	I.1. Consignor				I.2. IMSOC Reference						
	Name						I.2.a. Local Reference				
	Address										
	Country			ISO Cod	e						
	I.5. Consignee						I.3. Central competent authority				
	Name						I.4. Local competent authority				
en	Address										
nu	Country			ISO Cod	e						
Part I : Details of consignment	I.7. Country of orig	gin				ISO Code	I.9. Country of	destinatio	n		ISO Code
of (I.8. Region of origi	n				Code	I.10. Region of	destinatio	n		
ls (I.11. Place of Dispa	itch					I.12. Place of d	estination			
tai	Name						Name				
De	Address						Address				
Ξ	Approval Number	r					Approval Nur	nber			
art]	Country			ISO	Code		Country			ISO Code	
Pŝ	I.13. Place of Loadi	ing					I.14. Date and time of departure				
	Name										
	Address										
	Approval Number	ſ									
	Country			ISO	Code						
	I.15. Means of Tran	nsport					I.16 Entry Poir	ıt			
	Mode	Internation	nal	Identificatio	on						
		transport document									
	I.18. Transport cor	nditions					I.17. Accompanying documents				
	Frozen	Chilled \Box		Ambient 🗆		Controlled	Accompanyi ng Date of issue document				
					t	temperature \Box					
							reference				
							Country		Place o issue	of	
	I.19. Container No	/ Seal No									
	I.20. Certified as										
	Human consumpti	ion 🗆									
	I.21. For transit thi	rough a thir	rd coun	try			I.22. For transit through Member State(s)				
	Country			ISO Code							
	EU Exit Authority			BCP code			Country ISO Code				
	EU Entry Authority			BCP code							
	I.23. Total number	of package	s		I.25. To	otal net weight	I.25. Total gross weight				
	128 Description of	f consignme	nt								
	I.28. Description of consignment 1. 21 MISCELLANEOUS EDIBLE PREPARATIONS 2106 Food preparations not elsewhere specified or included										
	Commodity		Manu	facturing pla	nt	Package coun	t	Nature of	commodity	Net weight	
	Batch number										

E	UROPEAN UNION					GBHC088E (v1.3)			
	II. Health info	ormation							
	I. the unde	rsigned offi	cial veterina	rian/official inspector h	Level of the second sec				
ation	II.1.	I am awar 853/2004, ii production described a from (an) e	e of the rele n particular of the comp above were	vant provisions of Regu Article 6.1(b) on the ori posite products describe produced in accordance nt(s) implementing a pro	lations (EC) No 178/2002, (EC) igin of the products of animal ed above and certify that the c e with those requirements, in p ogramme based on the HACCP	origin used in the omposite products particular that they come			
ifica	II.2.	the compos	site products	s described above conta	in:				
Part II: Certification	(1)	either □ [11.2.A	animal hea	Meat products. treated stomachs, bladders and intestines(2) in any quan animal health requirements in Commission Decision 2007/777/EC and co meat constituents which meet the criteria indicated below:					
Par		Species(A)	Treatment (B)	Origin(C) Approved I	Establishment(s)(D)				
	_								
			(A)	bladders and intestine bison, Bubalus bubalis and goats (Capra hircu asinus and their crossh domestic rabbits, PFG farmed non-domestic a domestic animals othe	relevant species of meat prod s where BOV = domestic bovir and their crossbreds): OVI = c s); EQI = domestic equine anir oreds), POR = domestic porcine = domestic poultry and farmed animals other than suidae and r than suidae and solipeds; SU n-domestic solipeds, WL = wil	e animals (Bos taurus, Bison lomestic sheep (Ovis aries) nals (Equus caballus, Equus e animals (Sus scrofa); RM = d feathered game, RUF solipeds; RUW = wild non- W = wild non-domestic			
			(B)		for the required treatment as ex II to Decision 2007/777/EC.	specified and defined in			
			(C)	Insert the ISO code of t stomachs, bladders and 2007/777/EC and, in the relevant meat constitu Decision 2007/777/EC of	the country of origin of the me d intestines as listed in Annex e case of regionalization by re- ents, the region as indicated in or Great Britain.	II, Part 2 to Decision tained EU law for the n Part 1 of Annex II to			
				, .	of the meat products must be c	ç			
					s the country of export in box	1.7,			
				— Great Brita					
				meat produ Decision 20 product is j	ntry or parts thereof authorise ucts treated with treatment A a 007/777/EC, where the third co produced is also authorised to ucts treated with that treatmen	as set out in Annex II to untry where the composite export to Great Britain			
			(D)		er of the establishments of orig ders and intestines contained port to GB.	_			

