products specify the egg content percentage.							
Par	t II:							
(¹)	Keep as appropriate.							
(²)	Meat products as laid down in point 7.1 of Annex I to Regulation (E in point 7.9 of Annex I to Regulation (EC) No 853/2004 that have 2007/777/EC.							
(³)	By way of derogation from point 4, carcasses, half carcasses or containing no specified risk material other than the vertebral column							
	When removal of the vertebral column is not required, carcasses of shall be identified by a blue stripe on the label referred to in Reg		nals containing vertebral column					
	The number of bovine carcasses or wholesale cuts of carcasses, f where removal of the vertebral column is not required shall be added in case of imports.							
(⁴)	Only applicable to imports of treated intestines.							
(⁵)	By way of derogation from point 3, carcasses, half carcasses or containing no specified risk material other than the vertebral column							
	When removal of the vertebral column is not required, carcasses of shall be identified by a clearly visible blue stripe on the label reference.		nals containing vertebral column					
	Specific information on the number of bovine carcasses or wholess and from which removal of the vertebral column is not required sha 136/2004 in case of imports.							
(⁶)	Raw milk and dairy products means, raw milk and dairy products f No $853/2004$.	for human consumption as defined in point 7.	2 of Annex I to Regulation (EC)					
(7)	Date or dates of production. Imports of raw milk and dairy product for exportation to the European Union of the third country or par measures have been adopted by the European Union against im	rt thereof mentioned under I.7 and I.8, or di	uring a period where restrictive					
(8)	Number of the fishery product establishment authorised to export	to the EU.						
(₉)	Country of origin authorised to export to the EU.							
(10)	In case of composite products containing only egg or fishery products	ducte the cignoture of an official Inspector of	un he accented					

(I) (11) The removal of specified risk material is not required if the products of bovine, ovine and caprine animal origin 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 colour of the signature shall be different to that of the printing. The same rule applies to stamps other than those embossed or watermark.

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COUNTRY	Composite products	Composite products intended for human consumptio			
II. Health information	II.a. Certificate reference No	II.b.			
Official veterinarian/Official inspector (10)	·				
Name (in capital letters):	Qualification and title:	Qualification and title:			
Date:	Signature:				
Stamp:					

Status:

Point in time view as at 31/01/2020.

Changes to legislation:

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