	II. Health information							
			and/or intestines used i	rom bovine, ovine or caprine in the preparation of the mea ect to the following conditions of origin:	t products and/or treated			
uc	(1)	○ [(El)	-	country or a region classified in accordance with Decision ountry or region posing a negligible BSE risk:				
Part II: Certification			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 orig do not contain and are not derived from specified risl material as defined in point 1 of Annex V to Regulatio (EC) No 999/2001 of the European Parliament and of t Council (11);				
			3.	the products of bovine, ovine do not contain and are not de separated meat obtained from or caprine animals, except for and caprine animal origin de were born, continuously rear country or region classified i 2007/453/EC as a country or r BSE risk in which there have cases;	erived from mechanically m the bones of bovine, ovine or products of bovine, ovine erived from animals that red and slaughtered in a n accordance with Decision region posing a negligible			
				the animals from which the p and caprine animal origin ar slaughtered after stunning by the cranial cavity or killed by slaughtered by laceration aft nervous tissue by means of a instrument introduced into t the animals were born, conti slaughtered in a country or r accordance with Decision 200 region posing a negligible BS	e derived, were not y means of gas injected into y the same method or er stunning of central n elongated rod-shaped he cranial cavity, except if nuously reared and egion classified in 07/453/EC as a country or			
			5.	if the animals, from which th and caprine animal origin ar country or region classified i 2007/453/EC as posing an und animals were not fed with m greaves, as defined in the Ter of the World Organisation fo products were produced and ensures that it did not contai with nervous and lymphatic deboning process.]	e derived, originate from a n accordance with Decision letermined BSE risk, those eat-and-bone meal or crestrial Animal Health Code r Animal Health, and the handled in a manner which n and was not contaminated			
	(1)	or • [(E.2)		ntry or a region classified in a cy or region posing a controll				
				the animals, from which the and caprine animal origin ar mortem and post mortem ins after stunning by laceration means of an elongated rod-sl introduced into the cranial cavit	e derived, have passed ante spection and were not killed of central nervous tissue by naped instrument avity, or by means of gas			

ROPEAN UNION						ODIIC	2088E (v1.3)
II. Health information							
(1)(4)			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, or mechanically separated meat obtained from the bones of bovine, ovine or caprine animals			
(1)(4)			3.	country or treated int	a region with estines have b	a negligible BSE risk	, imports of
				(a)	accordance v	vith Decision 2007/45	53/EC as a
				(b)	bovine, ovine derived, wer slaughtered i negligible BS	e and caprine animal e born, continuously n the country or reg E risk and have pass	l origin are reared and ion with a ed ante
(1)				(c)	or region wh	ere there have been	
					d f a d	ate from which the b eeding of ruminants nd-bone meal and gr erived from ruminat	oan on the with meat- reaves
					c c s ii	aprine animal origin ontain and are not d pecified risk materia n point 1 of Annex V	do not erived from l as defined to
(1) or	0 [(E.3)						Decision
		1.	animal orig greaves de Animal hea	gin are deriv rived from a alth Code of	ved, were not ruminants, as the World Or	fed meat-and-bone n defined in the Terres ganisation for Anima	neal or strial ll Health,
		2.	animal orig laceration shaped ins	gin are deriv of central ne trument int	ved, were not ervous tissue roduced into t	killed, after stunning by means of an elong	g, by gated rod-
		3.			, ovine and ca	prine animal origin a	are not
			(a)	-		_	f Annex V to
			(b)	nervous ar deboning	nd lymphatic t	issues exposed durin	g the
			(c)	mechanically separated meat obtained from bones obvine, ovine or caprine animals.			
	(1)(4)	(1)(4)	 (1)(4) (1) or ○ [(E.3) for imports 2007/453/E 1. 2. 	 (1)(4) (1) or o [(E.3) for imports from a coul 2007/453/EC as a count 1. the animal origing reaves de Animal hea and have p 2. the animal origing reaves de Animal hea and have p 2. the animal origing reaves de Animal o	 (1)(4) (1)(4) (1)(4) (1)(4) (1)(4) (1) (1)	 (1) (2) (3) (4) (4) (5) (6) (7) (7) (7) (8) (9) (9) (9) (9) (9) (9) (9) (9) (9) (9)	 the products of bovine, ovine and caprine and on to contain and are not derived from spectrum terial as defined in point 1 of Annex V to 5 (C(C) No 999/2001, or mechanically separated obtained from the bones of bovine, ovine or a animals (1)(4) In the case of intestines originally sourced for country or a region with a negligible BSE risk treated intestines have been subject to the followine, ovine and accordance with Decision 2007/45 (country or region posing a contro risk: (b) the country or region posing a contro risk: (c) (d) (f) (c) (f) (f)

_	LU	ROPEAN UNION		GBHC088E (V1.3)			
		II. Health information					
		(1)(4) 4.	regio	he case of intestines originally sourced from a country or a on with a negligible BSE risk, imports of treated intestines have n subject to the following conditions:			
	uo		(a)	the country or region was classified in accordance with Decision 2007/453/EC as a country or region posing a undetermined BSE risk;			
	Part II: Certification		(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;			
	Part	(1)	(c)	if the intestines are sourced from a country or region where there have been BSE indigenous cases:			
				 the animals were born after the date from which the ban on the feeding of ruminants with meat-and-bone meal and greaves derived from ruminants was enforced; or 			
				 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.] 			
		(1) and/or D Processed dairy products (6) in an amount of half or more of the substance of the composit [II.2.B product or not shelf stable dairy products in any quantity that:					
		(a)	have been produced in the country in the establishment (approval number of the establishment(s) of origin of the dairy products contained in the composite product authorised at the time of production for export of dairy products to Great Britain).				
				country of origin of the dairy products must be one of the wing:			
			-	the same as the country of export in box 1.7,			
			-	Great Britain.			
			-	a third country authorised to export to Great Britain milk and dairy products in Column A or B of Annex I to Regulation (EU) No 605/2010, where the third country where the composite product is produced is also authorised under the same conditions, to export to Great Britain milk and dairy products.			
		(EU	The country of origin indicated in box I.7 must be listed in Annex I to Regulation (EU) No 605/2010 and the treatment applied must conform to the treatment provided for in that list for the relevant country;				
		(b)	hav	e 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			
			(iii)	subject to regular veterinary inspections to ensure that they satisfy the animal health conditions laid down in Chapter I of Section IX of Annex III to Regulation (EC) No 853/2004 and in Directive 2002/99/EC;			
		(c)	are	dairy products made from raw milk obtained from:			

II. Health information (1) either o [cows, ewes, goats or buffaloes and prior to import into the territory of Great Britain have undergone or been produced from raw milk which has undergone (1) either \circ [a pasteurisation treatment involving a single heat treatment with a heating effect at least equivalent to that achieved by a pasteurisation process of at least 72°C Part II: Certification for 15 seconds and where applicable, sufficient to ensure a negative reaction to an alkaline phosphatase test applied immediately after the heat treatment;] (1) or \circ [a sterilisation process, to achieve an F0 value equal to or greater than three;] (1) or \circ [an ultra-high temperature (UHT) treatment at not less than 135°C in combination with a suitable holding time;] or \circ [a high temperature short time pasteurisation (1)treatment (HTST) at 72°C for 15 seconds or a treatment with an equivalent pasteurisation effect, applied to milk with a pH lower than 7.0 achieving, where applicable, a negative reaction to an alkaline phosphatase test] or \circ [a high temperature short time pasteurisation (1)treatment (HTST) at 72°C for 15 seconds, or a treatment with an equivalent pasteurisation effect, applied twice to milk with a pH equal to or greater than 7,0 achieving, where applicable, a negative reaction to an alkaline phosphatase test, immediately followed by (1)either \circ [lowering the pH below 6 for one hour:1 or o [additional heating equal to or greater (1)than 72°C, combined with desiccation;]] (1)or \circ [animals other than cows, ewes, goats or buffaloes and prior to import into the territory of Great Britain have undergone or been produced from raw milk which has undergone either o [a sterilisation process, to achieve an F0 value (1) equal to or greater than three;] or \circ [an ultra-high temperature (UHT) treatment at not (1)less than 135°C in combination with a suitable holding time;]] (d) were produced on or between and (7).] (1) and/or D Processed fishery products that originate from the approved establishment No (8) [II.2.C situated in the country (9) (1) and/or \Box Processed egg products that originate from the approved county (9) 1 [II.2.D were produced from eggs coming from an establishment which satisfies the requirements of Section X of Annex III to Regulation (EC) No 853/2004 which at the date of issue of the certificate is free from highly pathogenic avian influenza as defined in Regulation (EC) No 798/2008 and either (1)II.2.D.1 \circ [within a 10 km radius of which \Box [including, where appropriate, the territory of a neighbouring country,] there has been no outbreak of highly pathogenic avian influenza or Newcastle disease for at least the previous 30 days.] Or

(GB) Composite products intended for human consumption / GBHC088E (v1.3)

	II. Health information						
	(1)	II.2.D.2	○ [the egg	g products were processed:			
	(1)		either	\circ [liquid egg white was treated:		ited:	
	(1)			either • [with 55.6°C for 870 seconds.]		or 870 seconds.]	
	(1)			or	○ [with 56.7°C for 232 seconds.]		
Certification	(1)		or	\circ [10% salted yolk was treated with 62.2°C for 138 seconds.]		ed with 62.2°C for 138	
lific:	(1)		or	\circ [dried egg white was treated:		ed:	
ert	(1)			either	○ [with 67°C for	20 hours.]	
				or	○ [with 54.4°C fe	or 513 hours.]	
Part II:	(1)		or	○ [whole e	eggs were at least	treated:	
۳ ۵	(1)			either	○ [with 60°C for	188 seconds.]	
	(1)			or	 [completely completely completely] 	ooked.]	
		JUMP>	🗆 [whole	egg blends v	vere at least treate	ed]:	
	(1)			either	○ [with 60°C for	188 seconds.]	
	(1)			or	○ [with 61.1°C f	or 94 seconds.]	

	II. Health info	rmation								
	Notes									
	(*) Those countries subject to the transitional import arrangements include: an EU member State; Liechtenstein;									
	-	eland and S								
lon	References to European Union legislation within this certificate are references to direct EU legislation which has been retained in Great Britain (retained EU law as defined in the European Union (Withdrawal) Act 2018).									
icati	References to Great Britain in this certificate include Channel Islands and Isle of Man.									
ertif	Part I:	urt I:								
Part II: Certification	_	Box reference I.7:	Insert the ISO code of the country of origin of the composite product containing meat product, treated stomachs, bladders and intestines as listed in Annex II, Part 2 to Decision 2007/777/EC and/or for processed dairy products in Annex II to Commission Regulation (EU) No 605/2010 and/or for processed fishery products in Annex I and II to Commission Implementing Regulation (EU) 2019/626 and/or for processed egg products in Annex I part 1 to Commission Regulation (EC) No 798/2008.							
	_	Box reference I.11:	Name, address and registration/app production of the composite produ- same as the country of origin in bo	ct(s). Name of the country of						
		Box reference	Registration number (railway wage		•					
		I.15:	(aircraft) or name (ship). In the cas containers and their registration n must be indicated in box I.19. In ca the border control post of introduc	umber and where there is a s se of unloading and reloading	erial number of the seal it					
	_	Box reference I.16:	Do not use this box until the end of the transitional staging period.							
	_	Box reference I.19:	For containers or boxes, the contai included.	ner number and the seal nun	ıber (if applicable) must be					
	_	Box reference I.25:	Indicate total gross weight and total net weight.							
	_	Box reference I.28:	Use the appropriate Harmonised S as: 16.01; 16.02; 16.03; 16.04; 16.05;	2	e					
		Box reference I.28:	Manufacturing plant: insert the na establishments of production of the composite products containing me indicate "meat product". "treated s product containing dairy products containing processed fishery product composite product containing egg	e composite product(s). Natur at products, treated stomachs tomachs", "bladders" or "intes indicate "dairy product". In c acts specify whether aquacult	e of commodity in case of , bladders and intestines stines". In case of composite ase of composite product ure or wild origin. In case of					
	Part II:									
	(1)	Keep as app								
	(2)	stomachs, k	ucts as laid down in point 7.1 of An bladders and intestines as laid dowr ndergone one of the treatments laid	n in point 7.9 of Annex I to Reg	gulation (EC) No 853/2004					
	(3)	three whole	derogation from point 4, carcasses, esale cuts, and quarters containing cluding dorsal root ganglia, may be	no specified risk material oth						
		bovine anii	oval of the vertebral column is not r mals containing vertebral column sl on (EC) No 1760/2000.							

	II. Health info	rmation									
		The number of bovine carcasses or wholesale column is required as well as the number whe be added to the document referred to in Articl	ere removal of the vertebral co	olumn is not required shall							
	(4)	Only applicable to imports of treated intestine	28.								
cation	(5)	By way of derogation from point 3, carcasses, half carcasses or half carcasses cut into no more than three wholesale cuts, and quarters containing no specified risk material other than the vertebral column, including dorsal root ganglia, may be imported.									
Part II: Certification		When removal of the vertebral column is not required, carcasses or wholesale cuts of carcasses of bovine animals containing vertebral column shall be identified by a clearly visible blue stripe on the label referred to in Regulation (EC) No 1760/2000.									
Part]		Specific information on the number of bovine carcasses or wholesale cuts of carcasses, from which removal of the vertebral column is required and from which removal of the vertebral column is not required shall be added to the document referred to in Article 56 of Regulation (EU) 2017/625 in case of imports.									
	(6)	Raw milk and dairy products means, raw milk point 7.2 of Annex I to Regulation (EC) No 853/		n consumption as defined in							
	(7)	Date or dates of production. Imports of raw m obtained either prior to the date of authorisati part thereof mentioned under I.7 and I.8, or du adopted by Great Britain against imports of ra thereof.	ion for exportation to Great Br uring a period where restriction	ritain of the third country or ve measures have been							
	(8)	Number of the fishery product establishment a	authorised to export to Great	Britain.							
	(9)	Country of origin authorised to export to Grea	t Britain.								
	(10)	In case of composite products containing only Inspector can be accepted.	v egg or fishery products the si	gnature of an official							
	(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 other than those embossed or watermark.	to that of the printing. The sar	ne rule applies to stamps							
	Certifying Offi	Cer									
	Name (in cap		Qualification and title Signature								
	Date of signa Stamp	ture									